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Intranasal steroids versus placebo or no intervention for chronic rhinosinusitis

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ABSTRACT

Background
This review is one of six looking at the primary medical management options for patients with chronic rhinosinusitis.

Chronic rhinosinusitis is common and is characterised by inflammation of the lining of the nose and paranasal sinuses leading to nasal blockage, rhinorrhoea, facial pressure/pain and loss of sense of smell. The condition can occur with or without nasal polyps. The use of topical (intranasal) corticosteroids has been widely advocated for the treatment of chronic rhinosinusitis given the belief that inflammation is a major component of this condition.

Objectives
To assess the effects of intranasal corticosteroids in people with chronic rhinosinusitis.

Search methods
The Cochrane ENT Information Specialist searched the Cochrane ENT Trials Register; Central Register of Controlled Trials (CENTRAL 2015, Issue 8); MEDLINE; EMBASE; ClinicalTrials.gov; ICTRP and additional sources for published and unpublished trials. The date of the search was 11 August 2015.

Selection criteria
Randomised controlled trials (RCTs) with a follow-up period of at least three months comparing intranasal corticosteroids (e.g. beclomethasone dipropionate, triamcinolone acetonide, flunisolide, budesonide) against placebo or no treatment in patients with chronic rhinosinusitis.

Data collection and analysis
We used the standard methodological procedures expected by Cochrane. Our primary outcomes were disease-specific health-related quality of life (HRQL), patient-reported disease severity and the commonest adverse event - epistaxis. Secondary outcomes included general HRQL, endoscopic nasal polyp score, computerised tomography (CT) scan score and the adverse events of local irritation or other systemic adverse events. We used GRADE to assess the quality of the evidence for each outcome; this is indicated in italics.
Main results

We included 18 RCTs with a total of 2738 participants. Fourteen studies had participants with nasal polyps and four studies had participants without nasal polyps. Only one study was conducted in children.

Intranasal corticosteroids versus placebo or no intervention

Only one study (20 adult participants without polyps) measured our primary outcome disease-specific HRQL using the Rhinosinusitis Outcome Measures-31 (RSOM-31). They reported no significant difference (numerical data not available) (very low quality evidence).

Our second primary outcome, disease severity, was measured using the Chronic Sinusitis Survey in a second study (134 participants without polyps), which found no important difference (mean difference (MD) 2.84, 95% confidence interval (CI) -5.02 to 10.70; scale 0 to 100). Another study (chronic rhinosinusitis with nasal polyps) reported an increased chance of improvement in the intranasal corticosteroids group (RR 2.78, 95% CI 1.76 to 4.40; 109 participants). The quality of the evidence was low.

Six studies provided data on at least two of the individual symptoms used in the EPOS 2012 criteria to define chronic rhinosinusitis (nasal blockage, rhinorrhoea, loss of sense of smell and facial pain/pressure). When all four symptoms in the EPOS criteria were available on a scale of 0 to 3 (higher = more severe symptoms), the average MD in change from baseline was -0.26 (95% CI -0.37 to -0.15; 243 participants; two studies; low quality evidence). Although there were more studies and participants when only nasal blockage and rhinorrhoea were considered (MD -0.31, 95% CI -0.38 to -0.24; 1702 participants; six studies), the MD was almost identical to when loss of sense of smell was also considered (1345 participants, four studies; moderate quality evidence).

When considering the results for the individual symptoms, benefit was shown in the intranasal corticosteroids group. The effect size was larger for nasal blockage (MD -0.40, 95% CI -0.52 to -0.29; 1702 participants; six studies) than for rhinorrhoea (MD -0.25, 95% CI -0.33 to -0.17; 1702 participants; six studies) or loss of sense of smell (MD -0.19, 95% CI -0.28 to -0.11; 1345 participants; four studies). There was heterogeneity in the analysis for facial pain/pressure (MD -0.27, 95% CI -0.56 to 0.02; 243 participants; two studies). The quality of the evidence was moderate for nasal blockage, rhinorrhoea and loss of sense of smell, but low for facial pain/pressure.

There was an increased risk of epistaxis with intranasal corticosteroids (risk ratio (RR) 2.74, 95% CI 1.88 to 4.00; 2508 participants; 13 studies; high quality evidence).

Considering our secondary outcome, general HRQL, one study (134 participants without polyps) measured this using the SF-36 and reported a statistically significant benefit only on the general health subscale. The quality of the evidence was very low.

It is unclear whether there is a difference in the risk of local irritation (RR 0.94, 95% CI 0.53 to 1.64; 2124 participants; 11 studies) (low quality evidence).

None of the studies treated or followed up patients long enough to provide meaningful data on the risk of osteoporosis or stunted growth (children).

Other comparisons

We identified no other studies that compared intranasal corticosteroids plus co-intervention A versus placebo plus co-intervention A.

Authors’ conclusions

Most of the evidence available was from studies in patients with chronic rhinosinusitis with nasal polyps. There is little information about quality of life (very low quality evidence). For disease severity, there seems to be improvement for all symptoms (low quality evidence), a moderate-sized benefit for nasal blockage and a small benefit for rhinorrhoea (moderate quality evidence). The risk of epistaxis is increased (high quality evidence), but these data included all levels of severity; small streaks of blood may not be a major concern for patients. It is unclear whether there is a difference in the risk of local irritation (low quality evidence).

Plain language summary

Intranasal steroids versus placebo or no intervention for chronic rhinosinusitis

Review question

We reviewed the evidence for the benefits and harms of intranasal (in the nose) steroids given to people with chronic rhinosinusitis.
Background

Chronic rhinosinusitis is a common condition that is defined as inflammation of the nose and paranasal sinuses (a group of air-filled spaces behind the nose, eyes and cheeks). Patients with chronic rhinosinusitis experience at least two or more of the following symptoms for at least 12 weeks: blocked nose, discharge from their nose or runny nose (rhinorrhoea), pain or pressure in their face and/or a reduced sense of smell (hyposmia). Some people will also have nasal polyps, which are grape-like swellings of the normal nasal lining inside the nasal passage and sinuses. Topical (intranasal) corticosteroids are used with the aim of reducing inflammation in order to improve patient symptoms.

Study characteristics

We included 18 randomised controlled trials (RCTs) with a total of 2738 participants in this review. Most studies were relatively small, with as few as 9 or 10 patients per intervention arm. The largest study had 748 patients in total. Most were conducted in tertiary referral centres in northern Europe, the US and Canada. Fourteen studies only included participants with chronic rhinosinusitis with nasal polyps and four studies had participants without nasal polyps. Only one study was conducted in children. The studies looked at a range of types, doses and methods of administration (e.g. spray, drops) of intranasal corticosteroids.

Key results and quality of the evidence

One study (20 participants) reported no statistically significant difference in disease-specific health-related quality of life. Another measured general health-related quality of life and reported a statistically significant benefit only on a subscale for general health. Both studies recruited participants with chronic rhinosinusitis without nasal polyps. The quality of the evidence was very low (we have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of effect).

Disease severity was measured in one study (chronic rhinosinusitis without nasal polyps, 134 participants), which found no important difference. Another study (chronic rhinosinusitis with nasal polyps) reported an increased chance of improvement in the intranasal corticosteroids group. The quality of the evidence was low (our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect).

When each type of symptom was measured separately (nasal blockage, rhinorrhoea, loss of sense of smell, facial pain/pressure), benefit was shown in the intranasal corticosteroids group. The quality of the evidence was moderate for nasal blockage, rhinorrhoea and loss of sense of smell, but low for facial pain/pressure (moderate quality evidence means we are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different). The risk of nosebleeds is increased (high quality evidence), but this included all levels of severity; for some patients small streaks of blood may not be a major concern. It is unclear whether there is a difference in the risk of local irritation (low quality evidence).

Conclusions

Most of the evidence available was from studies in patients with chronic rhinosinusitis with nasal polyps. There is little information about quality of life and the quality of this evidence is very low. For disease severity, there seems to be improvement for all symptoms (low quality evidence), a moderate-sized benefit for nasal blockage and a small benefit for rhinorrhoea (moderate quality evidence). The risk of nosebleeds is increased (high quality evidence), but this included all levels of severity; for some patients small streaks of blood may not be a major concern. It is unclear whether there is a difference in the risk of local irritation (low quality evidence).
<table>
<thead>
<tr>
<th>Outcomes of participants (studies)</th>
<th>Relative effect (95% CI)</th>
<th>Anticipated absolute effects* (95% CI)</th>
<th>Quality</th>
<th>What happens</th>
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<tr>
<td></td>
<td></td>
<td>With placebo</td>
<td>With intranasal corticosteroids</td>
<td>Difference</td>
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<tr>
<td>Disease-specific HRQL measured as median change from baseline (RSOM-31)</td>
<td>-</td>
<td>Median 5 points lower</td>
<td>Median 62 points lower</td>
<td>-</td>
</tr>
<tr>
<td>Disease severity - measured as change from baseline using the Chronic Sinusitis Survey at 20 weeks</td>
<td>-</td>
<td>Chronic Sinusitis Survey score was 7.35</td>
<td>-</td>
<td>MD 2.84 higher (5.02 lower to 10.7 higher) than placebo</td>
</tr>
<tr>
<td>Disease severity - analysed as the proportion of patients who reported improvement on a global symptom score</td>
<td>RR 2.78 (1.76 to 4.40)</td>
<td>Study population</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Disease severity measured as average change from baseline at 12 to 20 weeks (range 0 to 3 points) for a combination of symptoms

- **All 4 EPOS domains**
  - Number of participants: 243 (2 RCTs)
  - MD: 0.26 lower (0.37 lower to 0.15 lower) LOW
  - The improvement in the intranasal corticosteroids group was higher (moderate effect size). Lower score = less severe symptoms

- **3 EPOS domains - nasal blockage, rhinorrhoea, loss of sense of smell**
  - Number of participants: 1345 (4 RCTs)
  - MD: 0.31 lower (0.38 lower to 0.23 lower) MODERATE
  - The improvement in the intranasal corticosteroids group was higher (moderate effect size)

- **2 EPOS domains - nasal blockage and rhinorrhoea**
  - Number of participants: 1702 (6 RCTs)
  - MD: 0.31 lower (0.38 lower to 0.24 lower) MODERATE
  - The improvement in the intranasal corticosteroids group was higher (moderate effect size)

## Disease severity measured as average change from baseline at 12 to 20 weeks (range 0 to 3 points) for individual symptoms

- **Nasal blockage**
  - Number of participants: 1702 (6 RCTs)
  - MD: 0.4 lower (0.52 lower to 0.29 lower) MODERATE
  - The improvement in the intranasal corticosteroids group was higher (moderate effect size)
<table>
<thead>
<tr>
<th>Outcome</th>
<th>Number of Participants</th>
<th>MD or RR</th>
<th>Comparison</th>
<th>Result</th>
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</thead>
<tbody>
<tr>
<td>Rhinorrhea</td>
<td>1702 (6 RCTs)</td>
<td>0.25 lower (0.33 lower to 0.17 lower)</td>
<td>MODERATE</td>
<td>The improvement in the intranasal corticosteroids group was higher (small effect size)</td>
</tr>
<tr>
<td>Loss of sense of smell</td>
<td>1345 (4 RCTs)</td>
<td>0.19 lower (0.28 lower to 0.11 lower)</td>
<td>MODERATE</td>
<td>The improvement in the intranasal corticosteroids group was higher (small effect size)</td>
</tr>
<tr>
<td>Facial pain/pressure</td>
<td>243 (2 RCTs)</td>
<td>0.27 lower (0.56 lower to 0.02 higher)</td>
<td>LOW</td>
<td>The improvement in the intranasal corticosteroids group was higher (moderate effect size)</td>
</tr>
</tbody>
</table>

Note: MD = Mean Difference; RR = Risk Ratio; SD = Standard Deviation; SD = Standard Error; CI = Confidence Interval; **= p < 0.05; *** = p < 0.01; **** = p < 0.001; n.s. = not significant; HRQL = Health-related quality of life; SF-36 = Short-Form 36-Item Health Survey; RCT = Randomized Controlled Trial.
### Local irritation of participants: 2124 (11 RCTs)

<table>
<thead>
<tr>
<th>Study Population</th>
<th>Adverse events - local irritation (per 1000)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>RR 0.94 (0.53 to 1.64)</td>
</tr>
<tr>
<td>Moderate</td>
<td>26 per 1000 (4 to 42)</td>
</tr>
<tr>
<td>Low</td>
<td>21 per 1000 (11 to 34)</td>
</tr>
<tr>
<td>Moderate</td>
<td>24 per 1000 (14 to 42)</td>
</tr>
<tr>
<td>Low</td>
<td>1 fewer per 1000 (12 fewer to 16 more)</td>
</tr>
<tr>
<td>Moderate</td>
<td>2 fewer per 1000 (14 fewer to 18 more)</td>
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</table>

**GRADE Working Group grades of evidence**

- **High quality**: We are very confident that the true effect lies close to that of the estimate of the effect.
- **Moderate quality**: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different from the estimate of the effect.
- **Low quality**: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect.
- **Very low quality**: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of the effect.

#### Adverse events - local irritation

- **OR**: odds ratio; **CI**: confidence interval; **HRQL**: health-related quality of life; **EPOS**: European Position Paper on Rhinosinusitis and Nasal Polyps 2012; **HR**: hazard ratio; **RCT**: randomised controlled trial; **RR**: risk ratio.
**BACKGROUND**

**Description of the condition**

Chronic rhinosinusitis is defined as inflammation of the nose and paranasal sinuses characterised by two or more symptoms, one of which must be nasal blockage/obstruction/congestion or nasal discharge (anterior/posterior nasal drip). The other possible symptoms include facial pain/pressure, reduction or loss of sense of smell (in adults) or cough (in children). Symptoms must have continued for at least 12 weeks. In addition, people must have either mucosal changes within the ostiomeatal complex and/or sinuses as evidenced by a computerised tomography (CT) scan and/or endoscopic signs of at least one of the following: nasal polyps, mucopurulent discharge primarily from middle meatus or oedema/mucosal obstruction primarily in the middle meatus (EPOS 2012). Chronic rhinosinusitis represents a common source of ill health; 11% of UK adults reported chronic rhinosinusitis symptoms in a worldwide population study (Hastan 2011). Symptoms, including nasal obstruction, nasal discharge, facial pain, anosmia and sleep disturbance, have a major impact on quality of life, reportedly greater in several domains of the SF-36 than angina or chronic respiratory disease (Gliklich 1995). Acute exacerbations, inadequate symptom control and respiratory disease exacerbation are common. Complications are rare, but may include visual impairment and intracranial infection.

Two major phenotypes of chronic rhinosinusitis have been identified based on the presence or absence of nasal polyps on examination. Nasal polyps are tumour-like hyperplastic swellings of the nasal mucosa, most commonly originating from within the ostiomeatal complex (Larsen 2004). Chronic rhinosinusitis with nasal polyps (CRSwNP) is diagnosed when polyps are seen (on direct or endoscopic examination) bilaterally in the middle meatus. The acronym CRSwNP is used for the condition in which no polyps are present. Although the aetiology of chronic rhinosinusitis is not fully understood, it may involve abnormalities in the host response to irritants, commensal and pathogenic organisms and allergens, obstruction of sinus drainage pathways, abnormalities of normal mucociliary function, loss of the normal mucosal barrier or infection. Two typical profiles may be observed with respect to inflammatory mediators; in eosinophilic chronic rhinosinusitis, which is typically associated with nasal polyps, high levels of eosinophils, immunoglobulin E (IgE) and interleukin (IL)-5 may be found, while in neutrophilic chronic rhinosinusitis, more often associated with chronic rhinosinusitis without polyps, neutrophils predominate, with elevated interferon (IFN) gamma, IL-8 and tumour necrosis factor (TNF) (EPOS 2012).

While treatment decisions should be made based on an understanding of the patient's chronic rhinosinusitis phenotype and likely aetiology, in practice treatment may be initiated without knowledge of the polyp status, particularly in primary care. This review (and most of its companion reviews) consider patients with and without polyps together in the initial evaluation of treatment effects. However, subgroup analyses explore potential differences between them.

The most commonly used interventions for chronic rhinosinusitis are used either topically (sprayed into the nose) or systemically (by mouth) and include steroids, antibiotics and saline.

**Description of the intervention**

Anti-inflammatory therapy plays a significant role in the treatment of chronic rhinosinusitis. This includes corticosteroids and low-dose macrolides. Topical corticosteroids are more widely used than oral steroids because treatment can be given for longer without significant adverse effects.

Intranasal corticosteroid therapy is often prescribed for patients with chronic rhinosinusitis, but with considerable variability in timing, frequency, dose, topical delivery method and the specific agent used (Benninger 2003; Spector 1998). The topical delivery method may affect the amount of steroid that comes into contact with the paranasal sinus mucosa (Grobler 2008; Harvey 2009). The simplest nasal delivery methods are drops, sprays, aerosols, nebulisers and atomisers. These contrast with methods involving direct sinus cannulation and nasal irrigation with squeeze bottles and neti pots, which are likely to provide better delivery to the sinuses, especially in the post-sinus surgery setting (Grobler 2008; Harvey 2009; Thomas 2013).

Classes of topical corticosteroid include first-generation intranasal steroids (beclomethasone dipropionate, triamcinolone acetonide, flunisolide and budesonide) and newer preparations (fluticasone propionate, mometasone furoate, ciclesonide and fluticasone furoate).

**How the intervention might work**

The use of topical (intranasal) corticosteroids has been widely advocated for the treatment of chronic rhinosinusitis given the belief that inflammation is a major component of this condition (Fokkens 2007; Hamilos 2000; McNally 1997). The mechanism of action is a combination of anti-inflammatory effects (for example, reducing pro-inflammatory, and increasing anti-inflammatory, gene transcription and reducing airway inflammatory cell infiltration) and suppression of the production of pro-inflammatory mediators, cell chemotactic factors and adhesion molecules (Mullol 2009). Several factors could affect the relative levels of effectiveness or harm from using intranasal corticosteroids. It has been suggested that the type of steroid, dose and method of delivery (which affects the bioavailability) may contribute to the relative effectiveness of the treatment. In addition, it is unclear whether patient characteristics will affect levels of effectiveness (i.e. whether they have polyps and whether they are adults or children). Another...
uncertainty is whether the duration of treatment is important. If heterogeneity in effects was indeed observed, we planned to explore these factors through subgroup analyses.

Why it is important to do this review

Intranasal corticosteroids are the mainstay and currently recommended treatment for chronic rhinosinusitis. This review incorporates an update of two previous Cochrane reviews (Kalish 2012; Snidvongs 2011). Unlike the previous reviews, this review focuses on the effects of intranasal corticosteroids when compared to no treatment or placebo, to establish their effectiveness in the treatment of chronic rhinosinusitis. The relative effects of different types, doses and methods of delivery are investigated in a separate review (Chong 2016a).

This review is one of a suite of reviews looking at management options for patients with chronic rhinosinusitis (Chong 2016a; Chong 2016b; Head 2016a; Head 2016b; Head 2016c), and we use the same outcome measures across the reviews. We have not included studies designed to evaluate interventions in the immediate peri-surgical period, which are focused on assessing the impact of the intervention on the surgical procedure or on modifying the post-surgical results (preventing relapse).

OBJECTIVES

To assess the effects of intranasal corticosteroids in people with chronic rhinosinusitis.

METHODS

Criteria for considering studies for this review

Types of studies

We included studies with the following design characteristics:
- randomised controlled trials, including cluster-randomised trials and quasi-randomised trials (cross-over trials were only to be included if the data from the first phase were available); and
- patients were followed up for at least two weeks.

We excluded studies with the following design characteristics:
- randomised patients by side of nose (within-patient controlled) because it is difficult to ensure that the effects of any of the interventions considered can be localised; or
- perioperative studies, where the sole purpose of the study was to investigate the effect of intranasal corticosteroids on surgical outcome.

Types of participants

Patients with chronic rhinosinusitis, whether with or without polyps.

We excluded studies that included a majority of patients with:
- cystic fibrosis;
- allergic fungal sinusitis/ eosinophilic fungal/mucinous rhinosinusitis;
- aspirin-exacerbated respiratory disease;
- antrochoanal polyps (benign polyps originating from the mucosa of the maxillary sinus);
- malignant polyps;
- primary ciliary dyskinesia;
- a history of surgery for nasal polyps within six weeks of entry to the study.

Types of interventions

All intranasal corticosteroids; this included nasal sprays and nasal drops.

First-generation intranasal corticosteroids:
- Beclomethasone dipropionate
- Triamcinolone acetonide
- Flunisolide
- Budesonide

Second-generation intranasal corticosteroids:
- Ciclesonide
- Fluticasone furoate
- Fluticasone propionate
- Mometasone furoate
- Betamethasone sodium phosphate

If other interventions were used, these should have been used in both treatment arms. Allowed co-interventions included:
- nasal saline irrigation;
- antibiotics;
- intermittent nasal decongestants.

The comparators were placebo or no intervention.

The main comparison pair was:
- intranasal corticosteroids versus placebo or no intervention.

Other possible comparison pairs included:
- intranasal corticosteroids plus co-intervention A versus placebo plus co-intervention A.

This review is part of a larger series of six reviews for the treatment of chronic rhinosinusitis.
- Intranasal steroids versus placebo or no intervention for chronic rhinosinusitis (this review).
- Different types of intranasal steroids for chronic rhinosinusitis (Chong 2016a). This review compares different classes, doses and delivery methods of intranasal corticosteroids for chronic rhinosinusitis.
• Short-course oral steroids alone for chronic rhinosinusitis (Head 2016a). This review compares short-course oral steroids alone with placebo or no intervention, or against other pharmacological interventions such as antibiotics or nasal saline irrigation.
• Short-course oral steroids as an adjunct therapy for chronic rhinosinusitis (Head 2016b). This review compares oral steroids where they have been used as add-on therapy to other treatments for chronic rhinosinusitis (such as intranasal corticosteroids, antibiotics, or saline solution).
• Saline irrigation for chronic rhinosinusitis (Chong 2016b). This review compares nasal saline irrigation for chronic rhinosinusitis with both placebo/no intervention and with intranasal corticosteroids or antibiotics.
• Systemic and topical antibiotics for chronic rhinosinusitis (Head 2016c). This review compares both topical and systemic antibiotics with placebo/no treatment, two different antibiotics with each other and antibiotics with intranasal corticosteroids.

Types of outcome measures
We analysed the following outcomes in the review, but we did not use them as a basis for including or excluding studies.

Primary outcomes
• Health-related quality of life, using *disease-specific* health-related quality of life scores, such as the Sino-Nasal Outcome Test-22 (SNOT-22), Rhinosinusitis Outcome Measures-31 (RSOM-31) and SNOT-20.
• Disease severity, as measured by patient-reported symptom score (such as the Chronic Sinusitis Survey (CSS) questionnaire and visual analogue scales).
• Significant adverse effect: epistaxis.

Secondary outcomes
• Health-related quality of life, using *generic* quality of life scores, such as the SF-36, EQ-5D and other well-validated instruments.
• Other local adverse effects: local irritation (including oral thrush, sore throat and other local nasal irritation such as dryness, itchiness etc.).
• Other systemic adverse effects:
  - in children - stunted growth (minimum time point: six months of treatment and follow-up);
  - in adults - osteoporosis.
• Nasal endoscopic score (depending on population, either nasal polyps size score or endoscopy score, e.g. Lund-Mackay/Lund-Kennedy).
• Computerised tomography (CT) scan score (e.g. Lund-Mackay).

Outcomes were measured at three to six months, six to 12 months and more than 12 months. For adverse events, we analysed data from the longest time periods.

Search methods for identification of studies
The Cochrane ENT Information Specialist conducted systematic searches for randomised controlled trials and controlled clinical trials. There were no language, publication year or publication status restrictions. The date of the search was 11 August 2015.

Electronic searches
The Information Specialist searched:
• the Cochrane Register of Studies ENT Trials Register (searched 11 August 2015);
• the Cochrane Central Register of Controlled Trials (CENTRAL 2015, Issue 7);
  - Ovid MEDLINE (1946 to July week 5 2015);
    - Ovid MEDLINE (In-Process & Other Non-Indexed Citations) (searched 11 August 2015);
    - PubMed (as a top up to searches in Ovid MEDLINE) (searched 11 August 2015);
• Ovid EMBASE (1974 to 2015 week 32);
• ClinicalTrials.gov, www.clinicaltrials.gov (search via the Cochrane Register of Studies) (searched 11 August 2015);
• World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP) (searched 11 August 2015);
• Google Scholar (searched 11 August 2015).

The Information Specialist modelled subject strategies for databases on the search strategy designed for CENTRAL. Where appropriate, they were combined with subject strategy adaptations of the highly sensitive search strategy designed by Cochrane for identifying randomised controlled trials and controlled clinical trials (as described in the *Cochrane Handbook for Systematic Reviews of Interventions* Version 5.1.0, Box 6.4.b. (*Handbook 2011*). Search strategies for major databases including CENTRAL are provided in Appendix 1.

Searching other resources
We scanned the reference lists of identified publications for additional trials and contacted trial authors where necessary. In addition, the Information Specialist searched PubMed, *The Cochrane Library* and Google to retrieve existing systematic reviews relevant to this systematic review, so that we could scan their reference lists for additional trials.

Data collection and analysis
Selection of studies
At least two review authors independently screened all titles and abstracts of the studies obtained from the database searches to identify potentially relevant studies. At least two review authors evaluated the full text of each potentially relevant study to determine if it met the inclusion and exclusion criteria for this review. We resolved any differences by discussion and consensus, with the involvement of a third author for clinical and methodological input where necessary.

Data extraction and management
Two review authors independently extracted data from each study using a standardised data collection form (see Appendix 2). Whenever a study had more than one publication, we retrieved all publications to ensure complete extraction of data. Where there were discrepancies in the data extracted by different review authors, we checked these against the original reports and resolved differences by discussion and consensus, with the involvement of a third author or a methodologist where appropriate. We contacted the original study authors for clarification or for missing data whenever possible. If differences were found between publications of a study, we contacted the original authors for clarification. We used data from the main paper(s) if no further information was found.

We included key characteristics of the studies, such as study design, setting, sample size, population and how outcomes were defined or collected in the studies. In addition, we also collected baseline information on prognostic factors or effect modifiers. For this review, this included:
- presence or absence of nasal polyps;
- baseline polyp score (where appropriate);
- whether the patient has had previous sinus surgery.

For the outcomes of interest to the review, we extracted the findings of the studies on an available case analysis basis; i.e. we included data from all patients available at the time points based on the treatment randomised whenever possible, irrespective of compliance or whether patients had received the treatment as planned. In addition to extracting pre-specified information about study characteristics and aspects of methodology relevant to risk of bias, we extracted the following summary statistics for each trial and each outcome:
- For continuous data: the mean values, standard deviations and number of patients for each treatment group. Where endpoint data were not available, we extracted the values for change from baseline. We analysed data from measurement scales such as SNOT-22 and EQ-5D as continuous data.
- For binary data: the numbers of participants experiencing an event and the number of patients assessed at the time point.
- For ordinal scale data: if the data appeared to be approximately normally distributed or if the analysis that the investigators performed suggested parametric tests were appropriate, then we treated the outcome measures as continuous data. Alternatively, if data were available, we planned to convert into binary data.

We prespecified the time points of interest for the outcomes in this review. While studies may have reported data at multiple time points, we only extracted the longest available data within the time points of interest. For example, for 'short' follow-up periods, our time point was defined as three to six months post-randomisation. If a study reported data at three, four and six months, we only extracted and analysed the data for the six-month follow-up.

Assessment of risk of bias in included studies
Two review authors independently assessed the risk of bias of each included study. We followed the guidance in the Cochrane Handbook for Systematic Reviews of Interventions (Handbook 2011) and we used the Cochrane 'Risk of bias' tool. With this tool we assessed the risk of bias as 'low', 'high' or 'unclear' for each of the following six domains:
- sequence generation;
- allocation concealment;
- blinding of participants, personnel and outcome assessment;
- incomplete outcome data;
- selective reporting;
- other sources of bias.

Measures of treatment effect
We summarised the effects of dichotomous outcomes (e.g. proportion of patients with symptom resolution) as risk ratios (RR) with CIs. For the key outcomes that we presented in the 'Summary of findings' table, we also expressed the results as absolute numbers based on the pooled results and compared to the assumed risk. We also planned to calculate the number needed to treat to benefit (NNTB) using the pooled results. The assumed baseline risk is typically either (a) the median of the risks of the control groups in the included studies, this being used to represent a 'medium risk population' or, alternatively, (b) the average risk of the control groups in the included studies is used as the 'study population' (Handbook 2011). If a large number of studies were available, and where appropriate, we also planned to present additional data based on the assumed baseline risk in (c) a low-risk population and (d) a high-risk population.

For continuous outcomes, we expressed treatment effects as a mean difference (MD) with standard deviation (SD) or as standardised mean difference (SMD) if different scales had been used to measure the same outcome. We provided a clinical interpretation of the SMD values.

Unit of analysis issues
This review did not use data from phase II of cross-over studies or from studies where the patient was not the unit of randomisation, i.e. studies where the side (right versus left) was randomised.
Dealing with missing data

We tried to contact study authors via email whenever the outcome of interest was not reported, if the methods of the study suggested that the outcome had been measured. We did the same if not all data required for meta-analysis had been reported, unless the missing data were standard deviations. If standard deviation data were not available, we approximated these using the standard estimation methods from P values, standard errors or 95% CIs if these are reported as detailed in the Cochrane Handbook for Systematic Reviews of Interventions (Handbook 2011). If it was impossible to estimate these, we contacted the study authors. Apart from imputations for missing standard deviations, we conducted no other imputations. However, we carried out calculations relating to disease severity (measured by patient-reported symptom scores) as most of the data measured individual symptoms rather than using validated instruments (see ‘Imputing total symptom scores’ below). We extracted and analysed data for all outcomes using the available case analysis method.

Imputing total symptom scores

Where a paper did not present information for the total disease severity in terms of patient-reported symptom scores but did present data for the results of individual symptoms, we used the symptoms covering the important domains of the EPOS chronic rhinosinusitis diagnosis criteria (EPOS 2012) to calculate a total symptom score. The EPOS 2012 criteria for chronic rhinosinusitis require at least two symptoms. One of the symptoms must be either nasal blockage or nasal discharge; other symptoms can include facial pressure/pain, loss of sense of smell (for adults) or cough (for children). Where mean final values or changes from baseline were presented in the paper for the individual symptoms we summed these to calculate a ‘total symptom score’. We calculated standard deviations for the total symptom score as if the symptoms were independent, random variables that were normally distributed. We acknowledge that there is likely to be a degree of correlation between the individual symptoms, however we used this process because the magnitude of correlation between the individual symptoms is not currently well understood (no evidence found). If the correlation is high, the summation of variables as discrete variables is likely to give a conservative estimate of the total variance of the summed final score. If the correlation is low, this method of calculation will underestimate the standard deviation of the total score. However, the average patient-reported symptom scores have a correlation coefficient of about 0.5; if this is also applicable to chronic rhinosinusitis symptoms, the method used should have minimal impact (Balk 2012). As this method of calculation does not take into account weighting of different symptoms (no evidence found), we downgraded all the disease severity outcomes for lack of use of validated scales whenever this occurred.

Assessment of heterogeneity

We assessed clinical heterogeneity (which may be present even in the absence of statistical heterogeneity) by examining the included trials for potential differences between studies in the types of participants recruited, interventions or controls used and the outcomes measured.

We assessed statistical heterogeneity by visually inspecting the forest plots and by considering the Chi² test (with a significance level set at P value < 0.10) and the I² statistic, which calculates the percentage of variability that is due to heterogeneity rather than chance, with I² values over 50% suggesting substantial heterogeneity (Handbook 2011).

Assessment of reporting biases

We assessed reporting bias as between-study publication bias and within-study outcome reporting bias.

Outcome reporting bias (within-study reporting bias)

We assessed within-study reporting bias by comparing the outcomes reported in the published report against the study protocol, whenever this could be obtained. If the protocol was not available, we compared the outcomes reported to those listed in the methods section. If results are mentioned but not reported adequately in a way that allows analysis (e.g. the report only mentions whether the results were statistically significant or not), bias in a meta-analysis is likely to occur. We sought further information from the study authors. If no further information could be found, we noted this as being a ‘high’ risk of bias. Quite often there was insufficient information to judge the risk of bias; we noted this as an ‘unclear’ risk of bias (Handbook 2011).

Publication bias (between-study reporting bias)

We planned to assess funnel plots if sufficient trials (more than 10) were available for an outcome. If we had observed asymmetry of the funnel plot, we would have conducted more formal investigation using the methods proposed by Egger 1997.

Data synthesis

We conducted all meta-analyses using Review Manager 5.3 (RevMan 2014). For dichotomous data, we planned to analyse treatment differences as a risk ratio (RR) calculated using the Mantel-Haenszel methods. We analysed time-to-event data using the generic inverse variance method. For continuous outcomes, if all the data were from the same scale, we planned to pool mean values obtained at follow-up with change.
outcomes and report this as a MD. However, if the SMD had to be used as an effect measure, we did not plan to pool change and endpoint data.

When statistical heterogeneity is low, random-effects versus fixed-effect methods yield trivial differences in treatment effects. However, when statistical heterogeneity is high, the random-effects method provides a more conservative estimate of the difference.

Subgroup analysis and investigation of heterogeneity

We planned to conduct some subgroup analyses regardless of whether statistical heterogeneity was observed, as these are widely suspected to be potential effect modifiers. For this review, this included:

- phenotype of patients: whether patients have chronic rhinosinusitis without nasal polyps, chronic rhinosinusitis with nasal polyps, a mixed group or the status of polyps is not known or not reported. We planned to undertake the subgroup analysis because although there appears to be a considerable overlap between the two forms of chronic rhinosinusitis with regards to inflammatory profile, clinical presentation and effect of treatment (Cho 2012; DeMarcantonio 2011; Ebbens 2010; Fokkens 2007; Ragab 2004; Ragab 2010; van Drunen 2009), there is some evidence pointing to differences in the respective inflammatory profiles (Kern 2008; Keswani 2012; Tan 2011; Tomassen 2011; Zhang 2008; Zhang 2009), and potentially even differences in treatment outcome (Ebbens 2011).

We planned to present the main analyses of this review according to the subgroups of phenotypes of chronic rhinosinusitis in forest plots and all other subgroup analysis results in tables. When studies had a mixed group of patients, we analysed the study as one of the subgroups (rather than as a mixed group) if more than 80% of patients belonged to one category. For example, if 81% of patients had chronic rhinosinusitis without nasal polyps, we analysed the study as that subgroup.

In addition to polyps status, we also planned to conduct the following subgroup analyses in the presence of statistical heterogeneity:

- patient age (children versus adults);
- dose;
- duration of treatment;
- method of delivery.

Sensitivity analysis

We planned to carry out sensitivity analyses to determine whether the findings were robust to the decisions made in the course of identifying, screening and analysing the trials. We planned to conduct sensitivity analysis for the following factors, whenever possible:

- impact of model chosen: fixed-effect versus random-effects model;
- risk of bias of included studies: excluding studies with high risk of bias (we defined these as studies that had a high risk of allocation concealment bias and a high risk of attrition bias (overall loss to follow-up of 20%, differential follow-up observed);
- how outcomes were measured: we planned to investigate the impact of including data where the validity of the measurement is unclear.

If any of these investigations found a difference in the size of the effect or heterogeneity, we would mention this in the Effects of interventions section.

GRADE and 'Summary of findings' table

We used the GRADE approach to rate the overall quality of evidence for each outcome using the GDT tool (http://www.guidelinedevelopment.org/) for the main comparison pairs listed in the Types of interventions section. The quality of evidence reflects the extent to which we are confident that an estimate of effect is correct and we applied this in the interpretation of results. There are four possible ratings: 'high', 'moderate', 'low' and 'very low'. A rating of 'high' quality evidence implies that we are confident in our estimate of effect and that further research is very unlikely to change our confidence in the estimate of effect. A rating of 'very low' quality implies that any estimate of effect obtained is very uncertain.

The GRADE approach rates evidence from RCTs that do not have serious limitations as high quality. However, several factors can lead to the downgrading of the evidence to moderate, low or very low. The degree of downgrading is determined by the seriousness of these factors:

- study limitations (risk of bias);
- inconsistency;
- indirectness of evidence;
- imprecision;
- publication bias.

The 'Summary of findings' table presents only the seven top priority outcomes (disease-specific health-related quality of life, disease severity score, adverse effects and generic quality of life score). We did not include the outcomes of endoscopic score and CT scan score in the 'Summary of findings' table.
The searches retrieved a total of 2470 references after removal of duplicates. We identified one additional reference from other sources. We screened the titles and abstracts and subsequently removed 2297 references. We assessed 87 full texts for eligibility. We excluded 45 studies (55 references), with reasons. We included 18 studies (24 references). We identified three ongoing studies and there are five studies awaiting assessment because we cannot locate the full-text papers.

A flow chart of study retrieval and selection is provided in Figure 1.
Figure 1. Process for sifting search results and selecting studies for inclusion.

3770 records identified through database searching

1 additional record identified through other sources

2470 records after duplicates removed

2470 records screened

2297 records discarded

87 full-text articles assessed for eligibility

45 studies (55 articles) excluded, with reasons
5 articles awaiting assessment
3 ongoing studies

18 studies (24 articles) included in qualitative synthesis

13 studies included in quantitative synthesis (meta-analysis)
Included studies

Design

All studies included studies were double-blinded randomised controlled trials. All but three studies (Holmberg 1997; Lang 1983; Lund 2004) had a treatment and follow-up duration of between 12 to 16 weeks. Holmberg 1997 and Lund 2004 had a treatment duration of 20 and 26 weeks respectively. Some of these studies had multiple treatment arms: four compared different doses of intranasal corticosteroids against placebo (Chur 2013; Penttilla 2000; Small 2005; Stjarne 2006), whereas Johansen 1993 compared two different types of delivery methods (nasal spray versus nasal drops) against placebo.

Setting

Most studies were published more than 10 years ago (Aukema 2005; Holmberg 1997; Holopainen 1982; Johansen 1993; Keith 2000; Lang 1983; Lund 1998; Lund 2004; Parikh 2001; Penttilla 2000; Small 2005). Most of these were conducted in tertiary referral centres in northern Europe, the US and Canada. Chur 2013, Lund 2004, Small 2005 and Stjarne 2006 included other centres around the world, while Zhou 2015 was conducted in China.

Population and sample size

One study recruited 127 children aged 6 to 17 (Chur 2013); the rest were conducted in adults. Most studies were relatively small, with as few as 9 or 10 patients per intervention arm. The largest study was Zhou 2015, with 748 patients in total. The populations included in the studies with regards to phenotypes of chronic rhinosinusitis were:


Intervention and comparison

The following types of intranasal corticosteroids were used:

- Beclomethasone propionate: 400 µg per day delivered as nasal sprays in Holmberg 1997 and Lund 1998. Lang 1983 used nasal drops at 800 µg per day.
- Mometasone furoate nasal spray, given as 200 µg once daily in Small 2005, Stjarne 2006 and Stjarne 2006a. 400 µg per day was used in Mosges 2011, Zhou 2015 and in the higher-dose arm of Stjarne 2006. Chur 2013 used either 100 µg a day for children aged 6 to 11 or 200 µg per day for children aged 12 to 17. It also had a group using a higher dose, doubling the doses to 200 µg and 400 µg per day respectively.
- Budesonide nasal spray was used in Johansen 1993, Holopainen 1982 (400 µg per day) and Lund 2004 (128 µg per day).

Outcomes

Health-related quality of life, using disease-specific health-related quality of life scores

One study used the Rhinosinusitis Outcome Measure (RSOM-31) (Hansen 2010). Stjarne 2006a reported that scores for quality of life were recorded at every study visit using an investigator-administered scale with the following items: “nose breathing”, “experience of smell and taste”, “interference with daily activities caused by nasal symptoms” and “sleep disturbance”. It is unlikely that this is a validated quality of life instrument.

Disease severity, as measured by patient-reported symptom score

All studies but one (Lang 1983) measured disease severity as reported by patients, but only Lund 2004 used a validated scale (Chronic Sinusitis Survey - CSS). However, Lund 2004 reported that at the time of the study, a validated version of the Hungarian questionnaire was not available (the study was conducted in 19 centres, six of which were in Hungary). Most studies presented the results (partially) in graphs as median or mean values. The method of measurement was reported in several ways and differed in the choice of scale, type/combination of symptoms measured, timing of measures and scoring/analysis method (see Appendix 3). Due to these variations, it was unclear whether the total scores between studies were comparable with each other and we had to make modifications in order to standardise scores before meta-analysis could be conducted (see Methods and Potential biases in the review process).
Significant adverse effect: epistaxis
All but five studies reported the number of patients with epistaxis and these could be meta-analysed (Holmberg 1997; Johansen 1993; Lang 1983; Lund 1998; Parikh 2001). Holmberg 1997 and Johansen 1993 did not provide enough information about the types of adverse effects experienced and how many patients experienced them, whereas the other three studies were very small (fewer than 15 patients per treatment arm) and it was unclear whether epistaxis was not experienced by any patients or not reported. The studies included nosebleeds of all severities, including mild.

Health-related quality of life, using generic quality of life scores, such as the SF-36, EQ-5D and other well-validated instruments
Lund 2004 used the SF-36, but only reported for which subscale a statistically significant difference was found.

Other local adverse effects: local irritation (including oral thrush, sore throat and other local nasal irritation such as dryness, itchiness etc.)
A variety of types of possible local irritation symptoms were reported in the studies, ranging from “itchiness” to “nasal burns” and “nasopharyngeal pain/discomfort”. Some studies used broader classifications whereas others used more specific description. For example, Vlckova 2009 reported an unusual combination of “adverse events”, listing “rhinalgia” and “nasal discomfort” separately alongside headache; they also reported “rhinitis” as an adverse event alongside “sneezing” and “rhinorrhea”.

Other systemic adverse effects (in children - stunted growth, in adults - osteoporosis)
Only one study followed up patients in the longer term (Lang 1983), and this adverse effect was not reported or relevant in the other studies.

Nasal endoscopic score (either nasal polyp size score or endoscopy score, e.g. Lund-Kennedy)
Lund 1998 and Stjärne 2006a did not report any endoscopy results.
Studies that recruited patients with nasal polyps mostly used a 0- to 3-point scale. In most studies, this was the Lildholdt scale, but the definitions were not reported clearly in the other studies. Whenever studies reported these as mean values at endpoint or change from baseline, they summed the scores from both sides of the nose (0 to 6 range). However, no definitions were given in the studies that reported the results as proportions of people who had improved.
One study used a 0 to 4 scale (Holmberg 1997), after some modification to the Lildholdt scale. The investigators in Aukema 2005 scored the volume of polyps on a 0 to 10 cm visual analogue scale after conducting endoscopy.
Studies that included patients without polyps used the Lund-Kennedy scale (Hansen 2010; Mosges 2011; Parikh 2001). It was unclear whether Lund 2004 used endoscopy scores as an outcome - this was not reported.

Computerised tomography (CT) scan score (e.g. Lund-Mackay)
Only one study reported a CT scan score (Aukema 2005). The Lund-Mackay score was used (0 to 24 points, higher score = more severe).

Excluded studies
We excluded 55 papers (45 studies) after reviewing the full text. Further details of the reasons for exclusion can be found in the Characteristics of excluded studies table. We excluded 23 studies due to the population, most of these (19 studies) because all patients received surgery at the start of the trial and the intranasal steroids were used to try to prevent recurrence of polyps (Bross-Soriano 2004; Cassano 1996; Dijkstra 2004; Drettnar 1982; el Naggar 1995; Gulati 2001; Hartwig 1988; Jorissen 2009; Jurkiewicz 2004; Kang 2008; Karlsson 1982; Malmberg 1988; Passali 2003; Rotenberg 2011; Rowe-Jones 2005; Slifirski 2009; Stjärne 2009; Vento 2012; Virolainen 1980). Other reasons for excluding studies based on the population were related to patients not meeting the current criteria for chronic rhinosinusitis (ALA 2015), chronic allergic or bacterial sinusitis where less than 50% of patients had chronic disease (Cuenant 1986), aspirin-induced asthma and chronic eosinophilic rhinitis (Mastalerz 1997), and recurrent or chronic maxillary sinusitis (Qvarnberg 1992). We excluded three studies because the studies were designed to look at the impact of intranasal corticosteroids on outcomes when given perioperatively; all patients underwent surgery during the period of the trial (Albu 2010; Ehnhage 2009; Saunders 1999). Two studies compared steroids with placebo or no treatment but we excluded them as they used a catheter or tube to administer steroids directly to the participant’s sinuses (Furukido 2005; Lavigne 2002). One study was a clinical trial that appeared to meet the inclusion criteria but the clinical trials registry website stated that the trial had been terminated early (Optinose 2012). The reason for early termination is not provided. We excluded the remaining 16 papers as they did not meet the minimum requirements for the duration of treatment and follow-up. Ten of these studies had a follow-up time of one month or less (Chalton 1985; Johansson 2002; Kapucu 2012; Keith 1995; Lildholdt 1995; Mygind 1975; Ruhno 1990; Taub 1968; Toft 1982; Wang 2015), whereas the remaining six treated and followed up patients for between six and eight weeks (Filiaci 2000; Jankowski 2001; Jankowski 2009; Man 2013; Meltzer 1993; Tös 1998).
**Ongoing studies**

We identified three relevant ongoing studies, all of which are in adults with chronic rhinosinusitis with nasal polyps (NCT01622569; NCT01624662; NCT01013701). Two of these are large, multicentre trials each with a planned population of over 300 patients (NCT01622569; NCT01624662). These two trials will make the same comparisons, comparing three different doses of fluticasone propionate (400 µg bid, 200 µg bid and 100 µg bid) with placebo. All of the arms will use a novel bi-directional device. The studies were completed in October 2015 but no study data were available at the time of writing this review. The other trial compares fluticasone furoate with placebo for 16 weeks (NCT01013701). The trial information was registered in 2009 and the ClinicalTrials.gov website reports that the recruitment status of the trial is unknown as the information has not been recently validated. We attempted to contact the investigators but we did not receive a response.

**Risk of bias in included studies**

We included 18 studies in this review. Nine of these had low risk of bias for both selection and blinding (Keith 2000; Lund 2004; Mosges 2011; Parikh 2001; Penttila 2000; Small 2005; Stjarne 2006; Stjarne 2006a; Zhou 2015). Lang 1983 was only available as an abstract and therefore there was insufficient information to judge the risk of bias for most domains. We did most of the ratings based solely on the study report(s), as the trials were not registered and no protocols were available.

See Figure 2 for the 'Risk of bias' graph (our judgements about each risk of bias item presented as percentages across all included studies) and Figure 3 for the 'Risk of bias' summary (our judgements about each risk of bias item for each included study).
Figure 3. 'Risk of bias' summary: review authors' judgements about each risk of bias item for each included study.
Allocation

Sequence generation
Six of the studies only stated that the trials were randomised but did not provide further information about how sequence generation was conducted. We therefore judged them to be at an unclear risk of bias (Hansen 2010; Holmberg 1997; Holopainen 1982; Johansen 1993; Lang 1983; Vlckova 2009).

Another two studies also did not provide details of their randomisation procedures, but we judged this to be a low risk of bias because these studies were conducted quite recently as multinational trials and therefore should have used adequate methodology to ensure adequate sequence generation (Penttilla 2000; Small 2005). The rest of the studies either specified that the randomisation was conducted by another unit supporting clinical trials (the pharmacy for Parikh 2001) or provided a clear description of how computerised sequence generation was used.

Allocation concealment
We also rated nine studies as low risk of bias for allocation concealment (Keith 2000; Lund 2004; Mosges 2011; Parikh 2001; Penttilla 2000; Small 2005; Stjarne 2006; Stjarne 2006a; Zhou 2015). As for sequence generation, we considered the large multinational RCTs to have adequate methodology although they did not provide specific information about the allocation concealment method (Penttilla 2000; Small 2005; Stjarne 2006a). An exception to this is Chur 2013, which we rated as unclear risk of bias for allocation concealment. Despite the adequate methods used to generate a random sequence, blocked randomisation was used and the effectiveness of the blinding was unclear in the absence of a 'double-dummy' design in this multi-arm trial.

We rated the other studies as unclear risk of bias due to lack of information.

Blinding
The ratings for the risk of performance bias versus detection bias were closely related for this review. Most of the outcomes were assessed by the patients and the overall risks of bias were low when both participants and investigators were adequately blinded. We did not find information to suggest that the clinicians could have obtained extra information from blood tests etc. to allow them to 'guess' which treatment the patients were allocated. We considered the majority (16) of the studies to be at low risk of bias for blinding. We considered two studies to be at unclear risk: Chur 2013 did not use a double-dummy design to mask the regimens and there was no information in Lang 1983.

Incomplete outcome data
We considered only four trials to have a low risk of attrition bias: Holopainen 1982; Keith 2000; Vlckova 2009 and Zhou 2015. Eleven studies had a high risk of attrition bias due to unbalanced drop-out rates between the placebo and treatment groups, with higher drop-out rates in the placebo groups (Aukema 2005; Hansen 2010; Penttilla 2000, Small 2005; Stjarne 2006; Stjarne 2006a), an overall high rate of drop-outs (Lund 2004), or a combination of these factors (Holmberg 1997; Lund 1998; Parikh 2001). In Mosges 2011, only 10% of patients did not complete the full study, but it was unclear why the study only included 75% of the sample in intention-to-treat (ITT) analysis due to “major protocol violation” and 61% in the per protocol population due to “minor protocol violation”.

The risk of attrition bias was unclear for three of the included studies (Chur 2013; Johansen 1993; Lang 1983). These studies did not provide enough information to adequately judge the risk. For example, Johansen 1993 reported that 5/91 (5.5%) participants did not complete the study. There was no information on how many were randomised to each group in Johansen 1993 and so it is difficult to determine whether this could have affected the results.

Selective reporting
Many of the study reports presented effectiveness outcomes only in graphs and only provided limited, selective information, for example P values or mean values only when statistical significance was noted.

We considered only two studies to have low risk of bias, with all expected outcomes reported (Small 2005; Zhou 2015).

We considered the risk of selective reporting to be high in eight studies (Aukema 2005; Chur 2013; Holmberg 1997; Holopainen 1982; Lang 1983; Lund 1998; Mosges 2011; Parikh 2001). These studies either presented outcomes that were not pre-specified in the methods section or downplayed and provided insufficient information about prespecified outcomes. Others used an unclear or arbitrary method to combine data or report some of the outcomes (or both).

The primary endpoint in Chur 2013 was "safety" (cortisol levels) and despite presenting the mean change values for effectiveness outcomes, they did not provide any P values or standard deviations. The study's rationale for collecting the data and not fully reporting them was: "No statistical analysis of efficacy end points was pre-specified in the study protocol, and only descriptive efficacy statistics were collected." We observed that these values (mean changes) were similar between groups and unlikely to be statistically significant and so poor reporting due to lack of beneficial effects cannot be ruled out.
We considered the remaining eight studies to be at unclear risk. There was not enough information in the methods and/or protocol and we found it difficult to judge whether there was a risk of reporting bias.

Other potential sources of bias

Use of validated outcome measures
The lack of use of validated outcome measures is a major bias concern in this review. If an instrument is insensitive to measuring differences, this biases towards a finding of ‘no difference’ in the studies and also in this review.

None of the included studies mentioned using validated outcome measures, for either of the primary effectiveness outcomes (disease-specific health-related quality of life and disease severity/symptom scores).

Almost all studies attempted to measure change in symptom scores as measured by patients, but none reported validation of the instruments being used. Most studies used a 0 to 3 scale in the “diaries”, but used different methods to calculate the result (period of time, combination of scores). There is no evidence that a 0- to 3-point scale, especially when used as a single scale, has the sensitivity to distinguish between groups of patients who improved versus those who did not (discriminant validity) or whether the different method of scoring was valid.

The scales used to measure nasal polyps were generally well described. However, it is again unclear whether a 0- to 3-point scale has the discriminant validity to detect a difference in the small trials seen.

Effects of interventions

See: Summary of findings for the main comparison Intranasal corticosteroids for people with chronic rhinosinusitis

Where the range of scales and values for minimal important differences were unclear, we used the standardised mean difference (SMD) as a guide to estimate the effect sizes. As suggested in the Cochrane Handbook for Systematic Reviews of Interventions (Handbook 2011), we used standard rules of thumb in the interpretation of effect sizes (SMD, or Cohen’s effect size of < 0.41 = small, 0.40 to 0.70 = moderate, > 0.70 = large) (Cohen 1988). Established scales such as the SF-12 may have other rules of thumb to estimate the minimal important difference (MID = 0.5 SMD) and we use these to guide our interpretation whenever available (Jaeschke 1989; Revicki 2008).

Although we had planned to present data for patients with our without polyps in subgroups as a visual comparison, this was not necessary for the effectiveness outcomes because no more than one study contributed to the analysis, with the exception of polyps score data.

Intranasal corticosteroids compared to placebo or no intervention

Three of the 18 studies did not contribute any data for meta-analysis (Holmberg 1997; Johansen 1993; Parikh 2001). In our protocol, we had specified that results from studies of participants with chronic rhinosinusitis with nasal polyps and without nasal polyps would be presented as subgroups in the forest plots (Chong 2015). However, this was only possible for the outcomes of epistaxis and local irritation. Of the four studies in patients with chronic rhinosinusitis without nasal polyps (Hansen 2010; Lund 2004; Mosges 2011; Parikh 2001), only Lund 2004 contributed data for the effectiveness outcomes (disease severity).

Health-related quality of life, using disease-specific health-related quality of life scores

Hansen 2010 (chronic rhinosinusitis without nasal polyps) was the only study reporting the used of a disease-specific health-related quality of life questionnaire, the Rhinosinusitis Outcome Measures-31 (RSOM-31). The median change from baseline (median 178 points) was -62 points for the intranasal corticosteroids group (n = 10) and -5 points for the placebo group (n = 10, from a median baseline score of 187 points). The difference was not statistically significant (Mann-Whitney U test).

Disease severity, as measured by patient-reported symptom score (such as the Chronic Sinusitis Survey (CSS) questionnaire and visual analogue scales)

Chronic Sinusitis Survey

Lund 2004 (chronic rhinosinusitis without nasal polyps) reported that at the time of the study, a validated version of the Hungarian questionnaire was not available (the study was conducted in 19 centres, six of which were in Hungary). Only the mean change and 95% confidence interval (CI) for each group was reported, without a sample size. We imputed the sample size based on the number of patients available at the end of the study and it seems that all available patients had filled in the CSS. The mean difference (MD) was 2.84 points (95% CI -5.02 to 10.70; 134 participants) between groups (range 0 to 100, higher values = better) (Analysis 1.1). The magnitude of the difference is insignificant.

Global rating scale

Vlckova 2009 used a five-point global rating scale (very much improved, improved, same, worse or very much worse). At 12 weeks, 76% of patients in the treatment group were “improved” or “very much improved” compared to 27% of patients in the placebo group (risk ratio (RR) 2.78, 95% CI 1.76 to 4.40; 109 participants; one study) (Analysis 1.2).
Combined symptom scores
Six studies provided information on mean and standard deviation values, or had enough information in their graphs for us to estimate these values for various symptom scores (Aukema 2005; Lund 2004; Small 2005; Sjarne 2006; Vlckova 2009; Zhou 2015). Of these, only Vlckova 2009 and Lund 2004 (chronic rhinosinusitis without nasal polyps) reported an overall combined score for all four groups of symptoms covered by EPOS 2012 (nasal blockage, nasal discomfort (facial and sinus pain/pressure) and rhinitis symptoms (nasal secretion, itching, irritation and sneezing), and loss of sense of smell). Parikh 2001 also included these symptoms in their diary card and symptom scores, but it is less clear how these were calculated. Although the means and standard deviations were reported for this study, we did not include it in the meta-analysis because the values were obviously skewed (mean of 29.4 (standard deviation (SD) 37) for intranasal corticosteroids group (n = 9), mean of 16.9 (SD 48.5) (n = 13) for placebo group, P value = 0.39 using Mann Whitney U test).

Therefore we only included six studies in our analysis. We report the average values when all four types of symptoms mentioned in EPOS were measured versus when only three types (loss of sense of smell, nasal blockage and nasal discharge) and two types of symptoms (nasal blockage and nasal discharge) were measured. All studies used 0- to 3-point scales in their diaries, except for Aukema 2005, which used a 0 to 100 visual analogue scale (VAS) measured during follow-up. All studies reported change from baseline, except for Aukema 2005, which reported the mean difference at the end of the study. To allow for ease of interpretation, we converted these 0 to 100 VAS scores into 0 to 3 by a division of 33.333. All studies used the ‘usual dose’ of intranasal steroids, except for Vlckova 2009, which only used a higher dose of fluticasone propionate (800 µg/day), delivered in two divided doses. Two studies, Small 2005 and Sjarne 2006, had two treatment arms using mometasone furoate nasal spray with low (200 µg/day) and high (400 µg/day) doses. Only one study included patients without nasal polyps (Lund 2004). All studies were conducted in adults. The pooled results are as follows:

- Combined symptom score for four EPOS domains, average score: MD -0.26 (95% CI -0.37 to -0.15; 243 participants; two studies; I² = 46%), scale range: 0 to 3, lower = better, indicating less severe symptoms in the intranasal corticosteroids group (Analysis 1.3).
- Combined symptom score for three EPOS domains (nasal blockage, rhinorrhoea and loss of sense of smell), average score: MD -0.31 (95% CI -0.38 to -0.23; 1345 participants; four studies; I² = 0%), scale range: 0 to 3, lower = better (Analysis 1.3).
- Combined symptom score for two EPOS domains (only nasal blockage and rhinorrhoea), average score: MD -0.31 (95% CI -0.38 to -0.24; 1702 participants; six studies; I² = 0%), scale range: 0 to 3, lower = better (Analysis 1.3).

The observed mean differences correspond to a moderate effect size (the SMD was about 0.4 for the two and three domain average symptom scores and 0.6 for the four domain average symptom score). The quality of the evidence is low for the four domain scores and moderate for the three and two domains scores (facial pain/pressure not considered), with the main concerns being the use of non-validated symptom scores and a high risk of reporting bias (many studies did not publish the results in detail and this could be linked to a lack of observed efficacy).

Individual symptom scores
- Nasal blockage: MD -0.40 (95% CI -0.52 to -0.29; 1702 participants; six studies; I² = 47%) (Analysis 1.4).
- Rhinorrhoea: MD -0.25 (95% CI -0.33 to -0.17; 1702 participants; six studies; I² = 6%) (Analysis 1.4).
- Loss of sense of smell: MD -0.19 (95% CI -0.28 to -0.11; 1345 participants; four studies; I² = 0%) (Analysis 1.4).
- Facial pain/pain/pressure: MD -0.27 (95% CI -0.56 to 0.02; 243 participants; two studies; I² = 78%). Of these two studies, Lund 2004 included patients with chronic rhinosinusitis without nasal polyps, whereas Vlckova 2009 included patients with chronic rhinosinusitis with nasal polyps. Due to the differences in type, dose and delivery method (fluticasone propionate 800 µg per day using breadth actuated inhaler versus budesonide 128 µg per day as a nasal spray), the source of heterogeneity was unclear.

We used a random-effects model to conduct the analysis; if a fixed-effect model is used the statistical significance of the pooled MD for facial pain is -0.24 (95% CI -0.37 to -0.11). The quality of the evidence is moderate for nasal blockage, rhinorrhoea and loss of sense of smell, but low for facial pain/pressure.

Significant adverse effect: epistaxis
The risk of epistaxis was higher in the intranasal corticosteroids group (RR 2.74, 95% CI 1.88 to 4.00; 2508 participants; 13 studies; I² = 0%) (Analysis 1.5). The quality of the evidence is high.

Health-related quality of life, using generic quality of life scores, such as the SF-36, EQ-5D and other well-validated instruments
Lund 2004 used the SF-36, but only stated that there was a statistically significant improvement in the general health subscale in patients on intranasal corticosteroids compared to placebo. Apart from stating that no other significant differences were observed, no other details were reported.

Other local adverse effects: local irritation (including oral thrush, sore throat and other local nasal irritation such as dryness, itchiness etc.)
It is unclear whether there is an important difference in the risk of local irritation between participants taking intranasal corticosteroids or placebo (RR 0.94, 95% CI 0.53 to 1.64; 2124 participants; 11 studies; I² = 0%) (Analysis 1.6). The quality of the evidence is low (we are uncertain about this estimate), because the reporting of local irritation effects varied a lot between studies. This was sometimes finely split into many types of local irritation and we could only use the numbers for the most commonly reported types of irritation to avoid double counting in this review. The actual event rate for all types of local irritation is higher than reported in this analysis.

**Other systemic adverse effects (in children - stunted growth, in adults - osteoporosis)**

None of the studies treated or followed up patients for long enough to report these adverse events.

**Nasal endoscopic score (depending on population, either nasal polyps size score or endoscopy score, e.g. Lund-Mackay/Lund-Kennedy)**

Three studies reported polyps score results as the mean change from baseline on a 0- to 6-point scale. Five studies reported this as the proportion of patients who had an improvement. Vlckova 2009 reported both the mean improvement and the proportion of patients with an improvement. Aukema 2005 measured polyps size on a 0 to 10 cm visual analogue scale and seems to have reported the values at the end of the study. We did not include this in the analysis as it was unclear what the scale was and whether it was valid. The MD was -24.70 (95% CI -48.00 to -1.40, n = 47) and we observed heterogeneity when it was combined with the other studies.

**Chronic rhinosinusitis with nasal polyps - reduction in polyps size**

The MD in polyps score was -0.58 (95% CI -0.90 to -0.26; 1417 participants; four studies; I² = 83%) indicating less severity for the intranasal corticosteroids group (Analysis 1.7). All reported the sum of polyps score from both sides of the nose (range 0 to 6). One study, Vlckova 2009, had an effect that was larger than the other studies, with a mean difference of -1.21 (95% CI -1.56 to -0.86) points between treatment arms in the reduction of polyps size score. When this study is removed, the heterogeneity is resolved and the observed effect size is smaller (MD -0.35, 95% CI -0.47 to -0.24; 1308 participants; three studies; I² = 0%).

Five studies reported the proportion of participants who had an improvement in their polyps scores. The chance of an improvement was higher in patients on intranasal corticosteroids (RR 1.77, 95% CI 1.06 to 2.95; 676 participants; five studies; I² = 66%) (Analysis 1.8). The observed heterogeneity was resolved when we removed a study with an outlier effect (Vlckova 2009), but the RR became slightly smaller (RR 1.46, 95% CI 1.12 to 1.90; 567 participants; four studies; I² = 0%).

Using a method recommended in the *Cochrane Handbook for Systematic Reviews of Interventions*, we converted the continuous outcome data into proportions and analysed them together using the generic inverse variance method. For Vlckova 2009, which reported both the mean difference and proportions, we used the values reported for proportions to avoid double counting and to minimise imputations. The overall pooled odds ratio (OR) was 2.07 (95% CI 1.48 to 2.91; 1984 participants; eight studies) (Analysis 1.9). The observed heterogeneity was quite substantial (> 50%). We observed that Vlckova 2009 seemed to have a larger effect than the other studies. If this study is removed from the analysis, the OR is slightly lower (OR 1.71, 95% CI 1.43 to 2.04; 1875 participants; seven studies) and heterogeneity is no longer observed.

**Chronic rhinosinusitis without nasal polyps - endoscopy score**

The studies that included patients without polyps, such as Hansen 2010, Mosges 2011 and Parikh 2001, used a modified Lund-Kennedy score. However, the results were either only partially reported or could not be meta-analysed due to highly skewed distribution (very small sample sizes). Lund 2004 did not report the endoscopy score as an outcome and it was unclear whether this was measured.

Parikh 2001 reported the percentage of change and standard deviation. This was -22.3% (SD 61.8) (n = 9) for the intranasal corticosteroids group and +19.9% (SD 58.3) (n = 13). This was not statistically significant (Mann Whitney U test). Mosges 2011 showed the total endoscopy score for redness, oedema and discharge on a chart. There were no statistically significant differences between groups. Similarly, Hansen 2010 reported no statistically significant differences between groups for different aspects of the assessment, except for oedema.

**Computerised tomography (CT) scan score (e.g. Lund-Mackay)**

Only one study reported the CT scan score (Aukema 2005). The MD was -4.82 (95% CI -7.27 to -2.37; 47 participants; one study; I² = 0%) (Analysis 1.10). The Lund-Mackay score was used (0 to 24 points, higher score = more severe).

**DISCUSSION**

**Summary of main results**
This review includes a total of 18 studies. However, many studies were very small and did not measure and/or report data in a way that allowed for meta-analysis.

Only one very small study (n = 20) reported disease-specific health-related quality of life, using the Rhinosinusitis Outcome Measures-31 (RSOM-31). Lund 2004 described using the SF-36, but reported that only the general health subscale showed a statistically significant difference between groups.

We found that intranasal corticosteroids improved patient symptom scores, when these were measured as a combined score, with scores for individual items or as a global score (non-validated scales). Apart from rhinorrhoea and loss of sense of smell, which had small effect sizes when measured as individual items, the effect sizes observed using other parameters corresponded to moderate effect sizes. However, because these data could only be obtained from a few studies (many studies did not report enough detail when the results were not ‘significant’) there is a risk of reporting bias.

Epistaxis was probably the most consistently reported outcome, with 13 out of 18 studies reporting this. The risk was increased in patients using intranasal corticosteroids (risk ratio (RR) 2.74, 95% confidence interval (CI) 1.88 to 4.00; 2508 participants; 13 studies). However, local irritation was very inconsistently reported, with studies using various definitions to report the data. The RR obtained was 0.94 (95% CI 0.53 to 1.64; 2124 participants; 11 studies; I² = 0%) and it is unclear whether there is an important increase in risk. None of the studies treated or followed up patients for long enough to provide meaningful data on osteoporosis risk or risk of stunted growth.

There seemed to be an increased odds of polyp improvement in people who had chronic rhinosinusitis with nasal polyps. However, the results for endoscopy score were not well reported in studies that included participants with chronic rhinosinusitis without polyps.

**Overall completeness and applicability of evidence**

A good range of types and doses of intranasal corticosteroids are included in this review. However, the main body of evidence is in patients with chronic rhinosinusitis with polyps. Studies in chronic rhinosinusitis without polyps tended to be smaller, had poorer reporting and did not contribute enough information to the meta-analyses to allow us to evaluate whether there are differences in effectiveness between these subgroups. Only one of these studies was conducted in children (up to 18 years old) and therefore it is again unclear whether the evidence is applicable to children. Most studies were conducted for three to four months, with the exception of three studies that were conducted for 20 weeks (Aukema 2005), 26 weeks (Lund 2004), and two years (Lang 1983), respectively. Therefore, it is unclear whether the observed effectiveness is maintained if intranasal corticosteroids are used over longer periods and whether these benefits can be maintained after patients stop treatment.

Most of the studies did not make any reference to whether saline irrigation could be used. However, Vlckova 2009 specified that patients could not use nasal saline. This study showed larger effect sizes than the other studies and unlike most studies, where participants in both the intervention and placebo groups showed some improvement over time, the participants in the placebo group in this study got worse with time. It is unclear whether this is a random observation or due to the patient population, the high doses used or the exclusion of normal saline.

**Quality of the evidence**

The studies included in this review were relatively well conducted, with most using good methodology for selection of patients and blinding.

However, we had serious concerns about how the effectiveness outcomes were measured, analysed and reported. The validity of the tools used to measure outcomes is a major concern. Most studies did not use validated tools to measure and score symptom severity scales and this reduces our confidence in the estimates of effect. Moreover, there is also a possibility of reporting bias, as studies only tended to report enough detail to allow for meta-analysis when they found a statistical significant result. These reasons significantly reduced our confidence in the estimates of effect sizes and we rated the quality of the evidence as moderate for disease severity outcomes. The exception was those outcomes involving facial pain/pressure, but there was some unexplained heterogeneity and there were fewer data. For quality of life, the quality of the evidence is very low.

The reporting methodology for ‘local irritation’ was also a concern and along with the imprecision observed we considered the evidence for this outcome to be of low quality.

However, we considered the evidence for the risk of epistaxis to be of high quality; this is an outcome that appears to have been collected and reported consistently across studies.

**Potential biases in the review process**

There were two major challenges to meta-analysis in the review of effectiveness outcomes: 1) the lack of use of validated tools and variations in how outcomes were measured and reported and 2) the outcomes were often not fully reported.

**Lack of use of validated tools and variations in how outcomes were measured and reported**

The lack of use of validated instruments to measure patient-important outcomes, such as the impact on quality of life and disease severity, is probably the single most important issue that hampers
our ability to meta-analyse results or to compare results between studies. Studies used a variety of scales, timings and analysis methods to measure different combinations of symptoms. Some studies may have measured one group of symptoms (e.g. nasal discharge) as several separate items (anterior rhinorrhoea, post-nasal drip), but not measured other types of symptoms at all (e.g. loss of sense of smell and facial pain/pressure). In the absence of evidence for both disease-specific health-related quality of life and disease severity measured with validated tools (only one study respectively, Hansen 2010 and Lund 2004, reported these), some ‘standardisation’ of these measures had to be conducted in order to allow the results to be pooled. We took the decision to combine the scores for individual symptoms to create a total symptoms score. The methods we used to do this are described in the methods section (Dealing with missing data). The symptoms we included were based on the EPOS 2012 diagnostic criteria, but this score was not a validated measure and the correlation between symptoms was not accounted for in the results.

In addition to making an assumption of no correlation between symptoms, we also had to make other decisions to standardise the data:

- Most studies only specified that diaries were completed in the morning, whereas others were completed in the morning and evening and may, or may not, have reported the results separately. Where studies presented both morning and evening scores, we only used the morning score values in this review, to allow for standardisation across studies. We avoided taking an average of morning and evening data and estimating the standard deviations, since these should be taken as paired data and this may further overestimate the size of the standard deviation (see below for considerations when imputing standard deviations).
- Some studies only measured the outcomes at the endpoint. We had to assume that the difference between changes and endpoints would produce a similar mean difference between groups and therefore could be pooled. We made this assumption for Aukema 2005.
- Aukema 2005 used a 10 cm visual analogue scale. Since this was only one small study, we had to assume that the scales were linear and could be converted to a 0- to 3-point scale; we tested this in a sensitivity analysis. Although the effect size observed for this study was larger, it did not have an impact on the results.

Fortunately, most of the studies had measured symptom severity using diaries with a 0- to 3-point scale and we could use the mean differences to look at the size of the effect. However, what is a minimal important difference (MID) for a 0- to 3-point scale is not known and to assist interpretation we still had to look at the standardised mean difference (SMD) and used Cohen’s effect size as a rule of thumb. Bearing in mind that we did factor in correlations (which will result in smaller standard deviations), this means that the effect sizes could be larger than estimated. Nevertheless, we still found moderate to large effect sizes in disease severity scores. In fact, a moderate effect size was found in Lund 2004 using the individual or combined scores, even though the study did not find a clinically important difference using the Chronic Sinusitis Survey (CSS), a validated tool.

Local irritation is another outcome where there were many variations in reporting, in terms of categorisation and descriptions used. Where studies reported more than one type of local irritation (e.g. nasal burning and nasal irritation were both reported), we took the data for the outcome with the higher total event rate. If rates were the same for both outcomes, we chose the one terminology that was closest to the description of general irritation (e.g. nasal irritation would be used in this example), with the review author blinded to the data.

Outcomes were often not fully reported

Many of the data for the included studies were presented in the papers in graphs or charts. Where this was the case, we extracted the data from the paper using an online program (http://arohatgi.info/WebPlotDigitizer/app/). There will inevitably be a degree of error in using these data, both from inaccuracies during the printing process and the process used to collect the data. Where P values were reported as less than a certain threshold (e.g. P value < 0.05), calculations were based on P value = 0.05. This is a conservative estimate for standard deviation values and we were conscious of the need to minimise imputations as much as possible. For example, Small 2005 had two arms (a high-dose and a low-dose group) and only reported the P values compared to placebo for many outcomes. Rather than trying to estimate what these P values may be when ‘combined’ and risk inflating the standard deviation further, we entered this study twice as a high-dose and low-dose group, and halved the sample size in the placebo group to avoid double-counting.

Agreements and disagreements with other studies or reviews

This review aimed to answer the clinical question of whether intranasal corticosteroids are effective in patients who have chronic rhinosinusitis. It is one of a series of reviews looking at the (relative) effectiveness and safety of different medical interventions for chronic rhinosinusitis. Although the intranasal steroids included in this review comprised different types of molecules, doses, regimens and delivery methods, we observed no statistical heterogeneity that suggested that these factors could result in different levels of effectiveness and adverse events such as epistaxis and local irritation. This observation about the impact of variations in the types and doses of intranasal corticosteroids is consistent with our companion review, which looks at different types and doses of intranasal corticosteroids (Chong 2016a). Chong 2016a also did not find any important differences between high and low doses and there was very little evidence to draw conclusions about the other aspects. However, Chong 2016a
did find it possible that the risk of adverse effects is increased when higher doses are used, whereas in this review we did not observe any heterogeneity.

This review only compared intranasal steroids against placebo. We have compared short-course oral steroids against intranasal corticosteroids in Head 2016a and Head 2016b; antibiotics against intranasal corticosteroids in Head 2016c; and nasal saline against intranasal corticosteroids in Chong 2016b. When compared to oral corticosteroids, patients who received oral corticosteroids instead of intranasal corticosteroids seemed to benefit more in terms of reduced disease severity and polyps size for short follow-up periods (two to three weeks), but it was uncertain whether the benefit persisted (difference minimises by three months) (Head 2016a). The antibiotics review only found one very small study comparing antibiotics (12 weeks of 250 mg clarithromycin) against 200 µg of mometasone furoate spray and did not find an important difference between groups for overall disease severity, but found the endoscopy score to be slightly better in the antibiotics group (Head 2016c). The saline review only found one very small study comparing intranasal corticosteroids versus nebulised saline; intranasal corticosteroids were much more effective than nebulised saline (Chong 2016b). However, most of the evidence for intranasal corticosteroids compared against other interventions is of very low quality (we have very little confidence in the effects estimated - the evidence is inconclusive).

Unlike previous reviews of intranasal steroids, which focused on either patients with polyps (Kalish 2012) or without polyps (Snidvongs 2011), this review includes all types of chronic rhinosinusitis patients. However, we limited inclusion to studies that had followed up patients for at least three months and we excluded patients who had just undergone surgery. The impact of intranasal steroids in reducing recurrence in patients who have just had sinus surgery is not the clinical question addressed by this review.

**AUTHORS’ CONCLUSIONS**

**Implications for practice**

Patients with chronic rhinosinusitis with nasal polyps showed an improvement in symptoms on intranasal corticosteroids. However, data are lacking for patients without polyps and it is unclear whether they derive a similar level of benefit.

The risk of adverse effects such as epistaxis and local irritation is increased in people taking intranasal corticosteroids, although the severity of the epistaxis is unknown, as is whether patients discontinue usage as a result. If epistaxis is limited to streaks of blood in the mucus it may be tolerated by the patient and be safe to continue treatment. However, it may be a factor that affects compliance. Ensuring good technique in usage of spray by patients may help to reduce these effects, especially where spraying of the septum is avoided. Different intranasal corticosteroid delivery nozzles may also have a bearing on this.

**Implications for research**

As of August 2015, most studies of intranasal steroids have been conducted in participants with chronic rhinosinusitis with nasal polyps. The evidence we found suggests that chronic rhinosinusitis patients show an improvement in symptoms with intranasal corticosteroids. Recent international trials using the Optinose device (Navigate trials I and II) have been completed and they include different doses within their protocols and a comparison with placebo. Further information will therefore be forthcoming once these results are published (NCT01622569; NCT01624662).

Future research should recruit patients with chronic rhinosinusitis diagnosed using the EPOS 2012 criteria and include both patients with and without nasal polyps (stratified randomisation by subgroup). Intranasal corticosteroids should be compared against placebo and this should be considered against a background of nasal irrigation, including in the placebo arm. The intervention and follow-up should be carried out for at least three or six months, since intranasal corticosteroids are used as a long-term treatment for a chronic condition.

A key area of weakness across all of the included studies was the absence of both disease-specific and generic health-related quality of life tools as outcome measures. It is recommended that any future research uses primary outcome measures that are relevant to patients and any disease-specific instruments used should be validated in people with chronic rhinosinusitis. Many studies chose to use polyp scores as their primary outcome measure yet the correlation between endoscopic results and patient symptoms is unclear. The methods for defining and recording adverse events should be considered at the protocol stage and adverse events recorded should include epistaxis and local irritation; longer-term effects such as osteoporosis should also be considered.

This review is one of a suite of reviews of medical treatments for chronic rhinosinusitis, each of which features its own research recommendations. Across all reviews, key features of future research are as follows:

- Trials should be adequately powered and imbalances in prognostic factors (for example, prior sinus surgery) must be accounted for in the statistical analysis.

- Study participants should be diagnosed with chronic rhinosinusitis using the EPOS 2012 criteria and should primarily be recruited based on their symptoms. Different patient phenotypes (that is, those with and without nasal polyps) should be recognised and trials should use stratified randomisation within these subgroups or focus on one or other of the phenotypes.
• Studies should focus on outcomes that are important to patients and use validated instruments to measure these. Validated chronic rhinosinusitis-specific health-related quality of life questionnaires exist, for example the Sino-Nasal Outcome Test-22 (SNOT-22). Patients may find dichotomised outcomes easiest to interpret; for example the percentage of patients achieving a minimal clinically important difference (MCID) or improvement for that outcome. Such MCIDs or cut-off points should be included in the study protocol and clearly outlined in the methods section.

• Trials and other high-quality studies should use consistent outcomes and adhere to reporting guidelines, such as CONSORT, so that results can be compared across future trials. The development of a standardised set of outcomes, or core outcome set, for chronic rhinosinusitis, agreed by researchers, will facilitate this process.

ACKNOWLEDGEMENTS

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Intranasal steroids versus placebo or no intervention for chronic rhinosinusitis (Review)

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## Characteristics of included studies [ordered by study ID]

### Aukema 2005

<table>
<thead>
<tr>
<th>Methods</th>
<th>Double-blinded, parallel-group RCT, with 12 weeks duration of treatment and follow-up</th>
</tr>
</thead>
</table>
| Participants                  | **Location**: Netherlands, single side  
|                               | **Setting of recruitment and treatment**: patients awaiting FESS at a university hospital |
|                               | **Sample size**:  
|                               | • Number randomised: 27 in intervention, 27 in comparison  
|                               | • Number completed: 26 in intervention, 21 in comparison  
|                               | **Participant (baseline) characteristics**:  
|                               | • Age: median 44 (range 18 to 68)  
|                               | • Gender: 28 males, 26 females  
|                               | • Main diagnosis: polyposis and/or CRS requiring FESS  
|                               | • Polyps status: CT score $\geq 3$ on at least one side on Lund-Mackay  
|                               | • Previous sinus surgery status: 78% in treatment, 88% in placebo  
|                               | • Other important effect modifiers, if applicable:  
|                               | ◦ Atopy (increased specific serum IgE): 19% in treatment and 26% in placebo group  
|                               | **Inclusion criteria**: Patient on waiting list for FESS (recent CT score of $\geq 3$ on one side, prior treatment with corticosteroid spray $\geq 3$ months, total VAS score of $\geq 200$ mm on 6 scales (range of 0 to 100 mm)) 
|                               | “...investigator had observed and approved the administration method (of the single dose nasal drops)” during the run-in period |
| Interventions                 | **Intervention (n = 27)**: Nasules (fluticasone propionate single dose nasal drops, administered once daily at night)  
|                               | **Comparator group (n = 27)**: placebo  
|                               | Patients were instructed to lie on their back with their head hanging down in a vertical position on the edge of a bed while administering the drops  
|                               | **Use of additional interventions (common to both treatment arms)**: not described |
| Outcomes                      | **Outcomes of interest in the review**:  
|                               | Primary outcomes:  
|                               | 1. Disease severity as measured by VAS of 0 to 100 mm for 6 symptoms (nasal blockage, rhinorrhea, facial pain, mucus in throat, loss of sense of smell and headache)  
|                               | 2. Significant adverse effect: epistaxis  
|                               | Secondary outcomes:  
|                               | 3. Other adverse effects: local irritation  
|                               | 4. Endoscopy score (polyp volume) using a VAS  
|                               | 5. CT scan score measured using Lund-Mackay score (max 24 points)  
|                               | **Other outcomes reported by the study**:  
|                               | • Number of patients who “finally needed FESS” (after a minimum follow-up of 6 months, median of 20 months); but actual length in both arms not reported  
|                               | • PNIF |
Notes

Study had a 2-week run-in period; patient’s method of application was assessed before randomisation

<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>
| Random sequence generation (selection bias)     | Low risk           | Quote: “Medications were numbered by means of computerized randomization and were assigned in numeric order.”
Comment: likely to be adequately generated |
| Allocation concealment (selection bias)         | Unclear risk       | Quote: “All investigations were carried out by one investigator”
Comment: no specific description provided. Sequence generation adequate; likely to depend on whether medications and packaging looked identical. Baseline risks potentially different |
| Blinding of participants and personnel          | Low risk           | Quote: “All investigations were carried out by one investigator...Double-blind randomization to FPNDs or placebo took place after the investigator had observed and approved the administration method ... Randomisation codes were not disclosed until a year after all the patients had finished the study”
Comment: likely to be adequate |
| Blinding of outcome assessment (detection bias) | Low risk           | Quote: “All investigations were carried out by one investigator......Randomisation codes were not disclosed until a year after all the patients had finished the study”
Comment: likely to be adequate |
| Incomplete outcome data (attrition bias)        | High risk          | Quote: “At the end of the study, 6 patients of the placebo group and 1 of the FPND group had dropped out. Of the placebo group, 5 patients dropped out because they had a lot of complaints and did not want to complete the study. Three were prematurely scheduled for FESS (2 after 6 weeks and 1 after 2 weeks of treatment). Two patients resumed intranasal corticosteroids (after 4 and 8 weeks of treatment). One patient received oral steroids from her pul- |
monologist because she did not tolerate inhaled steroids (after 8 weeks of treatment) and was scheduled for FESS as well. The one dropout in the FPND group failed to return for the last visit, despite repeated requests. At the second-to-last visit, he was not much improved compared with inclusion, and he finally underwent surgery.”

Comment: it is unclear whether those patients considered “dropped out” were included in the analysis of the results for symptom score. The drop-out was not balanced and was very likely to be related to lack of efficacy

| Selective reporting (reporting bias) | High risk | Comment: methods section did not specify which and how outcomes would be reported. Numerical data for 3 symptom scores (facial pain, loss of sense of smell and headache) was not shown. The total score on the VAS across 6 symptoms was not reported in the results (although it was used as a criterion to determine eligibility for surgery)

Methods states that “Mann Whitney test was used to test a difference in CT score between treatment groups at the end of study”, and “Wilcoxon signed rank test to test changes in CT score from baseline in either treatment group” but results reports mean difference and SD

| Other bias | Unclear risk | Quote: “VAS of 0-100 mm”, “Lund Mackay”

Comment: standard scales used for measuring outcomes. Inadequate information on the baseline characteristics to judge. There were slightly more current smokers, atopic patients and patients who had had surgery in the placebo group. Baseline data for outcomes not reported; e.g. “…the CT scores in the FPND patients were better at the start of the study” (page 1022)
### Methods

<table>
<thead>
<tr>
<th>Participants</th>
<th>4 arm, &quot;double blind&quot;, international, multicentre, parallel-group RCT, with 4 months' duration of treatment and follow-up</th>
</tr>
</thead>
</table>

| Location: 9 countries: Colombia, Guatemala, Honduras, Panama, Peru, Russia, South Africa, Ukraine, United States. No. of sites not presented |
| Setting of recruitment and treatment: not stated |
| Sample size: 6 to 11 years |
| Number randomised (6 to 11 years): 18 in intervention 1, 18 in intervention 2, 10 in comparison |
| Number completed (6 to 11 years): no information |
| Number randomised (12 to 17 years): 32 in intervention 1, 33 in intervention 2, 16 in comparison |
| Number completed (12 to 17 years): no information |

<table>
<thead>
<tr>
<th>Interventions</th>
<th>6 to 11 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervention 1 (n = 18): mometasone furoate nasal spray, 100 µg once per day for 4 months</td>
<td></td>
</tr>
<tr>
<td>Intervention 2 (n = 18): mometasone furoate nasal spray, 100 µg twice per day for 4 months</td>
<td></td>
</tr>
<tr>
<td>Comparator group (n = 9): placebo once or twice daily (combined), for 4 months</td>
<td></td>
</tr>
</tbody>
</table>

| Participants (baseline) characteristics: 6 to 11 years |
| Age: twice daily group 9.6, once daily group 9.7, PL group 12.7 |
| Gender M/F: twice daily group 5/13, once daily group 8/10, PL group 12/14 |
| Main diagnosis: nasal polyps |
| Polyps status: 100% with polyps |
| Previous sinus surgery status: no information |
| Other important effect modifiers: asthma, eosinophilic |

| Participants (baseline) characteristics: 12 to 17 years |
| Age: twice daily group 14.4, once daily group 14.4, PL group 12.7 |
| Gender M/F: twice daily group 15/18, once daily group 14/18, PL group 12/14 |
| Main diagnosis: bilateral nasal polyps |
| Polyps status: 100% with polyps |
| Previous sinus surgery status: no information |
| Other important effect modifiers: asthma, eosinophilic |

### Inclusion criteria:

- Children aged 6 to 17 years with nasal polyposis

### Exclusion criteria:

- Children younger than 6 years. Antrochoanal polyps, cystic fibrosis, acute rhinosinusitis, rhinitis medicamentosa, dyskinetic ciliary syndromes and aspirin allergy

Participants with asthma who received inhaled corticosteroids were required to be on no more than a moderate dosage regimen as defined by the 2005 Global Initiative for Asthma Guidelines (GINA) for 1 month before screening and to remain on it throughout the study (16); other forms of corticosteroids were prohibited.
Intervention 1 (n = 26): mometasone furoate nasal spray, 200 µg once per day for 4 months
Intervention 2 (n = 32): mometasone furoate nasal spray, 200 µg twice per day for 4 months
Comparator group (n = 16): placebo once or twice daily (combined) for 4 months
Use of additional interventions (common to both treatment arms): inhaled corticosteroids for patients with asthma (up to the equivalent of moderate dosage regimen according to GINA 2005)

Outcomes of interest in the review:
All outcomes were measured at 4 months
Primary outcomes
1. Disease severity, measured as participant-rated signs/symptoms including nasal congestion/obstruction, anterior rhinorrhoea/postnasal drip and loss of sense of smell; rated daily by participants on a 4-point scale
2. Significant adverse effect: epistaxis
Secondary outcomes:
3. Other adverse effects: local irritation (including oral thrush, sore throat)
4. Polyps size; no details on scores used
Other outcomes reported by the study:
(Primary outcome) Effects on hypothalamic-pituitary-adrenal (HPA) axis function (24-hour urinary free cortisol change from baseline and 24-hour urinary free cortisol corrected for creatinine/adverse events
Investigator-evaluated polyp size (on a 4-point scale)
Investigator assessment of overall therapeutic response (on a 5-point scale ranging from 0 (complete relief) to 4 (no relief)

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “Subjects were randomly assigned to one of four treatment groups in a 4:4:1: 1 ratio... stratified by age” Comment: pg 34, col 1, para 4</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Comment: no information about allocation concealment provided</td>
</tr>
</tbody>
</table>
| Blinding of participants and personnel (performance bias) | Unclear risk       | Quote: “received MFNS 200 mcg once daily, MFNS 200 mcg twice daily, placebo once daily, or placebo twice daily” Comment: the abstract mentioned “double-blind” and a placebo was used. However, instead of using a double-dummy design, where all participants received the medication twice daily (with a placebo
given for those who had once daily treat-
ment), groups either had medication once
or twice daily. Therefore, there was no
blinding of participants in terms of know-
ing whether they were on the once daily or
twice daily regimen

| Blinding of outcome assessment (detection bias) | Unclear risk | Quote: as above
Comment: most of the outcomes are pa-
tient-reported and therefore blinding of
outcome assessment is affected |
| All outcomes |

| Incomplete outcome data (attrition bias) | Unclear risk | Comment: no information about loss to
follow-up or exclusion. However, only 119/
127 randomised patients (93%) were in-
cluded in their primary endpoint analy-
sis. There were more exclusions/drop-outs
from the 100 µg group compared with the
higher-dose group (6 (12%) versus 1) but
no reasons were provided
Adverse effects and symptoms were re-
ported based on 127 participants. It is un-
clear whether there were any imputations |
| All outcomes |

| Selective reporting (reporting bias) | High risk | Quote: “No statistical analysis of effi-
cacy end points was pre-specified, in the
study protocol, and only descriptive effi-
cacy statistics were collected.”
Comment: we identified the protocol
(NCT00378378) and the purpose was "to
evaluate the safety and efficacy of Na-
sonex® (Mometasone Furoate Nasal Spray
(MFNS)) in the treatment of nasal polyps
in pediatric subjects between the ages of 6
and less than 18 years old. Safety will be
the primary focus of this study.” The study
only reported the change from baseline in
points and percentages but not the standard
deviations and P values. The values from
the treatment groups were very similar to
the placebo group for some outcomes (e.g.
for rhinorrhoea -43% for once daily versus
-42%) and poor reporting due to lack of
beneficial effects cannot be ruled out
Results for the "young (6-11 years)” group
and the "older (12-17 years)” group were
pooled together for the adverse events re-
sults and compared against the results sep-
arated by age group. This does not appear
### Hansen 2010

**Methods**
- 2-arm, double-blind, placebo-controlled, parallel-group RCT with 12 weeks of treatment and 14 weeks of follow-up

**Participants**
- **Location:** Netherlands, single site
- **Setting of recruitment and treatment:** ENT clinic in the Netherlands (Academic Medical Centre, Amsterdam)
- **Sample size:**
  - **Number randomised:** 10 in intervention, 10 in comparison
  - **Number completed:** 9 in intervention, 7 in comparison
- **Participant (baseline) characteristics:**
  - Age, mean (range): FP: 49.2 (25 to 61); PL: 46.7 (37 to 62)
  - Gender, M/F: FP: 6/4; PL: 8/2
  - Main diagnosis: recalcitrant CRS without nasal polyps or only cobble-stoned mucosa
    - Polyps status: no nasal polyps
    - Previous sinus surgery status: all had surgery before
      - Sinus surgery, median (range): FP: 4 (1 to 10); PL: 3 (1 to 8)
      - Polypectomy (%): FP: 2 (20); PL: 0
  - Previous courses of steroids: no information
  - Other important effect modifiers:
    - Current asthma, n (%): FP: 4 (40); PL: 3 (30)
    - Allergy, n (%): FP: 5 (50); PL: 5 (50)
    - ASA intolerance, n (%): FP: 3 (30); PL: 1 (10)
- **Inclusion criteria:**
  - Between 18 and 65 years of age
  - Chronic rhinosinusitis defined as at least a 12-week history of 2 or more of: blockage/congestion, discharge: anterior/post nasal drip, facial pain/pressure, reduction or loss of sense of smell and either mucopurulent discharge from the middle meatus or oedema/mucosal obstruction primarily in the middle meatus
- **Exclusion criteria:**
  - Visible nasal polyps on endoscopy, except cobble-stoned mucosa
  - Surgical treatment for nasal polyps during the previous 3 months
  - A diagnosis of cystic fibrosis
  - Depot or oral steroids during the previous 2 months
  - A requirement for more than 1000 µg beclometasone (or equivalent) per day for the treatment of asthma, or not on a stable dose for ≥ 3 months

**Interventions**
- **Intervention (n = 10):** fluticasone propionate, administered using a breath actuated inhaler (Optinose) 400 µg twice daily (800 µg total daily dose), duration of treatment
- **Comparator group (n = 10):** matching placebo, administered twice daily

**Use of additional interventions (common to both treatment arms):**
Participants using saline rinses were permitted to continue to do so; 5 in the placebo...
Continued

A total of 50 participants were included in the review: 27 in the control group and 7 in the treatment group. Participants continued using nasal saline irrigation twice daily during the study. Loratadine 10 mg tablets were provided as rescue medication. If a participant experienced a severe acute nasal blockage, the investigator could authorize the use of a short course of oxymetazoline drops or spray for a maximum of 7 consecutive days and a maximum total of 10 days during the treatment period. Oxymetazoline was not to be used within 24 hours of a scheduled study visit.

Outcomes of interest in the review:

Primary outcomes:
1. Health-related quality of life, disease-specific using RSOM-31. A reduction in the average total symptom impact score > 1 is considered clinically relevant.
2. Disease severity symptom score, measured using:
   a. A 10 cm VAS (not troublesome to most troublesome imaginable) “How troublesome are your symptoms of rhinosinusitis?”
   b. A diary (0 to 3 scale): 0 (none), 1 (mild - symptoms present but not troublesome), 2 (moderate - symptoms frequently troublesome but not interfering with daily activity or night-time sleep) or 3 (symptoms troublesome and interfering with daily activity or night-time sleep) to record nasal blockage, nasal discomfort and rhinitis symptoms. Participants also recorded sense of smell: 0 (normal), 1 (slightly impaired), 2 (moderately impaired) or 3 (absent).
3. Significant adverse effect: epistaxis

Secondary outcomes:
4. Endoscopy (polyps size or overall score) using the Lund-Mackay score

Other outcomes reported by the study:
- Peak nasal inspiratory flow (PNIF)
- Acoustic rhinometry
- MRI scans of the paranasal sinuses
- Use of rescue medication

Notes:
Study had a 14- to 16-day treatment-free run-in period at the beginning

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation</td>
<td>Unclear risk</td>
<td>Quote: “…subjects who met the eligibility criteria were randomized 1:1 …” Comment: no information on randomisation method provided</td>
</tr>
<tr>
<td>Allocation concealment</td>
<td>Unclear risk</td>
<td>Comment: no information on how to maintain allocation concealment</td>
</tr>
<tr>
<td>Blinding of participants and personnel</td>
<td>Low risk</td>
<td>Quote: “…The Opt-FP and placebo devices were identical in appearance. The spray pump in the Opt-FP contained … Placebo matched FP exactly, except for the active ingredient... To deliver a dose of FP</td>
</tr>
</tbody>
</table>

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Hansen 2010  *(Continued)*

<table>
<thead>
<tr>
<th>Source of bias</th>
<th>Risk of bias</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Quote: as above</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td>Comment: key outcomes are patient-reported - should remain well blinded until end of study</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Quote: “…The majority of subjects (9 Opt-FP 7 PBO) completed the study.”</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td>Comment: the drop-out rate is high (20%) and unbalanced (10% in active group and 30% in placebo group) enough to affect the findings of this small study (results were presented as medians). All withdrawals were related to adverse effects or worsening of symptoms</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Comment: unclear why study reported “combined nasal symptoms” (rhinitis and blockage) only, whereas other symptoms (discomfort and smell) were reported separately. This was not prespecified. Reporting of some outcomes was confusing or incomplete. The results for the for “symptoms of rhinosinusitis” as measured on a 10 cm VAS were reported as having reduced 13 points in the treatment group. It was not clear if this was a mistake or whether it was 13%. For endoscopic evaluation, the study only showed data for the oedema score, which was statically significant, but not the nasal discharge score, which was not statistically significant</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>Comment: instruments for the main patient-reported symptoms were validated</td>
</tr>
</tbody>
</table>
### Methods

3-arm, double-blind, parallel-group RCT, with a 26-week duration of treatment and 2 additional weeks of follow-up

### Participants

**Location:** Sweden  
**Setting of recruitment and treatment:** outpatient clinics; single-centre  
**Sample size:**  
Number randomised: 19 in FP group, 18 in BDP group, 18 in PL group  
Number completed: 15 in FP group, 16 in BDP group, 11 in PL group  
**Participant (baseline) characteristics:**  
Age mean (range): FP group: 54 (27 to 74); BDP group: 49 (26 to 68); PL group: 47 (21 to 71)  
Gender (M/F): FP group: 15/4; BDP group: 13/5; PL group: 14/4  
Main diagnosis: bilateral polyposis with a polyp score of 1 or 2  
Polyps status: 100% with polyps  
Previous sinus surgery status: 100% had a history of at least 1 polypectomy within the previous 5 years  
Other important effect modifiers:  
- Positive skin prick test (%): FP group: 3 (16%); BDP group: 6 (33%); PL group: 5/18 (27%)  
**Inclusion criteria:** bilateral polyposis with a polyp score of 1 or 2  
**Exclusion criteria:** nasal polyposis with a score of 3 or 4 (or 0); concurrent nasal infection; an inability to cease treatment with systemic, inhaled or intranasal steroids or sodium cromoglycate on visit 1; had used antihistamines in the 48 hours prior to visit 1; had a contraindication to steroids or had any serious or unstable concurrent disease

### Interventions

**FP group (n = 19):** fluticasone propionate, aqueous nasal spray, 2 actuations of 50 µg each to each nostril morning and evening (400 µg/day) for 26 weeks  
**BDP group (n = 18):** beclomethasone dipropionate, aqueous nasal spray, 2 actuations of 50 µg each to each nostril morning and evening (400 µg/day) for 26 weeks  
**PL group (n = 18):** placebo, actuations to each nostril morning and evening, containing the same vehicle as the intervention solutions including benzalkonium chloride as a preservative, for 26 weeks  
**Use of additional interventions (common to all treatment arms):**  
A 4-week run-in period during which there was no treatment for polyposis except for rescue loratadine, which could be used by the patients  
All patients were supplied with rescue loratadine tablets to use as relief medication, 10 mg loratadine once daily. Any use of rescue medication was documented on the patients’ daily record cards

### Outcomes

**Outcomes of interest in the review:**  
**Primary outcomes:**  
1. Disease severity, measured by daily records of all nasal symptoms including: nasal blockage; sense of smell; sneezing and rhinorrhea using a 4-point rating system (0 = no symptoms; 1 = mild symptoms; 2= moderate symptoms; 4 = severe symptoms)  
2. Physician assessment of symptoms. No details were provided on how these were measured. Measured at 26 weeks  
3. Significant adverse effect: epistaxis  
**Secondary outcomes:**  
4. Polyp size by endoscopy (0- to 4-point scale)
Other outcomes reported by the study:
5. Polyp score
6. Peak nasal inspiratory flow
7. Physician’s assessment of change in symptoms

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Quote: “randomized”</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Comment: pg 271, col 1, para 3</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>Comment: no information provided in the paper</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Low risk</td>
<td>Comment: no further information. Should also be low if there is adequate blinding</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>High risk</td>
<td>Comment: 13/54 patients (24%) did not complete trial; 4/19 in fluticasone, 2/18 in beclomethasone, 7/18 (39%) in placebo group. Uneven drop-out numbers (very high in placebo group)</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>Quote: “The primary efficacy endpoint was the physician’s assessments of symptoms and polyp score on all clinic visits”</td>
</tr>
</tbody>
</table>

Notes

Risk of bias

Holmberg 1997 (Continued)
Holmberg 1997  (Continued)

polyps (only “significant” for visit 5 on beclomethasone, not for fluticasone). In addition, there were some outcomes that seemed to have arbitrary, non-predefined cut-off points (% of days with symptom score < 2 in results). The denominator for the reported symptom scores outcome measures is not identified.

Other bias

High risk

Comment: primary outcome of physician assessment of outcomes was not well described in the paper with little information on the criteria used or any validation/inter-rater reliability

Holopainen 1982

Methods

2-arm, double-blind, parallel-group RCT, with a 16-week duration of treatment and follow-up

Participants

Location: Sweden, no information on number of sites
Setting of recruitment and treatment: unclear
Sample size:
Number randomised: 10 in intervention, 9 in comparison
Number completed: 10 in intervention, 8 in comparison
Participant (baseline) characteristics:
Age mean (range): group A: 43.5 (26 to 60); group B: 40 (18 to 62)
Gender (M/F): group A: 6/4; group B: 4/5
Main diagnosis: perennial, intrinsic nasal symptoms associated with small nasal polyps
Polyps status: 100% with polyps
Previous sinus surgery status: unclear (there is a comment, “When necessary the number of polyps was reduced by surgical measures so that the test solution could easily be administered”, but no further details are given)
Other important effect modifiers: none provided
Inclusion criteria: no further details available
Exclusion criteria: none stated

Interventions

Intervention (n = 10): budesonide nasal spray 400 µg daily, 2 puffs into each nostril, twice a day, for 16 weeks
Comparator group (n = 9): placebo nasal spray (same solvent as intervention but without the active ingredient), 2 puffs into each nostril, twice a day, for 16 weeks
Use of additional interventions (common to both treatment arms): all patients underwent a wash-out period of 2 weeks before the study

Outcomes

Outcomes of interest in the review:
Primary outcomes
1. Disease severity, measured by patient-reported symptom score cards recording nasal blocking, running, sneezing, itching and side effects according to a 0 to 3 scale daily for 2 weeks prior to check-up. Last check-up at 16 weeks
2. Disease severity, measured by physician rhinoscopy to assess mucosal congestion and nasal discharge.
3. Significant adverse effect: epistaxis
Secondary outcomes:
4. Size of polyps (on a 0 to 3 scale)
Other outcomes reported by the study:
- Saccharin test for measuring mucociliary activity
- Nasal smear for evaluating epithelial changes
- Biopsy of the nasal polyps
- Plasma cortisol determination
- Peak nasal inspiratory flow

Notes
Although the paper states “When necessary the number of polyps was reduced by surgical measures so that the test solution could be easily administered”, there was no report of this having been carried out.

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Quote: “…randomly assigned…” Comment: no further information</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Comment: no information</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>Quote: “The placebo was identical with the active spray but without budesonide… the other a correspondent dose of only the solvent” Comment: identical-looking and solvent used</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Low risk</td>
<td>Comment: low risk, since blinding is adequate</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>Low risk</td>
<td>Quote: “One patient with severe nasal blocking and obstructing polyps had to withdraw from the trial after 12 weeks of placebo treatment.” This patient was not reported in the safety outcomes Comment: only 1 drop-out (5%). Unlikely to have an important impact on outcomes</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>Comment: methods section reports that nasal discharge would be physician-assessed by rhinoscopy (pg 222, bullet point 1). No results are reported for this outcome</td>
</tr>
</tbody>
</table>
### Holopainen 1982 (Continued)

<table>
<thead>
<tr>
<th>Other bias</th>
<th>Unclear risk</th>
<th>Comment: no information regarding the validation of the disease severity measures. Only limited information provided in the study about baseline characteristics, pre-randomisation procedures etc</th>
</tr>
</thead>
</table>

### Johansen 1993

<table>
<thead>
<tr>
<th>Methods</th>
<th>3-arm, double-blind, multicentre, parallel-group RCT, with 3 months duration of treatment and follow-up</th>
</tr>
</thead>
</table>
| Participants | **Location:** 4 sites in Denmark, 1 site in Sweden  
**Setting of recruitment and treatment:** unclear  
**Sample size:**  
**Number randomised:** 91 (numbers allocated to each group unknown)  
**Number completed:** 86 (numbers allocated to each group unknown)  
**Participant (baseline) characteristics:**  
- Age median (range): 52 (18 to 78)  
- Gender (M/F): 70/21  
- Main diagnosis: eosinophilic nasal polyps with polyp score of 2 or less on each side  
- Polyps status: 100% with polyps  
- Previous sinus surgery status: not provided in the paper  
- Other important effect modifiers:  
  - 22 patients had asthma (allocation between groups unknown)  
  - 8 patients were known to be acetylsalicylic acid (ASA) sensitive  
**Inclusion criteria:** clinical diagnosis of eosinophilic nasal polyps with polyp scores of 2 or less on each side. Eosinophilic polyps was confirmed by nasal smear and/or biopsy.  
**Exclusion criteria:**  
Polyps surgically removed within 2 months  
Neutrophilic polyps  
Systemic or topical nasal corticosteroid therapy within 2 months |
| Interventions | **Group A (n = unknown):** budesonide aqua (Rhinocort Aqua), 50 µg in each nostril x 2, twice daily (400 µg/day), 3 months  
**Group B (n = unknown):** budesonide aerosol (Rhinocort Aerosol), 50 µg in each nostril x 2, twice daily (400 µg/day), 3 months  
**Group C (n = unknown):** placebo (aqua or aerosol), unclear dose, 3 months  
**Use of additional interventions (common to all treatment arms):** no information was provided about additional interventions |
| Outcomes | **Outcomes of interest in the review:**  
**Primary outcomes:**  
1. Disease severity, measured weekly by patients. Symptoms included were nasal obstruction, sneezing and nasal secretions, recorded for each nasal cavity (scale 0 to 3). Change in sense of smell was recorded at clinical visits using a 0 to 3 scale  
2. Significant adverse effect: epistaxis |
Secondary outcomes:
3. Other adverse effects: local irritation (including oral thrush, sore throat)
4. Polyps size (assessed using a 0 to 3 scale - definitions provided)
Other outcomes reported by the study:
- Nasal and oral peak inspiratory flow

<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>
| Random sequence generation (selection bias)       | Unclear risk       | Quote: “...randomised…”  
Comment: mentioned in abstract but no further mention |
| Allocation concealment (selection bias)           | Unclear risk       | Comment: no information provided                                                      |
| Blinding of participants and personnel (performance bias) | Low risk          | Quote: “The patients were treated with either budesonide aqua (Rhinocort Aqua) or budesonide aerosol (Rhinocort Aerosol) , 50 mcg x 2 in each nostril, twice daily = 400 mcg/day or placebo aqua or aerosol.”  
Comment: there should be adequate blinding for treatment versus placebo, but not for different forms of treatment |
| Blinding of outcome assessment (detection bias)    | Low risk           | Comment: no further information                                                      |
| Incomplete outcome data (attrition bias)          | Unclear risk       | Quote: “Five patients withdrew from the study…”  
Comment: low (5%) drop-out rate. No reasons given for withdrawals. Patients who withdrew were not included in any of the outcomes (including safety outcomes) |
| Selective reporting (reporting bias)              | Unclear risk       | Comment: all outcomes reported in the methods are mentioned in the results section, but numerical information for the results is not provided |
| Other bias                                        | High risk          | Comment: no comment on the validation of outcome measurements  
The paper does not provide clear background characteristics for each group. The number randomised to each group was not provided |
## Keith 2000

### Methods

2-arm, double-blind, multicentre, parallel-group RCT, with 12-week duration of blinded treatment and 12-week duration of open treatment, with the active intervention followed by a final assessment 2 weeks after treatment had completed.

### Participants

**Location:** 11 sites in Canada and Finland

**Setting of recruitment and treatment:** outpatient clinics

**Sample size:**

- **Number randomised:** 52 in intervention, 52 in comparison
- **Number completed:** 51 in intervention, 47 in comparison

**Participant (baseline) characteristics:**

- **Age (mean ± SD):** FP group: 49 ± 12; PL group: 47 ± 13
- **Gender (M/F):** FP group: 38/14; PL group: 35/17
- **Main diagnosis:** small or medium bilateral nasal polyposis
- **Polyps status:** 100% with polyps
- **Previous sinus surgery status (n (%)):** FP group: 38 (73%); PL group: 34 (65%)
- **Other important effect modifiers:**
  - 22% were atopic and allergic to one or more allergens
  - 2 patients in the study were reportedly aspirin-sensitive (no information on which groups)
  - 52% in FP group and 44% in PL group had a polyposis history of > 10 years

**Inclusion criteria:**

- Outpatients aged 16 years and over with bilateral nasal polyposis.
- Polyps were graded by clinical assessment during rhinoscopic examination. Patients with a severity score of 1 (small) or 2 (medium) were included.

**Exclusion criteria:**

- Large (grade 3) polyps, indicating severe nasal obstruction
- Surgical treatment for nasal polyps during the last 3 months
- Cystic fibrosis
- Purulent nasal infection
- Allergic rhinitis
- Any disease likely to interfere with the study parameters or which gave evidence of any serious or unstable concurrent disease or psychological disorder
- Hypersensitivity or contraindication to steroids
- Currently receiving inhaled corticosteroids or those who had received depot or oral steroids during the previous 3 months
- Unable to cease treatment with intranasal steroids, or inhaled or intranasal sodium cromoglycate, at the screening visit
- Astemizole during the last 6 weeks or other antihistamines within the last 48 hours, or received any other research medication during the previous month
- Pregnant, lactating or, in the investigator's opinion, were not taking adequate contraceptive measures to avoid becoming pregnant during the study
- Had not correctly completed the daily diary card during the run-in period

### Interventions

**Intervention (n = 52):** fluticasone propionate (unpreserved), nasal drops using head down and forwards position, 400 µg divided between both nostrils, once daily in the morning for 12 weeks

**Comparator group (n = 52):** placebo nasal drops using head down and forwards position, to both nostrils once daily in the morning for 12 weeks

**Use of additional interventions (common to both treatment arms):** loratadine tablets were provided as rescue medication for the relief of troublesome symptoms of rhinitis, to be used as needed, at a maximum dose of 10 mg once daily. No other medication was provided.
Outcomes

**Outcomes of interest in the review:**

**Primary outcomes**

1. Disease severity, patient-reported through daily diaries (nasal blockage, nasal discomfort and rhinitis symptoms) measured on a 4-point scale (0 = none, 1 = mild, 2 = moderate, 3 = severe) at 12 weeks. Sense of smell was recorded as 0 (normal), 1 (slightly impaired), 2 (moderately impaired) or 3 (absent).

2. Significant adverse effect: epistaxis

**Secondary outcomes:**

3. Other adverse effects: local irritation (including oral thrush, sore throat)

4. Polyp size (0 to 3 score): 0 (no polyps), 1 (mild polyposis) = small polyps not reaching the upper edge of the inferior turbinate and causing only slight obstruction, 2 (moderate polyposis) = medium polyps reaching between the upper and lower edge of the inferior turbinate and causing troublesome obstruction, 3 (severe polyposis) = large polyps reaching below the lower edge of the inferior turbinate and causing almost/total obstruction

**Other outcomes reported by the study:**

- Peak nasal inspiratory flow
- Olfactory function
- Daily use of loratadine tablets

Notes

- Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: &quot;… computer randomized number …”</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comment: proper sequence generation</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Quote: “… Each investigator was given a block of treatments, precoded with computer randomized numbers, which were assigned in ascending numerical order as patients presented.”</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comment: allocation concealment likely to be well maintained despite blocked randomisation; adequate blinding</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Quote: &quot;… FPND (400 mg unit dose preservative-free suspension) and placebo solution were supplied in identical opaque nasal drop containers, in a foil pack …”</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td>Comment: adequate blinding</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Comment: most outcomes are patient-reported therefore blinding was likely to be well maintained</td>
</tr>
</tbody>
</table>
Incomplete outcome data (attrition bias)  
All outcomes  
Low risk  
Comment: 6/104 (5.8%) patients did not complete the study, 2 of these patients required polypectomy and so the last available outcome measures were reported for the efficacy outcomes. All patients were included in the safety analysis.

Selective reporting (reporting bias)  
Unclear risk  
Comment: results are presented in a different format to those presented in the methods section. For example the methods provides scales for each symptom but the results presents the percentage of time the value was below a certain value on the scale. It was unclear if this change was pre-specified in the protocol or whether the study authors decided once the results had been processed.

Other bias  
Unclear risk  
Comment: no information is available regarding the validation of the scales used.

Lang 1983

Methods  
2-arm, double-blind, parallel-group RCT, with a 2-year duration of treatment and follow-up

Participants  
Location: unclear  
Setting of recruitment and treatment: unclear  
Sample size:  
Number randomised: 14 in intervention, 18 in comparison  
Number completed: unclear number of participants completed  
Participant (baseline) characteristics:  
Age (mean): 42  
Gender (M/F): 17/15  
Main diagnosis: clinical simple nasal polyps  
Polyps status: 100% with polyps  
Previous sinus surgery status: no details  
Other important effect modifiers: no details  
Inclusion criteria: no details  
Exclusion criteria: no details

Interventions  
Intervention (n = 14): beclomethasone dipropionate 400 µg twice daily for 2 years. No information on method of administration other than “Beconase topically in the nose”  
Comparator group (n = 18): placebo nasal insufflation twice daily for 2 years  
Use of additional interventions (common to both treatment arms): none mentioned
### Outcomes of interest in the review:

**Primary outcomes**
1. Disease severity, measured by subjective assessment of nasal obstruction, sneezing and nasal discharge. No details of scale used. Assessment made every 4 weeks for 2 years

**Secondary outcomes:**
2. Size of nasal polyps - reported as the number of patients with "resolution"

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
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<td>Unclear risk</td>
<td>Support for judgement</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Quote: “… randomly allocated …” Comment: no mention of method of randomisation</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>Comment: no information provided</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>Quote: “This allocation was kept blind from both patient and investigator.” Comment: mentions a &quot;placebo&quot; was used but there was no information about precautions taken to make the products as similar as possible</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>Quote: “This allocation was kept blind from both patient and investigator.” Comment: method not specified</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>No information was provided about losses to follow-up. Unlikely to have had no losses during a 2-year study</td>
</tr>
<tr>
<td>Other bias</td>
<td>High risk</td>
<td>Comment: abstract only. No information provided about adverse events. Very limited information provided</td>
</tr>
</tbody>
</table>
### Methods

**Location:** UK  
**Setting of recruitment and treatment:** tertiary referral centre (Royal National ENT Hospital London)  
**Sample size:**  
**Number randomised:** 10 in fluticasone propionate, 10 in beclomethasone dipropionate, 9 in placebo  
**Number completed:** unclear, likely to be all  
**Participant (baseline) characteristics:**  
- **Age (mean, range):** 52 (32 to 71), 46 (22 to 67) and 50 (27 to 69) in fluticasone propionate, beclomethasone dipropionate and placebo arms  
- **Gender (M/F):** 7/3, 9/1 and 7/2 in fluticasone propionate, beclomethasone dipropionate and placebo arms  
- **Main diagnosis:** “severe polyposis”  
- **Polyps status:** all had polyps, median total polyps score of 4 (both nostrils) using Lund-Mackay CT score  
- **Previous sinus surgery status:** 66% had surgery (7/10 in fluticasone propionate and beclomethasone dipropionate arms, 5/9 in placebo)  
- 59% had condition for more than 10 years  
- All had allergy  
**Inclusion criteria:**  
- Older than 16 years with a diagnosis of bilateral nasal polyposis requiring surgical intervention, meeting one or more of the following criteria:  
  - a total polyp score of 4 or higher plus a CT scan score > 12;  
  - a total polyp score of 3 or higher, a nasal blockage score of 2 or higher, plus a CT scan score > 12; and  
  - a total polyp score of 2 or higher, a nasal blockage score of 2 or higher, a CT scan > than 12, plus an UPSIT score > 32  
**Exclusion criteria:**  
- Concurrent purulent nasal infection  
- A requirement for more than 1000 µg beclomethasone (or equivalent) per day for the treatment of asthma  
- An inability to cease treatment with parenteral and intranasal corticosteroids or cromolyn sodium (sodium cromoglycate) at visit 1, used astemizole in the 6 weeks before the study or other antihistamines in the 48 hours before visit 1, or a contraindication to corticosteroid medications

### Participants

**Interventions**  
**Intervention 1 (n = 10):** fluticasone propionate aqueous nasal spray 400 µg per day, 2 actuations into each nostrils morning and night  
**Intervention 2 (n = 10):** beclomethasone dipropionate aqueous nasal spray 400 µg per day, 2 actuations into each nostrils morning and night  
**Comparator (n = 9):** placebo 2 sprays into each nostril twice a day  
**Use of additional interventions (common to both treatment arms):** terfenadine 60 mg as rescue medicine

### Interventions

**Outcomes**  
**Outcomes of interest in the review:**  
**Primary outcomes:**  
- Disease severity - collected patient diaries on a 0 to 4 scale for different symptoms, but only partially reported symptom-free days
### Secondary outcomes:
- Adverse events - local irritation
- Endoscopy - polyps size (scale not reported)

### Other outcomes reported by the study:
- PNIF, physician-reported score for symptom severity

#### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “Patients were randomly allocated, using a computer-generated random code and a block size of 6, to receive 1 of 3 treatments”</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Quote: “Patients were randomly allocated, using a computer-generated random code and a block size of 6, to receive 1 of 3 treatments”</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comment: method not specified; blocked randomisation but adequate blinding</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Quote: “The placebo was identical to the active formulations with the active ingredient omitted and was indistinguishable from the active treatments, which were themselves identical in appearance, taste, and smell.”</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comment: there was a 4-week pre-treatment period where all patients were exposed to the placebo, but blinding should still be adequate</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Comment: the same investigator did all the clinical assessments at all visits, but an identical placebo was used</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Quote: &quot;last value carried forward technique&quot; was used. Drop-outs not balanced: 3/10 in fluticasone, 0/10 in beclomethasone and 4/9 in placebo</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>Comment: patient-reported symptoms (using diaries) were collected, but it was not specified how these were planned to</td>
</tr>
</tbody>
</table>
be reported. Study only reported percentage of patients with 100% of days without nasal blockage, and the median % of days without nasal symptoms (different criteria). Other outcomes not reported at all.

There was also a higher percentage of patients in the fluticasone group (70%) compared to 33% and 30% in the beclomethasone and placebo groups, but details were not reported. Only stated that 1 of the adverse events in the fluticasone group (throat irritation) was “predictable”.

Other bias

<table>
<thead>
<tr>
<th>High risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quote: “overall rhinitis symptoms (sneezing, rhinorrhoea, nasal itching)”</td>
</tr>
<tr>
<td>Comment: symptoms scores (by patients and clinicians) were used but no mention of validation. Some items seem to be single symptom (e.g., nasal blockage), but others seem to encompass a few things (e.g. “overall rhinitis symptoms”)</td>
</tr>
<tr>
<td>Quote: “There was evidence, particularly from the acoustic rhinometric and PNIF data, that the patients randomly allocated to receive BDANS had milder symptoms than those randomly allocated to receive FPANS or placebo, even though all patients had been listed for surgical treatment on an equal basis before the study.”</td>
</tr>
<tr>
<td>Comment: baseline symptoms and other assessment scores were not reported. Unable to judge for other aspects</td>
</tr>
</tbody>
</table>

### Lund 2004

<table>
<thead>
<tr>
<th>Methods</th>
<th>Double-blind, multicentre, parallel-group RCT, with 20 weeks of treatment</th>
</tr>
</thead>
</table>
| Participants | **Location:** multicentre (19), UK (7), Hungary (6), South Africa (6)  
**Setting of recruitment and treatment:** all were ENT specialists, except 1 (in South Africa)  
**Sample size:**  
**Number randomised:** 81 in intervention, 86 in comparison  
**Number completed:** 67 in intervention, 67 in comparison  
**Participant (baseline) characteristics:**  
Age (mean): group A: 38; group B: 43  
Gender (M/F): group A: 35/46; group B: 41/45  
Main diagnosis: patients aged 18 years or over with chronic rhinosinusitis  
Polyps status: 0% with polyps |
Previous sinus surgery status: not provided

**Inclusion criteria:**
No nasal polyposis
\( \geq 18 \) years, with \( \geq 12 \) weeks with at least 2 major symptoms
Patients with a symptom score of \( \geq 2 \) on a 4-point scale for at least 1 of the symptoms for \( \geq 4 \) of 7 days during the last 7 days of the run-in period

**Exclusion criteria:**
Sinonasal surgery within the previous 12 months

### Interventions

**Intervention (n = 81):** budesonide aqueous nasal spray, 128 \( \mu \)g (64 \( \mu \)g in each nostril twice daily), for 20 weeks

**Comparator group (n = 86):** placebo nasal spray, for 20 weeks

**Use of additional interventions (common to both treatment arms):**
During the first 2 weeks of the run-in period, all patients received co-amoxiclav 250/125 mg three times daily, or 500 mg erythromycin twice daily
The same antibiotics could be given for 2 weeks as needed to treat exacerbations (defined as episodes of worsening symptoms requiring a course of antibiotic therapy)

### Outcomes

**Outcomes of interest in the review:**
Primary outcomes
1. Disease severity, measured by Chronic Sinusitis Survey (CSS) questionnaire at baseline and 20-week time point (note: English-speaking participants only as the survey was not validated in Hungarian) and patient-reported scores for individual symptoms
2. Disease severity, measured by patient-reported symptoms (facial pain, pressure or headache; facial congestion, nasal obstruction or blockage; nasal discharge; impairment of sense of smell). A combined symptom score (sum of the scores for the 4 domains of the symptoms above)
3. Significant adverse effect: epistaxis

Secondary outcomes:
4. Health-related quality of life, measured with the SF-36 at baseline and 20-week time point
5. Other adverse effects: local irritation (including oral thrush, sore throat)

Other outcomes reported by the study:
- Compliance with medication
- Overall patient-reported evaluation of efficacy
- Peak nasal inspiratory flow
- Skin prick test before and after treatment
- Blood tests

### Notes

“Financial support” from AstraZeneca

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “Randomisation was performed in balanced blocks of 4 by means of a computer program at the Department of Biostatistics…”</td>
</tr>
</tbody>
</table>
### Lund 2004 (Continued)

<table>
<thead>
<tr>
<th>Bias Type</th>
<th>Risk Level</th>
<th>Details</th>
</tr>
</thead>
</table>
| Allocation concealment (selection bias)        | Low risk   | Quote: “Patients were allocated a treatment number in consecutive order and randomisation was performed in balanced blocks of 4 by means of a computer program…”  
“The treatment codes were known only to the persons responsible for packaging, who were not involved in the study in any other way” |
| Blinding of participants and personnel (performance bias) | Low risk   | Quote: “BANS and placebo aqueous sprays were identical in appearance and were both administered via the same vehicle.”  
“The treatment codes were known only to the persons responsible for packaging, who were not involved in the study in any other way” |
| Blinding of outcome assessment (detection bias) | Low risk   | Quote: “The treatment codes were known only to the persons responsible for packaging, who were not involved in the study in any other way” |
| Incomplete outcome data (attrition bias)       | High risk  | Comment: 14/81 (17.3%) in intervention arm and 19/86 (22.1%) in comparison arm did not complete the study (overall rate 20%). No reasons for dropping out were provided |
| Selective reporting (reporting bias)           | Unclear risk | Comment: the methods section outlines the individual symptoms assessed, but the results are also presented as a “combined score” for which no information is available as to how it was calculated. Additionally, although all outcomes presented in the methods section are mentioned in the outcomes, this is not always in great detail, for example the results for the SF-36 were presented as “There was a significant improvement in the general health sub-scale of the SF-36 questionnaire in the BANS treated group compared with placebo, but no other significant differences were observed.” |
### Lund 2004

**Other bias**

| Unclear risk | Comment: no information about the validation of the combined symptom score. There is some information about validation of the chronic sinusitis score. 2 subgroups were reported (those with CT evidence of opacification and allergic versus non-allergic patients). Not all outcomes are presented for these subgroups. There were pre-randomisation procedures but these excluded patients with less symptomatic disease. |

---

### Mosges 2011

**Methods**

| 2-arm, double-blind, multicentre, parallel-group RCT, with a 16-week duration of treatment and follow-up |

**Participants**

| Location: Germany, 9 sites  
Setting of recruitment and treatment: ear, nose and throat departments in university hospitals or ENT specialists' practices  
Sample size:  
Number randomised: 30 in intervention, 30 in comparison  
Number completed: 29 in intervention, 30 in comparison  
Participant (baseline) characteristics:  
Age: group A: 40 (19 to 63); group B: 44 (22 to 64)  
Gender (M/F): group A: 10/19; group B: 17/13  
Main diagnosis: chronic rhinosinusitis (symptoms for a period longer than 8 weeks, or more than 4 episodes of a minimum length of 10 days each, over a 1-year period; Lund score \( \leq 10 \))  
Polyps status: 0% with polyps (exclusion criteria)  
Previous sinus surgery status: Discussion states “… only around 10% underwent preceding surgical treatment.” However, surgery within 6 months and extensive surgery were exclusion criteria  
Other important effect modifiers: Discussion states: “Allergy was present in only around 1 in 3 patients, who were distributed evenly between both treatment groups.”. No other details provided  
Inclusion criteria:  
Patients aged 18 to 65 years  
Clinical diagnosis of chronic sinusitis (a total symptom score of at least 5) confirmed at baseline by a coronal CT scan not older than 6 months. The scan was evaluated using the Lund scale  
Exclusion criteria:  
Nasal polyps visible on endoscopic examination at baseline  
Patients with pansinusitis, Lund score > 10  
Undergone nasal surgery within 6 months prior to study enrolment  
Patients who had undergone sinus surgery with opening of the lateral nasal wall at any time  
Inhalant or intranasal steroid therapy within 2 weeks prior to screening; systemic steroid |
therapy within 8 weeks prior to screening; antihistamine use within 12 hours to 14 days prior to screening depending on medication; regular use of decongestants within 24 hours to 3 days depending on medication; acute sinusitis or concurrent acute nasal infection, or upper respiratory tract infection, ongoing or within 2 weeks prior to screening

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Intervention (n = 29): mometasone furoate nasal spray, 200 µg twice daily (morning and evening) to the lateral nasal wall (not to the septum), in the ‘vertex to floor’ position, over 16 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Comparator group (n = 30): placebo nasal spray, twice daily (morning and evening) to the lateral nasal wall (not to the septum), in the ‘vertex to floor’ position, over 16 weeks</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Outcomes of interest in the review:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Primary outcomes</td>
</tr>
<tr>
<td></td>
<td>1. Total symptom score (TSS): sum of the 5 individual symptom score values (rhinorrhoea, postnasal drip, nasal obstruction, facial pain or pressure, and headache). Each symptom score was assessed on a 4-point scale (0 = no symptoms; 1 = mild; 2 = moderate; 3 = severe)</td>
</tr>
<tr>
<td></td>
<td>2. Significant adverse effect: epistaxis</td>
</tr>
<tr>
<td></td>
<td>Secondary outcomes:</td>
</tr>
<tr>
<td></td>
<td>3. Endoscopy score (endoscopic evaluation at every visit)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Other outcomes reported by the study:</th>
</tr>
</thead>
<tbody>
<tr>
<td>4. Patient evaluation of therapeutic response</td>
</tr>
<tr>
<td>5. Compliance of medication used (measured by weighing bottles)</td>
</tr>
<tr>
<td>6. Other adverse events</td>
</tr>
</tbody>
</table>

| Notes | - |

### Risk of bias

<table>
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<tr>
<th>Bias</th>
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<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “All qualified patients were randomized, according to a computer-generated code in a 1:1 ratio…”</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Quote: “all patients who met the protocol requirements were assigned a randomization number that corresponded to the treatment unit they were given… The randomization schedule for blinding of treatments was maintained by the sponsor” Comment: no description of allocation concealment but should be low risk, as there is adequate sequence generation and double-blinding</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Quote: “The bottles of MFNS or placebo for the 16-week treatment period had iden-</td>
</tr>
</tbody>
</table>
Mosges 2011  (Continued)

| All outcomes |  | The randomization schedule for blinding of treatments was maintained by the sponsor, and was disclosed only after study completion and database closure."
|  |  | Comment: sealed envelopes containing individual patient allocations were provided as a safety measure but whether any of these were opened is not stated; it is not likely to be enough to affect blinding since only 6 (10%) withdrew |

| Blinding of outcome assessment (detection bias) | Low risk | Comment: as above |

| Incomplete outcome data (attrition bias) | High risk | Comment: only 6 patients dropped out from the study. However, the study excluded patients with "major protocol violation" (not defined) resulting in an “ITT” population of only 75% and an even lower per protocol population (39% excluded) |

| Selective reporting (reporting bias) | High risk | The following outcomes are stated in the methods section but not reported in paper: "patient compliance, Rhinosinusitis Disability Index, Work Productivity and Activity Impairment questionnaires" Comment: pg 242, col 2, para 1 |

| Other bias | Unclear risk | Comment: there is no mention of the validation of the "total symptom score" measure |

Parikh 2001

| Methods | 2-arm, double-blind, parallel-group RCT, with a 16-week duration of treatment and follow-up |

| Participants | Location: UK, single site  Setting of recruitment and treatment: tertiary ENT clinic  Sample size:  Number randomised: 14 in intervention, 15 in comparison  Number completed: 9 in intervention, 13 in comparison  Participant (baseline) characteristics:  Age: FP: 45.1 ± 10.7; PL: 48 ± 20  Gender (M/F): FP: 2/7; PL: 7/6  Main diagnosis: chronic rhinosinusitis  Polyps status: FP: 2/9; PL: 2/13 |
Parikh 2001 (Continued)

| Previous sinus surgery status, mean ± SD: FP: 3 ± 6.2; PL: 2.8 ± 3.4 |
| Previous courses of steroids: no information |
| Other important effect modifiers |
| - Skin prick test positive: FP: 7/9; PL: 8/11 |
| - Asthma: FP: 2/9; PL: 3/13 |

**Inclusion criteria:**

- More than 3 months history of recurrent discoloured rhinorrhoea (> 2 weeks per episode), accompanied by more than 2 of the following symptoms: nasal obstruction, headache, facial pain, fever
- Endoscopy or CT scan evidence of CRS

**Exclusion criteria:**

- Acute exacerbation in the previous 2 weeks, on oral or depot corticosteroids in the previous 3 weeks, on INCS in the previous 2 weeks
- Other severe concurrent illness

### Interventions

| Intervention (n = 9): fluticasone propionate nasal spray, 400 µg per day, administered twice daily |
| Comparator group (n = 13): placebo, administered twice daily |

**Use of additional interventions (common to both treatment arms):** not described

### Outcomes

**Outcomes of interest in the review:**

**Primary outcomes:**

1. Disease severity symptom score - measured using a 10 cm VAS for blockage, sense of smell, sneezing, discharge from the front of nose, discharge from the back of nose, headache, facial pain. Also used diary cards
2. Significant adverse effect: epistaxis

**Secondary outcomes:**

3. Endoscopy: Lund-Kennedy score
4. Local irritation - itchiness of the nose, throat and ear

**Other outcomes reported by the study:**

- Middle meatus swabs
- Acoustic rhinometry
- Blood tests

### Notes

- -

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “…randomisation code was generated and maintained by personnel in the pharmacy. The investigators were not involved in the process of randomisation” Comment: adequate randomisation</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Comment: no specific information; should be adequate since the investigators were not involved in randomisation and blind-</td>
</tr>
</tbody>
</table>
### Parikh 2001 (Continued)

<table>
<thead>
<tr>
<th>Blinding of participants and personnel (performance bias)</th>
<th>Low risk</th>
<th>Quote: “Placebo spray had benzalkonium chloride in the same concentration as fluticasone propionate, and other had rose scent to mask any differences in smell. The study medications were prepared and supplied by Glaxo…” Comment: identical-looking, with the same composition and smell masking</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Comment: as above, since most outcomes were patient-reported</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Comment: 5/14 (36%) in treatment group and 2/15 (13%) in placebo group dropped out. The percentage is high and unbalanced. Most did not turn up for follow-up</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>Adverse effects such as epistaxis and local irrigation were mentioned as measured by one of the VAS, but were not reported separately. Unclear which symptoms were added up into the overall score for “symptom score” or “diary card score”</td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>Comment: study used a 10 cm VAS for symptom scores. Although not formally validated, it should provide adequate discriminant validity for each item. Unclear which symptoms were added up into the overall score</td>
</tr>
</tbody>
</table>

### Penttilla 2000

<table>
<thead>
<tr>
<th>Methods</th>
<th>3-arm, single-blind, international, multicentre, parallel-group RCT, with a 12-week duration of treatment</th>
</tr>
</thead>
</table>
| Participants | **Location**: 12 centres in Denmark (3 centres), Finland (1 centre) and Sweden (1 centre)  
**Setting of recruitment and treatment**: outpatient clinics  
**Sample size**:  
**Number randomised**: 47 in 400 µg FPND twice daily, 48 in 400 µg FPND once daily, 47 in placebo  
**Number completed**: 45 in 400 µg FPND twice daily, 47 in 400 µg FPND once daily, 41 in placebo  
**Participant (baseline) characteristics**:  
Age: mean 51, range 22 to 83  
Gender: M/F; 107/35 (%M: 75.4%) |
Main diagnosis: nasal polyposis
Polyps status: 100% with polyps
Previous sinus surgery status: 72% previous polypectomy (not within 3 months of trial)

**Inclusion criteria:** at least 16 years old, bilateral mild or moderate nasal polyposis

**Exclusion criteria:** severe polyposis (large polyps reaching below the lower edge of the inferior turbinate, causing total obstruction), concurrent purulent nasal infection, unable to cease treatment with intranasal steroids or sodium cromoglycate during run-in period.

Also excluded: people currently receiving inhaled corticosteroids or who had received depot or oral steroids within previous 3 months, patients who had received astemizole in the 6 weeks prior to the first clinic visit, patients who had undergone nasal polyp surgery in the previous 3 months, patients with hypersensitivity or contraindication to steroids, patients with allergic rhinitis or any other disease likely to interfere with outcomes, patients who were pregnant, lactating or likely to become pregnant during the study period.

### Interventions

- **Intervention A (n = 47):** fluticasone propionate nasal drops (FPND), 400 µg twice daily for 12 weeks
- **Intervention B (n = 48):** fluticasone propionate nasal drops (FPND), 400 µg once daily for 12 weeks plus placebo drops once daily for 12 weeks
- **Comparator group C (n = 47):** placebo nasal drops twice daily for 12 weeks

**Process:** contents were divided between both nostrils (200 µg per nostril) in the head down and forward position

**Use of additional interventions (common to both treatment arms):** all patients underwent a 2-week run-in period during which they ceased all medication for polyposis except loratadine tablets for relief of troublesome symptoms (10 mg daily maximum)

Initial visit: physical and oropharyngeal examinations and details of clinical history
Initial and 12-week visit: blood and urine samples

### Outcomes

**Outcomes of interest in the review:**

**Primary outcomes**
1. Disease severity, measured by assessing nasal blockage (0 to 3 scale) and overall rhinitis symptoms including sneezing, rhinorrhoea and nasal itching (0 to 3 scale) and sense of smell (0 to 3 scale) at 12 weeks after treatment
2. Nasal blockage, overall rhinitis
3. Significant adverse effect: epistaxis

**Secondary outcomes:**
3. Other adverse effects: local irritation
5. Polyps size

**Other outcomes reported by the study:** peak nasal inspiratory flow (PNIF), olfactory function, rescue medication usage and adverse events

### Notes

-
<table>
<thead>
<tr>
<th>Area</th>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low</td>
<td>Quote: &quot;...double blind randomised treatment...&quot;, Figure 1, pg 95</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low</td>
<td>Comment: no further information provided, but this is an “international, multicentre” study in 12 centres across 3 countries with regional monitors. Should have adequate sequence generation procedures</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low</td>
<td>Quote: &quot;active and placebo nasal drops were provided in identical single-dose containers ...&quot;</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low</td>
<td>Comment: no further information provided. Should be adequate with use of adequate double-blinding</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High</td>
<td>Quote: &quot;Sixteen patients were withdrawn during the randomized treatment phase, the majority due to lack of efficacy (five placebo, one FP 400 mg o.d., two FP 800 mg b.i.d.) or adverse events (five placebo, one FP 400 mg o.d., two FP 400 mg b.i.d.). One patient in the placebo group withdrew due to requirement for polypectomy. Two patients withdrew during the open phase, one requiring a polypectomy, the other for unspecified reasons&quot;, pg 97, col 2</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear</td>
<td>Comment: all outcome measures in the methods were discussed in the results section. However the diary card data were interpreted by using different cut-off points, which do not appear to be pre-specified in the methods section. Significant results</td>
</tr>
</tbody>
</table>
### Penttila 2000 (Continued)

<table>
<thead>
<tr>
<th>Other bias</th>
<th>Unclear risk</th>
<th>Comment: no mention of validation of symptom criteria used for the primary outcomes</th>
</tr>
</thead>
</table>

### Small 2005

#### Methods

| 3-arm, double-blind, multicentre, parallel-group RCT, with 4-month duration of treatment and follow-up |

#### Participants

| Location: 44 medical centres “worldwide” |
| Setting: no information |
| Sample size: |
| Number randomised: 122 in 400 µg, 115 in 200 µg, 117 in placebo group respectively |
| Number completed: 109 in 400 µg, 101 in 200 µg, 95 in placebo group respectively |
| Participant (baseline) characteristics: |
| Main diagnosis: bilateral nasal polyps and clinically significant congestion obstruction |
| Age (mean): 400 µg: 48.3; 200 µg: 46.7; placebo: 47.5 |
| Gender (%M/%F): 400 µg: 61/39; 200 µg: 66/34; placebo: 61/39 |
| Polyps status: 100% with polyps |
| Previous sinus surgery status: no information |
| Other important effect modifiers: |
| Asthma history (%): 400 µg: 21; 200 µg: 18; placebo: 21 |
| Perennial allergic rhinitis history (%): 400 µg: 25; 200 µg: 20; placebo: 17 |
| Inclusion criteria: |
| ≥18 years with an endoscopically confirmed diagnosis of bilateral nasal polyps (at least 1 on a scale of 0 to 3) and clinically significant nasal congestion obstruction (average morning score 2 or higher on as scale of 0 to 3 for each of the last 7 days of the 14-day run-in period) |
| If had asthma, had a documented FEV1 ≥ 80% of the predicted value within the 6 months before screening and no asthma exacerbations within 30 days before screening. Those treated with inhaled corticosteroids were required to be on a moderate, stable regimen of beclomethasone dipropionate ≤ 800 mg/d or equivalent for 1 month before screening and to remain on a stable regimen throughout the study period |
| Exclusion criteria: |
| Seasonal allergic rhinitis within the past 2 years |
| Sinus or nasal surgery within the previous 6 months or 3 nasal surgeries (or any surgical procedure preventing an accurate grading of polyps) |
| Presumed fibrotic nasal polyps, or complete or near complete nasal obstruction |
| Nasal septal deviation requiring corrective surgery |
| Nasal septal perforation |
| Acute sinusitis, nasal infection or upper respiratory tract infection at screening or in the 2 weeks before screening; |
| ongoing rhinitis medicamentosa |
| Churg-Strauss syndrome |
Dyskinetic ciliary syndromes
Cystic fibrosis
Glaucoma or a history of posterior subcapsular cataracts; allergies to corticosteroids or aspirin, or any other clinically significant disease that would interfere with the evaluation of therapy

| Interventions | 400 µg group (n = 122): mometasone furoate nasal spray 200 µg twice daily (morning and evening) for 4 months |
| | 200 µg group (n = 115): mometasone furoate nasal spray 200 µg once daily (morning, matching placebo used in the evening) for 4 months |
| | Placebo group (n = 117): placebo nasal spray twice daily (morning and evening) for 4 months |
| Use of additional interventions (common to both treatment arms): | Acetaminophen (paracetamol) was encouraged for analgesic purposes; NSAIDs limited to 5 consecutive days if alternative analgesia was required. Antibiotics were administered for bacterial infections at the discretion of the principal investigator |
| | Concomitant medications that would interfere with study evaluations were not permitted, including nasal sodium cromolyn; nasal atropine or ipratropium bromide; corticosteroids (except oral inhaled corticosteroids for asthma or mild-strength or mid-strength topical corticosteroids for dermatologic purposes); antihistamines; decongestants; topical, oral or ocular anti-inflammatory drugs; or topical nasal or oral antifungal agents |

| Outcomes | Outcomes of interest in the review: |
| | Primary outcomes |
| | 1. Disease severity, patient evaluation of symptoms (congestion/obstruction, loss of sense of smell, anterior rhinorrhea and postnasal drip) measured daily on a diary card on a 4-point scale (0 = none, 3 = severe) |
| | 2. Significant adverse effect: epistaxis |
| | Secondary outcomes: |
| | 3. Other adverse effects: local irritation |
| | Other outcomes reported by the study: |
| | Polyps grade; bilateral score and proportion of patients demonstrating an improvement at endpoint |
| | Therapeutic response (rated by investigator) |
| | Peak nasal inspiratory flow |
| | Treatment compliance |
| | Number of withdrawals due to AE and events occurring in more than 2% of participants in any group |

Notes

Supported by a grant from the Schering-Plough Research Institute

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “…randomised in a 1:1:1 ratio to 3 treatment arms…”</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comment: no further information. However, this is a large multinational RCT and...</td>
</tr>
<tr>
<td>Bias</td>
<td>Risk</td>
<td>Comment</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-------</td>
<td>-------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Comment: no information provided. However, this is a multinational trial with adequate blinding and should have adequate sequence generation and allocation concealment procedures</td>
</tr>
</tbody>
</table>
| Blinding of participants and personnel (performance bias) | Low risk | Quote: “double blind, double dummy”; “…matching placebo nasal spray …”  
Comment: pg 1276, col 1, para 1 and 2. “Matching placebo spray” mentioned and those on the 200 µg/day regimen were also given placebo nasal spray for the evening |
| Blinding of outcome assessment (detection bias) | Low risk | Comment: no information. Likely to remain well blinded until end of study |
| Incomplete outcome data (attrition bias) | High risk | Quote: 305/354 patients (86%) patients “completed 4-month treatment period”  
Comment: higher % of patients not completing in the placebo group 22/117 (19%)  
; compared to the twice daily or once daily groups 13/122 (11%) and 14/114 (12%) , respectively. Study mentioned analyses based on “all randomised subjects” using the “ITT principle” and endpoint was “defined as the last non-missing reading for the subject” for bilateral polyps score; however, unlikely all were analysed as numbers do not tally exactly with the ”meta-analysis subsequently reported” |
| Selective reporting (reporting bias) | Low risk | Comment: all outcomes reported in the methods section were reported in the results section |
| Other bias | Unclear risk | Comment: no information about the validation of outcome measures |
## Methods

3-arm, double-blind, multicentre, parallel-group RCT, with a 4-month duration of treatment and follow-up

## Participants

| Location | 24 centres in 17 countries worldwide |
| Setting  | study conducted from 25 June 2001 to 20 January 2003 |
| Sample size: | |
| Number randomised: | 102 in 400 µg, 102 in 200 µg, 106 in placebo group, respectively |
| Number completed: | 93 in 400 µg, 94 in 200 µg, 87 in placebo group, respectively |
| Participant (baseline) characteristics: | |
| Main diagnosis: | bilateral nasal polyps and clinically significant congestion/obstruction |
| Age (mean): | 400 µg: 47.6; 200 µg: 47.2; placebo: 50.9 |
| Gender (%M/%F): | 400 µg: 62/38; 200 µg: 70/30; placebo: 65/35 |
| Polyps status: | 100% with polyps |
| Previous sinus surgery status: | not more than 3 times or within past 6 months |
| Other important effect modifiers: | |
| o Asthma history (%): | 400 µg: 19; 200 µg: 15; placebo: 17 |
| o Perennial allergic rhinitis history (%): | 400 µg: 18; 200 µg: 14; placebo: 22 |

### Inclusion criteria:

≥ 18 years with an endoscopically confirmed diagnosis of bilateral nasal and clinically significant nasal congestion/obstruction (average morning score 2 or higher on an scale of 0 to 3 for each of the last 7 days of the 14-day run-in period). If had asthma, had a documented FEV₁ ≥ 80% of the predicted value within the 6 months before screening and no asthma exacerbations within 30 days before screening. Those treated with inhaled corticosteroids were required to be on a moderate, stable regimen of beclomethasone dipropionate ≤ 800 mg/d or equivalent for 1 month before screening and to remain on a stable regimen throughout the study period.

### Exclusion criteria:

- Seasonal allergic rhinitis within the past 2 years
- Sinus or nasal surgery within the previous 6 months or 3 nasal surgeries (or any surgical procedure preventing an accurate grading of polyps)
- Presumed fibrotic nasal polyposis, or complete or near complete nasal obstruction
- Nasal septal deviation requiring corrective surgery or nasal septal perforation
- Acute sinusitis, nasal infection or upper respiratory tract infection at screening or in the 2 weeks before screening
- Ongoing rhinitis medicamentosa
- Churg-Strauss syndrome
- Dyskinetic ciliary syndromes
- Cystic fibrosis
- Glaucoma or a history of posterior subcapsular cataracts
- Allergies to corticosteroids or aspirin, or any other clinically significant disease that would interfere with the evaluation of therapy

## Interventions

| 400 µg group (n = 102): | mometasone furoate nasal spray 200 µg twice daily (morning and evening) for 4 months |
| 200 µg group (n = 102): | mometasone furoate nasal spray 200 µg once daily (morning, matching placebo used in the evening) for 4 months |
| Placebo group (n = 106): | placebo nasal spray twice daily (morning and evening) for 4 months |

Use of additional interventions (common to both treatment arms):
Acetaminophen (paracetamol) was encouraged for analgesic purposes; NSAIDs limited to 5 consecutive days if alternative analgesia was required. Antibiotics were administered for bacterial infections at the discretion of the principal investigator. Concomitant medications that would interfere with study evaluations were not permitted, including nasal sodium cromolyn; nasal atropine or ipratropium bromide; corticosteroids (except oral inhaled corticosteroids for asthma or mild-strength or mid-strength topical corticosteroids for dermatologic purposes); antihistamines; decongestants; topical, oral or ocular anti-inflammatory drugs; or topical nasal or oral antifungal agents.

Outcomes

**Outcomes of interest in the review:**

**Primary outcomes**

1. Disease severity, patient evaluation of symptoms (congestion/obstruction, loss of sense of smell, anterior rhinorrhea and postnasal drip) measured daily on a diary card on a 4-point scale (0 = none, 3 = severe)
2. Significant adverse effect: epistaxis
3. Other adverse effects: local irritation
4. Polyps grade; bilateral score and proportion of patients demonstrating an improvement at endpoint

**Secondary outcomes:**

- Other outcomes reported by the study:
  - Therapeutic response (rated by investigator)
  - Peak nasal inspiratory flow
  - Treatment compliance
  - Number of withdrawals due to AE and events occurring in more than 2% of participants in any group

Notes

Supported by a grant from the Schering-Plough Research Institute

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “Randomization was performed in blocks of 3 using random numbers generated by SAS function UNIFORM (SAS Institute, Cary, NC) with seed based on clock time. Randomization was stratified by the presence or absence of concurrent asthma.” Comment: computerised randomisation</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Comment: although randomisation was blocked, blinding should be adequate</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Quote: “double blind”; “…matching placebo nasal spray …” Comment: “Matching placebo spray” mentioned; dosing regimen the same across all groups</td>
</tr>
</tbody>
</table>
### Stjarne 2006 (Continued)

<table>
<thead>
<tr>
<th>Bias</th>
<th>Risk Level</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td>Low risk</td>
<td>Comment: no information. Likely to remain well blinded until end of study</td>
</tr>
<tr>
<td><strong>Incomplete outcome data (attrition bias)</strong></td>
<td>High risk</td>
<td>Quote: &quot;More than 85% of subjects completed the 4-month treatment period, with more than twice as many placebo recipients as active drug recipients discontinuing during the treatment phase (18% vs 8%).&quot; Comment: drop-out rates not balanced</td>
</tr>
<tr>
<td><strong>Selective reporting (reporting bias)</strong></td>
<td>Unclear risk</td>
<td>Comment: although all outcomes mentioned in the methods were reported, these were mostly not in sufficient detail (e.g. only P values)</td>
</tr>
<tr>
<td><strong>Other bias</strong></td>
<td>Unclear risk</td>
<td>Comment: no information about the validation of outcome measures</td>
</tr>
</tbody>
</table>

### Stjarne 2006a

**Methods**

Double-blind, multicentre, parallel-group RCT, with a 4-month duration of treatment and follow-up

**Participants**

**Location:** 12 centres in Denmark, Finland, Norway and Sweden  
**Setting:** outpatient clinics  
**Sample size:**  
**Number randomised:** 153 in 200 µg, 145 in placebo group, respectively (298)  
**Number completed:** 134 in 200 µg, 101 in placebo group, respectively (235)  
**Participant (baseline) characteristics:**  
Main diagnosis: bilateral nasal polyps and clinically significant congestion/obstruction  
Age (mean): 53 (range 20 to 86)  
Gender (%M/%F): 200 µg group: 74.5/25.5; placebo group: 71.7/28.3  
Polyps status: 100% with polyps  
Previous sinus surgery status: > 2 surgeries, 25.5% in 200 µg group 26.2% in placebo group  
Other important effect modifiers:  
**Inclusion criteria:**  
Age ≥ 18 years; a diagnosis of bilateral nasal polyps and clinically significant nasal congestion. Nasal congestion was defined as significant when the symptom score was ≥ 2 (on a scale of 0 to 3) for ≥ 4 days per week during the month before screening, at screening and at the baseline visit  
**Exclusion criteria:**  
Nasal polyp surgery within the 6 months before screening; unhealed nasal surgery or trauma; polyp size of 3 (on a scale of 0 to 3); the presence of polyps that could interfere with nasal spray application; significant nasal structural abnormalities; ongoing concurrent nasal infections; glaucoma with narrow anterior chamber angle of the eye; rhinitis medicamentosa; or hereditary mucociliary dysfunction
Interventions

**200 µg group (n = 153):** mometasone furoate nasal spray 200 µg once daily (morning, matching placebo used in the evening) for 16 weeks

**Placebo group (n = 145):** placebo nasal spray twice daily (morning and evening) for 16 weeks

Concomitant medications that would interfere with study evaluations were not permitted, including nasal sodium cromolyn; nasal atropine or ipratropium bromide; corticosteroids (except oral inhaled corticosteroids for asthma or mild-strength or mid-strength topical corticosteroids for dermatologic purposes); antihistamines; decongestants; topical, oral or ocular anti-inflammatory drugs; or topical nasal or oral antifungal agents.

Outcomes

**Outcomes of interest in the review:**

*Primary outcomes*
1. Disease severity - participants with improvement in score
2. Significant adverse effect: epistaxis

*Secondary outcomes:*
3. Other adverse effects: local irritation

*Other outcomes reported by the study:*
- Investigator-evaluated nasal congestion, sense of smell and rhinorrhea
- PNIF
- Treatment response score
- Olfactory threshold

Notes

Participants who met the eligibility criteria at the screening visit (visit 1) underwent a 2-to 4-week no treatment run-in period. Criteria to remain in study/numbers subsequently excluded not reported.

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “...randomized at the baseline visit (day 0, visit 2) according to a computer-generated code...”</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Comment: adequate randomisation</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Quote: “The randomization schedule for the blinded treatments was maintained by the sponsor and only disclosed after the study was completed and the database closed.”</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td>Comment: adequate allocation concealment</td>
</tr>
</tbody>
</table>
**Stjarne 2006a** *(Continued)*

<table>
<thead>
<tr>
<th>Source of bias and type of bias</th>
<th>Risk of bias</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>As key outcomes were patient-reported and there was adequate blinding, it is likely there was adequate blinding for outcome assessment</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Quote: “Of the 298 subjects randomized to treatment, 235 (78.9%) completed the study. Premature withdrawals were more common in the placebo group than in the MFNS group (30.3% vs 12.4% of subjects, respectively).” Comment: drop-out rates not balanced across groups</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Comment: all outcomes mentioned in the methods section were reported in the results section, although these were mostly not in sufficient detail</td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>Quote: “Approximately 10% of subjects were considered to be noncompliant with the dosing regimen (defined as missing study medication doses for ≥7 consecutive days up to a maximum of 10 days, using rescue medication for ≥10 days during treatment, or using prohibited concomitant medications).” Comment: there was a 2 to 4-week run-in period - criteria to remain in study not reported There was no information about the validation of instruments used to measure symptoms</td>
</tr>
</tbody>
</table>

**Vlckova 2009**

- **Methods**: Double-blind, placebo-controlled, multicentre, parallel-group RCT, with a 12-week duration of treatment and a 14-week duration of follow-up
- **Participants**
  - **Location**: Czech Republic, 5 sites
  - **Setting of recruitment and treatment**: 5 otorhinolaryngology hospital clinics
  - **Sample size**:
    - **Number randomised**: 54 in intervention, 55 in comparison
    - **Number completed**: 54 in intervention, 52 in comparison
  - **Participant (baseline) characteristics**:
    - Age, mean (range): FP: 48.9 (18 to 65), PL: 47.0 (23 to 63)
    - Gender (M/F): FP: 74/26; PL: 62/38
Main diagnosis: mild to moderate bilateral nasal polyposis
Polyps status: all; about 50% in each arm had mild, 50% had moderate grade polyps
Previous sinus surgery status: never had surgery FP: 23 (43%), PL: 15 (27%); had at least 4 surgeries FP: 4 (7%), PL: 7 (13%)
Previous courses of steroids: no information
Other important effect modifiers, if applicable (e.g. aspirin sensitivity, comorbidities of asthma): asthma: FP: 17%; PL: 18%

Inclusion criteria:
Age 18 to 65 years, a diagnosis of bilateral nasal polyposis graded as mild or moderate
Verified airflow through both nostrils, an ability to close the soft palate and the ability to trigger the breath-actuation mechanism of a device in accordance with the instructions for use

Exclusion criteria:
Large polyps (grade 3)
Nasal polypl surgery during the 3 months before screening
Cystic fibrosis, a purulent nasal infection, allergic rhinitis or other disease likely to interfere with the study parameters
Depot or oral steroids during the previous 3 months
Cleft palate
Concomitant medications that would interfere with study evaluations were not permitted

Interventions

Intervention (n = 54): fluticasone propionate 800 µg/day delivered with breath actuated inhaler twice a day
Comparator group (n = 55): placebo, twice a day

Use of additional interventions (common to both treatment arms):
Saline rinsing and devices that dilate the nostrils were prohibited
Loratadine 10 mg tablets were provided as rescue medication for the relief of troublesome acute allergic symptoms. If a participant experienced a severe acute nasal blockage, the investigator could authorise the use of a short course of oxymetazoline drops or spray for a maximum of 7 consecutive days and a total maximum of 10 days during the treatment period. Oxymetazoline was not to be used within 24 hours of a scheduled study visit

Outcomes

Outcomes of interest in the review:

Primary outcomes:
1. Disease severity symptom score, using both a global rating scale (very much improved; improved; same; worse or very much worse). Also used a diary with the following scoring system: 0 (none), 1 (mild - symptoms present but not troublesome), 2 (moderate - symptoms frequently troublesome but not interfering with daily activity or night time sleep) or 3 (symptoms troublesome and interfering with daily activity or night-time sleep) for nasal blockage, nasal discomfort (facial and sinus pain and pressure) and rhinitis (nasal secretion, itching, irritation and sneezing). Sense of smell was rated as follows: 0 (normal), 1 (slightly impaired), 2 (moderately impaired) or 3 (absent)
2. Significant adverse effect: epistaxis

Secondary outcomes:
3. Endoscopy: polyp size was graded for each nostril using the Lildholdt scale. Some authors classify polyps causing total obstruction as grade 4. The score was presented for each nostril, the worst affected nostril and the summed score for both nostrils
4. Adverse event: local irritation
5. CT scan
Other outcomes reported by the study:
- PNIF
- Use of rescue medications

Study had a 14- to 16-day treatment-free run-in
Compliance was high with 98.92% of administrations made in the Opt-FP treatment group and 99.05% made in the placebo group

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Quote: “…randomized in a 1:1 ratio” Comment: no further description of randomisation methods</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Comment: no information about allocation concealment</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>Quote: “…Opt-FP and placebo breath-actuated bi-directional delivery devices were identical in appearance…. The placebo aqueous nasal spray was formulated to match FP exactly, except for the active ingredient… To deliver a dose of FP 400 µg b.i.d. or matching placebo, the subjects made two administrations to each nostril in the morning and the evening…” Comment: identical-looking devices and same frequency of administration. Same formulation, except the omission of the active ingredient. Should be able to maintain good blinding</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Low risk</td>
<td>Quote: as above Comment: most are patient-reported outcomes therefore blinding should be adequate</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>Low risk</td>
<td>Quote: “A total of 106 subjects (97%) completed the study. Three subjects withdrew, all in the placebo group (one due to worsening of polyps, two withdrew consent).” Comment: low drop-out rate</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Comment: no protocol was available. Most outcomes are reported in graphs. Global improvement score dichotomised when reported (not pre-specified)</td>
</tr>
</tbody>
</table>
Other bias | Unclear risk | Comment: no mention of validation of questionnaires
---|---|---

Zhou 2015

Methods | 2-arm, double-blind, multicentre, parallel-group RCT, with a 16-week duration of treatment and follow-up
---|---

Participants | Location: China, 28 sites  
Setting of recruitment and treatment: not clear  
Sample size: 748  
Number randomised: 375 in intervention, 373 in comparison  
Number completed: 350 in intervention, 336 in comparison  
Participant (baseline) characteristics:  
- Mean age ± SD: intervention: 46.8 ± 13.5; control: 46.8 ± 13.8  
- Gender male (%): intervention: 224 (59.7%); control: 239 (64.1%)  
- Main diagnosis: Chinese patients with bilateral nasal polyps  
- Polyps status: 100% with polyps  
- Baseline total polyp size: intervention 3.6 ± 1.1; control 3.7 ± 1.1  
- % previous sinus surgery status: intervention: 24.5%; 22.8%  
- Other important effect modifiers:  
  - % concurrent asthma status: intervention: 2.7%; control: 3.5%  
  - % sensitivity to antigen: intervention: 4.3%; control: 5.1%

Inclusion criteria:  
- ≥ 18 years of age, Chinese  
- diagnosis of bilateral nasal polyps (graded on a scale of 0 to 3, see below)  
- clinically significant nasal congestion/obstruction (average morning score ≥ 2 for each of the last 7 days of the 14-day run-in period)  
- Patients with asthma were included if FEV₁ ≥ 80% within 6 months and no exacerbations within 30 days of screening. Using inhaled corticosteroids, on a moderate, stable regimen of beclomethasone dipropionate 800 mg/day or equivalent for 1 month before screening and remained on a stable regimen throughout the study.  
- Nasal polyp score: polyps were graded by size and extent in both the left and right nasal fossa on a scale of 0 to 3 (0 = no polyps; 1 = polyp in middle meatus, not reaching below the inferior border of the middle turbinate; 2 = polyp reaching below the inferior border of the middle turbinate but not the inferior border of the inferior turbinate; and 3 = large polyp reaching to or below the lower border of the inferior turbinate or polyps medial to the middle turbinate). The sum of left and right nasal fossa polyp scores gave the total bilateral polyps grade.

Exclusion criteria:  
- History of seasonal allergic rhinitis within the past 2 years  
- Sinus or nasal surgery within the previous 6 months  
- History of 3 or more nasal surgeries in the past  
- History of any procedure preventing an accurate grading of polyps  
- Presumed fibrotic nasal polyps, or complete/near-complete nasal obstruction  
- Nasal septal deviation requiring corrective surgery; nasal septal perforation  
- Acute sinusitis, nasal infection or upper respiratory tract/nasal infection at (or
within 2 weeks of) screening
- Ongoing rhinitis medicamentosa, Churg-Strauss syndrome, dyskinetic ciliary syndromes, cystic fibrosis
- Glaucoma or a history of posterior subcapsular cataracts; allergies to corticosteroids or aspirin
- Any other clinically significant disease that would interfere with the evaluation of therapy

Interventions

**Intervention (n = 375):** mometasone nasal spray, 400 µg per day (200 µg twice daily), 16 weeks

**Comparator group (n = 373):** matching placebo nasal spray, twice daily for 16 weeks

**Use of additional interventions:**
The following were prohibited: nasal sodium cromolyn; nasal atropine or ipratropium bromide; guaifenesin; oral/intramuscular/intranasal corticosteroids (except oral inhaled corticosteroids for asthma or mild-strength topical corticosteroids for dermatologic purposes); antihistamines; decongestants; topical, oral or ocular antiinflammatory drugs; nonsteroidal anti-inflammatory drugs; or topical nasal or oral antifungal agents

Outcomes

**Outcomes of interest in the review:**

**Primary outcomes:**
1. Disease severity symptom score; patients reported individual symptom scores at 16 weeks, measured on a scale of 0 to 3 (0 = none; 1 = mild; 2 = moderate; 3 = severe) for the following symptoms: nasal obstruction, anterior rhinorrhea, post-nasal drip, loss of sense of smell
2. Significant adverse effect: epistaxis

**Secondary outcomes:**
3. Endoscopy (nasal polyps size); measured at 16 weeks in both left and right nasal fossa on a scale of 0 to 3 (see inclusion criteria for details) and summed for each nasal fossa to give a total score
4. Adverse effects: local irritation

Other outcomes reported by the study:
- Therapeutic response on a qualitative scale ranging from “complete relief of symptoms” to “no relief”
- Compliance defined as 29% to 138% of reference study drug bottle weight

Notes

All patients had a 14-day, single-blind placebo run-in period where signs and symptoms of nasal polyps were evaluated
The compliance rate was similar; 95.5% (MFNS) and 94.8% (placebo)

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: “At baseline, patients were randomised (1:1 ratio) according to a computer-generated allocation schedule” Comment: adequate randomisation method</td>
</tr>
<tr>
<td>Domain</td>
<td>Bias Type</td>
<td>Risk</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>-----------------------</td>
<td>--------</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td></td>
</tr>
</tbody>
</table>

AE: adverse event  
ASA: acetylsalicylic acid  
BDP: beclomethasone dipropionate  
CRS: chronic rhinosinusitis  
CT: computed tomography  
d: day  
ENT: ear, nose and throat  
F: female
**Characteristics of excluded studies** [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALA 2015</td>
<td>POPULATION: children and adults with uncontrolled asthma and CRS. The population did not meet the EPOS definition of CRS</td>
</tr>
<tr>
<td>Albu 2010</td>
<td>DESIGN: perioperative treatment using topical steroids for improvement of surgical outcomes in patients undergoing surgery</td>
</tr>
<tr>
<td>Bross-Soriano 2004</td>
<td>POPULATION: all patients underwent endoscopic polypectomy at the start of the trial</td>
</tr>
<tr>
<td>Cassano 1996</td>
<td>POPULATION: all patients had surgery at the start of the trial</td>
</tr>
<tr>
<td>Chalton 1985</td>
<td>DURATION: treatment and follow-up only 4 weeks (betamethasone drops)</td>
</tr>
<tr>
<td>Cuenant 1986</td>
<td>POPULATION: chronic allergic or bacterial sinusitis</td>
</tr>
<tr>
<td>Dijkstra 2004</td>
<td>POPULATION: treatment started 1 week after FESS (continued for 1 year)</td>
</tr>
<tr>
<td>Drettner 1982</td>
<td>POPULATION: treatment started 4 weeks after polypectomy surgery (continued for 3 months)</td>
</tr>
<tr>
<td>Ehnhage 2009</td>
<td>DESIGN: perioperative study; patients received intranasal corticosteroids/placebo before FESS, and then carried on intranasal corticosteroids/placebo</td>
</tr>
<tr>
<td>el Naggar 1995</td>
<td>STUDY DESIGN: treatment started immediately after intranasal polypectomy; randomised by side of nose</td>
</tr>
<tr>
<td>Filiaci 2000</td>
<td>STUDY DESIGN: treatment and follow-up only 8 weeks (budesonide)</td>
</tr>
<tr>
<td>Furukido 2005</td>
<td>INTERVENTION: YAMIK sinus catheter for 4 weeks</td>
</tr>
<tr>
<td>Study</td>
<td>POPULATION</td>
</tr>
<tr>
<td>------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Gulati 2001</td>
<td>POPULATION: treatment started 1 week after polypectomy surgery (continued for 3 months)</td>
</tr>
<tr>
<td>Hartwig 1988</td>
<td>POPULATION: all patients had polypectomy and treatment started the day after surgery</td>
</tr>
<tr>
<td>Jankowski 2001</td>
<td>DURATION: treatment only 8 weeks (budesonide)</td>
</tr>
<tr>
<td>Jankowski 2009</td>
<td>DURATION: there was an intranasal corticosteroids (fluticasone) versus placebo comparison only for 8 weeks, subsequently all patients had intranasal corticosteroids (for 6 months)</td>
</tr>
<tr>
<td>Johansson 2002</td>
<td>DURATION: treatment and follow-up only 2 weeks</td>
</tr>
<tr>
<td>Jorissen 2009</td>
<td>POPULATION: all participants had FESS at the start of the trial</td>
</tr>
<tr>
<td>Jurkiewicz 2004</td>
<td>POPULATION: started treatment after endoscopic polypectomy</td>
</tr>
<tr>
<td>Kang 2008</td>
<td>POPULATION: study started directly after FESS</td>
</tr>
<tr>
<td>Kapucu 2012</td>
<td>DURATION: follow-up only 1 month (triamcinolone intra-polyp steroid injections or nasal spray versus placebo)</td>
</tr>
<tr>
<td>Karlsson 1982</td>
<td>POPULATION: included patients immediately after polypectomy</td>
</tr>
<tr>
<td>Keith 1995</td>
<td>DURATION: treatment and follow-up only 4 weeks (budesonide)</td>
</tr>
<tr>
<td>Lavigne 2002</td>
<td>INTERVENTION: sinus irrigation with steroids using MAST tube, for 3 weeks</td>
</tr>
<tr>
<td>Lildholdt 1995</td>
<td>DURATION: treatment and follow-up only 4 weeks</td>
</tr>
<tr>
<td>Malmberg 1988</td>
<td>POPULATION: all patients had polypectomy within 2 months of the start of the trial</td>
</tr>
<tr>
<td>Man 2013</td>
<td>DURATION: treatment and follow-up was for just 6 weeks. The intervention used was 3 mg of fluticasone in 240 ml of normal saline</td>
</tr>
<tr>
<td>Mastalerz 1997</td>
<td>POPULATION: aspirin-induced asthma and chronic eosinophilic rhinitis</td>
</tr>
<tr>
<td>Meltzer 1993</td>
<td>DURATION: treatment and follow-up only 7 weeks; patients with maxillary sinusitis had flunisolide spray versus placebo added to amoxicillin/clavulanate</td>
</tr>
<tr>
<td>Mygind 1975</td>
<td>DURATION: treatment and follow-up only 3 weeks</td>
</tr>
<tr>
<td>Optinose 2012</td>
<td>OTHER: RCT prematurely ended</td>
</tr>
<tr>
<td>Passali 2003</td>
<td>POPULATION: patients started treatment 1 month after surgery</td>
</tr>
<tr>
<td>Qvarnberg 1992</td>
<td>POPULATION: did not meet current CRS definitions - “recurrent or chronic maxillary sinusitis” (no nasal polyps)</td>
</tr>
</tbody>
</table>
Continued

<table>
<thead>
<tr>
<th>Study Reference</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rotenberg 2011</td>
<td>POPULATION: all patients underwent endoscopic sinus surgery at the start of the trial</td>
</tr>
<tr>
<td>Rowe-Jones 2005</td>
<td>POPULATION: all patients underwent endoscopic sinus surgery at the start of the trial</td>
</tr>
<tr>
<td>Ruhno 1990</td>
<td>DURATION: treatment and follow-up only 4 weeks</td>
</tr>
<tr>
<td>Saunders 1999</td>
<td>DESIGN: perioperative study: patients were given intranasal corticosteroids for 2 weeks before polypectomy</td>
</tr>
<tr>
<td>Slifirski 2009</td>
<td>POPULATION: all patients underwent surgery at the start of the trial before randomisation</td>
</tr>
<tr>
<td>Stjarne 2009</td>
<td>POPULATION: all patients underwent FESS at the start of the trial</td>
</tr>
<tr>
<td>Taub 1968</td>
<td>DESIGN: cross-over study. 4 weeks of treatment followed by dexamethasone spray/placebo, then switched. Unclear if there was a washout period. Results not presented separately</td>
</tr>
<tr>
<td>Toft 1982</td>
<td>DURATION: treatment only 2 weeks (beclomethasone). A cross-over study with 1-week follow-up</td>
</tr>
<tr>
<td>Tos 1998</td>
<td>DURATION: treatment and follow-up only 6 weeks (budesonide)</td>
</tr>
<tr>
<td>Vento 2012</td>
<td>POPULATION: all patients had surgery around 2 weeks before the treatment started</td>
</tr>
<tr>
<td>Virolainen 1980</td>
<td>POPULATION: all patients underwent radical ethmoidectomy surgery 2 days prior to treatment starting</td>
</tr>
<tr>
<td>Wang 2015</td>
<td>DURATION: treatment and follow-up only 14 days (budesonide nebulisation)</td>
</tr>
</tbody>
</table>

CRS: chronic rhinosinusitis  
EPOS: European Position Paper on Rhinosinusitis and Nasal Polyps 2012  
FESS: functional endoscopic sinus surgery  
RCT: randomised controlled trial

**Characteristics of studies awaiting assessment**  [ordered by study ID]

**Bachert 2004**

<table>
<thead>
<tr>
<th>Details</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
<td>-</td>
</tr>
<tr>
<td>Participants</td>
<td>-</td>
</tr>
<tr>
<td>Interventions</td>
<td>-</td>
</tr>
<tr>
<td>Outcomes</td>
<td>-</td>
</tr>
<tr>
<td>Notes</td>
<td>Conference proceeding: we cannot locate the abstract</td>
</tr>
<tr>
<td>Study</td>
<td>Methods</td>
</tr>
<tr>
<td>------------------</td>
<td>---------</td>
</tr>
<tr>
<td>Meln 2004</td>
<td>-</td>
</tr>
<tr>
<td>Pisano 2000</td>
<td>-</td>
</tr>
<tr>
<td>Riem 2005</td>
<td>-</td>
</tr>
<tr>
<td>Ygind 1996</td>
<td>-</td>
</tr>
</tbody>
</table>

Notes: We cannot locate the abstract for Pisano 2000, Riem 2005, and Ygind 1996.
### Characteristics of ongoing studies  *(ordered by study ID)*

#### NCT01013701

<table>
<thead>
<tr>
<th>Trial name or title</th>
<th>'Compare the effects of fluticasone furoate nasal spray vs placebo in patients with nasal polyoid disease'</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
<td>Double-blind, parallel assignment, randomised controlled trial</td>
</tr>
<tr>
<td>Participants</td>
<td>Adults with nasal polyps</td>
</tr>
</tbody>
</table>
| Interventions       | - Fluticasone furoate  
                      - Placebo                                                                                           |
| Outcomes            | To evaluate the effect of once daily nasal steroid therapy with fluticasone furoate nasal spray (110 µg/day) in suppressing nasal polyp-induced symptoms over the course of 16 weeks in patients presenting to the clinic with active nasal polyoid disease |
| Starting date       | 2009                                                                                                   |
| Contact information | Peter S. Creticos MD, Johns Hopkins University                                                          |
| Notes               | ClinicalTrials.gov website indicates that the recruitment status of this study is unknown because the information has not been verified recently. We tried to contact the study authors to find out more information but we did not receive a response |

#### NCT01622569

<table>
<thead>
<tr>
<th>Trial name or title</th>
<th>'Study evaluating the efficacy and safety of intranasal administration of 100, 200, and 400 µg of fluticasone propionate twice a day (bid) using a novel bi directional device in subjects with bilateral nasal polyposis followed by an 8-week open-label extension phase to assess safety'</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
<td>Double-blind, parallel assignment, randomised controlled trial</td>
</tr>
<tr>
<td>Participants</td>
<td>Adults with bilateral nasal polyposis</td>
</tr>
</tbody>
</table>
| Interventions       | - Fluticasone propionate 100 µg twice a day  
                      - Fluticasone propionate 200 µg twice a day  
                      - Fluticasone propionate 400 µg twice a day  
                      - Matching placebo  
                      For 16 weeks                                                                |
| Outcomes            | - Reduction of nasal congestion/obstruction symptoms  
                      - Reduction in total polyp grade (sum of scores from both nasal cavities)  
                      No secondary outcomes were listed in the trial registry entry                                                      |
| Starting date       | 2013                                                                                                               |
| Contact information | Optinose US Inc. No further details provided.                                                                      |
### NCT01622569 (Continued)

| Notes | Study has been listed as completed on the registry website (October 2015). No results are currently available |

### NCT01624662

| Trial name or title | 'Efficacy and safety study of intranasal administration of 100, 200, and 400 µg of fluticasone propionate twice a day (bid) using a novel bi directional device in subjects with bilateral nasal polyposis followed by an 8-week open-label extension phase to assess safety’ |
| Methods | Double-blind, parallel assignment, randomised controlled trial |
| Participants | Adults with bilateral nasal polyposis |
| Interventions | - Fluticasone propionate 100 µg twice a day  
- Fluticasone propionate 200 µg twice a day  
- Fluticasone propionate 400 µg twice a day  
- Matching placebo  
For 16 weeks |
| Outcomes | - Reduction of nasal congestion/obstruction symptoms  
- Reduction in total polyp grade (sum of scores from both nasal cavities)  
No secondary outcomes were listed in the trial registry entry |
| Starting date | 2012 |
| Contact information | Optinose US Inc. No further details provided. |
| Notes | Study has been listed as completed on the registry website (October 2015). No results are currently available |
### Comparison 1. Intranasal corticosteroids versus placebo

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Disease severity - measured as change from baseline using the Chronic Sinusitis Survey at 20 weeks</td>
<td>1</td>
<td>134</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>2.84 [-5.02, 10.70]</td>
</tr>
<tr>
<td>2 Disease severity - global symptom score, measured as proportion of patients who improved</td>
<td>1</td>
<td>109</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>2.78 [1.76, 4.40]</td>
</tr>
<tr>
<td>3 Disease severity - combination of individual symptom scores, measured on a 0 to 3 scale as change from baseline at 12 to 20 weeks</td>
<td>6</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>3.1 Average symptom score (4 EPOS domains)</td>
<td>2</td>
<td>243</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.26 [-0.37, -0.15]</td>
</tr>
<tr>
<td>3.2 Average symptom score (3 EPOS domains - nasal blockage, rhinorrhoea, loss of sense of smell)</td>
<td>4</td>
<td>1345</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.31 [-0.38, -0.23]</td>
</tr>
<tr>
<td>3.3 Average symptom score (2 EPOS domains - nasal blockage and rhinorrhoea)</td>
<td>6</td>
<td>1702</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.31 [-0.38, -0.24]</td>
</tr>
<tr>
<td>4 Disease severity - individual symptoms, measured as average change from baseline at 12 to 20 weeks (range 0 to 3 points)</td>
<td>6</td>
<td></td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>4.1 Nasal blockage</td>
<td>6</td>
<td>1702</td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>-0.40 [-0.52, -0.29]</td>
</tr>
<tr>
<td>4.2 Rhinorrhoea</td>
<td>6</td>
<td>1702</td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>-0.25 [-0.33, -0.17]</td>
</tr>
<tr>
<td>4.3 Loss of sense of smell</td>
<td>4</td>
<td>1345</td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>-0.19 [-0.28, -0.11]</td>
</tr>
<tr>
<td>4.4 Facial pain/pressure</td>
<td>2</td>
<td>243</td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>-0.27 [-0.56, 0.02]</td>
</tr>
<tr>
<td>5 Adverse events - epistaxis</td>
<td>13</td>
<td>2508</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>2.74 [1.88, 4.00]</td>
</tr>
<tr>
<td>5.1 With nasal polyps</td>
<td>10</td>
<td>2262</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>3.03 [2.00, 4.59]</td>
</tr>
<tr>
<td>5.2 Without nasal polyps</td>
<td>3</td>
<td>246</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>1.49 [0.59, 3.78]</td>
</tr>
<tr>
<td>6 Adverse events - local irritation</td>
<td>11</td>
<td>2124</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>0.94 [0.53, 1.64]</td>
</tr>
<tr>
<td>6.1 With nasal polyps</td>
<td>9</td>
<td>2045</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>0.93 [0.52, 1.67]</td>
</tr>
<tr>
<td>6.2 Without nasal polyps</td>
<td>2</td>
<td>79</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>1.02 [0.15, 6.84]</td>
</tr>
<tr>
<td>7 Endoscopy score (bilateral polyps score) measured change from baseline (range 0 to 6 points)</td>
<td>4</td>
<td>1417</td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>-0.58 [-0.90, -0.26]</td>
</tr>
<tr>
<td>7.1 With nasal polyps</td>
<td>4</td>
<td>1417</td>
<td>Mean Difference (IV, Random, 95% CI)</td>
<td>-0.58 [-0.90, -0.26]</td>
</tr>
<tr>
<td>8 Endoscopy score (polyps size) - measured as numbers with improvement at longest available follow-up</td>
<td>5</td>
<td>676</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.77 [1.06, 2.95]</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>8.1 With nasal polyps</td>
<td>5</td>
<td>676</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.77 [1.06, 2.95]</td>
</tr>
<tr>
<td>9 Endoscopy score (polyp size) - measured as numbers with improvement at longest available follow-up</td>
<td>8</td>
<td>1984</td>
<td>Odds Ratio (Random, 95% CI)</td>
<td>2.07 [1.48, 2.91]</td>
</tr>
<tr>
<td>9.1 With nasal polyps</td>
<td>4</td>
<td>1417</td>
<td>Odds Ratio (Random, 95% CI)</td>
<td>2.43 [1.46, 4.06]</td>
</tr>
<tr>
<td>9.2 With nasal polyps from dichotomous data</td>
<td>4</td>
<td>567</td>
<td>Odds Ratio (Random, 95% CI)</td>
<td>1.73 [1.19, 2.51]</td>
</tr>
<tr>
<td>10 CT score (overall) - measured using Lund-Mackay score (max 24 points) at 3 months</td>
<td>1</td>
<td>47</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-4.82 [-7.27, -2.37]</td>
</tr>
</tbody>
</table>

**Contributions of Authors**

Lee Yee Chong: scoped, designed and wrote the protocol, screened abstracts, extracted data, conducted the analysis and wrote up the review.

Karen Head: reviewed and edited the protocol, screened abstracts, extracted data, helped to check the analysis and contributed to the writing of the review.

Claire Hopkins: clinical guidance at all stages of project scoping and protocol development, full text screening, data analysis and interpretation of the review. Commented on drafts of the review.

Carl Philpott: clinical guidance at all stages of project scoping and protocol development, full text screening, data analysis and interpretation of the review. Contributed to the writing of the review.

Anne GM Schilder: commented on drafts and contributed to the writing of the review.

Martin J Burton: helped to draft the protocol; clinical guidance at all stages of project scoping and protocol development and contributed to the writing of the review.

**Declarations of Interest**

Lee Yee Chong: none known.

Karen Head: none known.

Claire Hopkins: I have received financial support from several companies involved in producing instruments for sinus surgery: Acclarent, Sinusys, Cryolife and Medtronic.

Carl Philpott: I have previously received consultancy fees from the companies Acclarent, Navigant, Aerin Medical and Entellus.

Anne GM Schilder: Professor Anne Schilder is joint Co-ordinating Editor of Cochrane ENT, but had no role in the editorial process for this review. Her evidENT team at UCL is supported by her NIHR Research Professorship award with the remit to develop a UK infrastructure and programme of clinical research in ENT, Hearing and Balance. Her institution has received a grant from GSK for a study on the microbiology of acute tympanostomy tube otitis.
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Internal sources

- No sources of support supplied

External sources

- National Institute for Health Research, UK.
  Funding to complete a suite of reviews on medical interventions for chronic rhinosinusitis in 2015/2016 (award reference 14/174/03), in addition to infrastructure funding for Cochrane ENT

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

In the protocol, the example given for local irritation was “including sore throat, oral thrush”. This has been expanded to include “and other local nasal irritation such as dryness, itchiness etc.”

Although we had planned to present data for chronic rhinosinusitis without nasal polyps and chronic rhinosinusitis with nasal polyps in subgroups as a visual comparison, this was not carried out except for the nasal polyps and adverse events outcomes. For all the other outcomes there was no more than one study of patients with chronic rhinosinusitis without nasal polyps available for analysis. We had footnoted this in the forest plots and highlighted it in the write up whenever this occurred.

As part of the discussions about the use of a total symptom score we noted that many papers within the suite of reviews did not present information for all four elements of the EPOS criteria for defining chronic rhinosinusitis (EPOS 2012). In particular, many studies that only included patients with nasal polyps did not present information on facial pressure or pain. We made the decision that where individual symptoms were recorded, they should be presented within the outcome of disease severity symptom score within the paper as this information would be useful for the reader.

INDEX TERMS

Medical Subject Headings (MeSH)

*Quality of Life; Administration, Intranasal; Adrenal Cortex Hormones [administration & dosage; adverse effects]; Beclomethasone [administration & dosage; adverse effects]; Budesonide [administration & dosage; adverse effects]; Chronic Disease; Fluticasone [administration & dosage; adverse effects]; Mometasone Furoate [administration & dosage; adverse effects]; Nasal Polyps [drug therapy]; Nasal Sprays; Placebos [administration & dosage]; Randomized Controlled Trials as Topic; Rhinitis [*drug therapy]; Severity of Illness Index; Sinusitis [*drug therapy]; Steroids [*administration & dosage; adverse effects]

MeSH check words

Adolescent; Adult; Child; Humans