Inhaled corticosteroids for subacute and chronic cough in adults (Review)

Johnstone KJ, Chang AB, Fong KM, Bowman RV, Yang IA

This is a reprint of a Cochrane review, prepared and maintained by The Cochrane Collaboration and published in The Cochrane Library 2013, Issue 3

http://www.thecochranelibrary.com

WILEY
# Table of Contents

- **HEADER** ........................................................................................................... 1  
- **ABSTRACT** ........................................................................................................ 1  
- **PLAIN LANGUAGE SUMMARY** ................................................................. 2  
- **SUMMARY OF FINDINGS FOR THE MAIN COMPARISON** ....................... 4  
- **BACKGROUND** ............................................................................................... 7  
- **OBJECTIVES** .................................................................................................... 7  
- **METHODS** ...................................................................................................... 8  
- **RESULTS** .......................................................................................................... 10  
  - Figure 1. .............................................................................................................. 11  
  - Figure 2. .............................................................................................................. 12  
  - Figure 3. .............................................................................................................. 15  
  - Figure 4. .............................................................................................................. 19  
  - Figure 5. .............................................................................................................. 20  
- **DISCUSSION** ................................................................................................. 21  
- **AUTHORS’ CONCLUSIONS** .......................................................................... 26  
- **ACKNOWLEDGEMENTS** .................................................................................. 27  
- **REFERENCES** ................................................................................................. 27  
- **CHARACTERISTICS OF STUDIES** .............................................................. 31  
- **DATA AND ANALYSES** .................................................................................. 48  
- **WHAT’S NEW** ................................................................................................. 51  
- **CONTRIBUTIONS OF AUTHORS** .................................................................. 52  
- **DECLARATIONS OF INTEREST** ................................................................... 52  
- **SOURCES OF SUPPORT** ............................................................................... 52  
- **DIFFERENCES BETWEEN PROTOCOL AND REVIEW** ............................... 53  
- **INDEX TERMS** ................................................................................................ 53
Inhaled corticosteroids for subacute and chronic cough in adults

Kate J Johnstone¹, Anne B Chang²,³,⁴, Kwun M Fong¹,⁵, Rayleen V Bowman¹,⁵, Ian A Yang¹,⁵

¹School of Medicine, The University of Queensland, Brisbane, Australia. ²Menzies School of Health Research, Charles Darwin University, Casuarina, Australia. ³Queensland Children’s Respiratory Centre, Royal Children’s Hospital, Brisbane, Australia. ⁴Queensland Children’s Medical Research Institute, The University of Queensland, Brisbane, Australia. ⁵Thoracic Medicine Program, The Prince Charles Hospital, Brisbane, Australia

Contact address: Kate J Johnstone, School of Medicine, The University of Queensland, The Prince Charles Hospital, Rode Rd, Brisbane, Queensland, 4032, Australia. kate.johnstone@uqconnect.edu.au.

Editorial group: Cochrane Airways Group.
Publication status and date: Edited (no change to conclusions), published in Issue 5, 2013.
Review content assessed as up-to-date: 13 December 2012.

Citation: Johnstone KJ, Chang AB, Fong KM, Bowman RV, Yang IA. Inhaled corticosteroids for subacute and chronic cough in adults. Cochrane Database of Systematic Reviews 2013, Issue 3. Art. No.: CD009305. DOI: 10.1002/14651858.CD009305.pub2.

Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

ABSTRACT

Background
Persistent cough is a common clinical problem. Despite thorough investigation and empirical management, a considerable proportion of those people with subacute and chronic cough have unexplained cough, for which treatment options are limited. While current guidelines recommend inhaled corticosteroids (ICS), the research evidence for this intervention is conflicting.

Objectives
To assess the effects of ICS for subacute and chronic cough in adults.

Search methods
We searched the Cochrane Airways Group Register of Trials, Cochrane Central Register of Controlled Trials, MEDLINE, EMBASE and ClinicalTrials.gov in December 2012 and conducted handsearches.

Selection criteria
Two authors independently assessed all potentially relevant trials. All published and unpublished randomised comparisons of ICS versus placebo in adults with subacute or chronic cough were included. Participants with known chronic respiratory disease and asthma were excluded. Studies of cough-variant asthma and eosinophilic bronchitis were eligible.

Data collection and analysis
Two authors independently extracted data pertaining to pre-defined outcomes. The primary outcome was the proportion of participants with clinical cure or significant improvement (over 70% reduction in cough severity measure) at follow up (clinical success). The secondary outcomes included proportion of participants with clinical cure or over 50% reduction in cough severity measure at follow up, mean change in cough severity measures, complications of cough, biomarkers of inflammation and adverse effects. We requested additional data from study authors.
Main results
Eight primary studies, including 570 participants, were included. The overall methodological quality of studies was good. Significant clinical heterogeneity resulting from differences in participants and interventions, as well as variation in outcome measures, limited the validity of comparisons between studies for most outcomes. Data for the primary outcome of clinical cure or significant (> 70%) improvement were available for only three studies, which were too heterogeneous to pool. Similarly, heterogeneity in study characteristics limited the validity of meta-analysis for the secondary outcomes of proportion of participants with clinical cure or over 50% reduction in cough severity measure and clinical cure. One parallel group trial of chronic cough which identified a significant treatment effect contributed the majority of statistical heterogeneity for these outcomes. While ICS treatment resulted in a mean decrease in cough score of 0.34 standard deviations (SMD -0.34; 95% CI -0.56 to -0.13; 346 participants), the quality of evidence was low. Heterogeneity also prevented meta-analysis for the outcome of mean change in visual analogue scale score. Meta-analysis was not possible for the outcomes of pulmonary function, complications of cough or biomarkers of inflammation due to insufficient data. There was moderate quality evidence that treatment with ICS did not significantly increase the odds of experiencing an adverse event (OR 1.67; 95% CI 0.92 to 3.04).

Authors’ conclusions
The studies were highly heterogeneous and results were inconsistent. Heterogeneity in study design needs to be addressed in future research in order to test the efficacy of this intervention. International cough guidelines recommend that a trial of ICS should only be considered in patients after thorough evaluation including chest X-ray and consideration of spirometry and other appropriate investigations.

Plain Language Summary
Inhaled corticosteroids for adults with cough lasting over three weeks

Background
There is often no obvious cause for coughs that last more than three weeks. Lack of a clear cause makes the cough difficult to treat. Current guidelines recommend that in many cases people with cough lasting longer than three weeks be given inhaled corticosteroids (ICS), which are commonly used to treat asthma and other diseases involving airway inflammation.

Review question
We wanted to find out if taking inhaled steroids in adults with cough lasting three weeks or longer were beneficial.

We looked at evidence from clinical trials. We analysed the effects of ICS compared with placebo on cough severity, lung function, complications of cough and airway inflammation, as well as the safety of this treatment.

Study characteristics
We found eight studies on 570 people with cough lasting over three weeks. Studies included different types of participants in terms of age, duration of coughing and risk factors for cough. Studies also varied in types of ICS, doses, treatment lengths and types of inhaler used. Cough severity was measured using different scales.

Key results
We looked at the proportion of people who were clinically cured or showed a significant improvement in cough severity as our primary outcome, but the data were too mixed to be able draw any conclusions. These differences between studies also prevented meaningful pooling of study results for proportion of people showing improvement in cough and average improvement in one specific type of cough scale. There was low quality evidence that ICS reduced cough severity score. There was not enough data about changes in pulmonary function, complications of cough and markers of inflammation to allow pooling of results. There was evidence of moderate quality that ICS treatment did not increase the risk of adverse events.

Conclusion and future work
This review has shown that the effects of ICS for subacute and chronic cough are inconsistent. Further studies with more consistent patient populations, interventions, outcome measures and reporting are needed to determine whether ICS help subacute and chronic cough in adults.
This Cochrane plain language summary was written in December 2012.
### Summary of Findings for the Main Comparison

Inhaled corticosteroids (ICS) compared to placebo for adults with subacute and chronic cough

**Patient or population:** adults with subacute and chronic cough  
**Settings:** all  
**Intervention:** ICS  
**Comparison:** placebo  

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Illustrative comparative risks* (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No of Participants (studies)</th>
<th>Quality of the evidence (GRADE)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Assumed risk</td>
<td>Corresponding risk</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>ICS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Primary outcome

Proportion of participants who achieved clinical cure or significant improvement (≥70% reduction in cough severity measure) at follow up (clinical success)  
Symptomatic cough severity measure as assessed by the patient  
Follow up: 2 to 4 weeks

See comment  
See comment  
Not estimable  
180 (3 studies)  
⊕⊕⃝⃝  
low

Meta-analysis not appropriate; heterogeneity explained by differences in study design and outcomes

#### Secondary outcomes

Proportion of participants who achieved clinical cure or > 50% reduction in cough severity measure at follow up  
Symptomatic cough

See comment  
See comment  
Not estimable  
230 (4 studies)  
⊕⊕⃝⃝  
low

Meta-analysis not appropriate; heterogeneity explained by differences in study design and outcomes
<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Description</th>
<th>Participants</th>
<th>Mean Change in Cough Score</th>
<th>Proportion with Adverse Effects of Treatment</th>
<th>GRADE Working Group grades of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proportion of participants with clinical cure at follow up</td>
<td>See comment</td>
<td>Not estimable</td>
<td>SMD -0.34 (-0.56 to -0.13)</td>
<td>116 per 1000 (108 to 285) OR 1.67 (0.92 to 3.04)</td>
<td>Low2,4</td>
</tr>
<tr>
<td>Mean change in cough score</td>
<td>Symptomatic cough severity measure as assessed by the patient</td>
<td>320 (4 studies)</td>
<td>346 (5 studies)</td>
<td>381 (4 studies)</td>
<td>moderate7</td>
</tr>
</tbody>
</table>

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

CI: Confidence interval; OR: Odds ratio

GRADE Working Group grades of evidence
- **High quality**: Further research is very unlikely to change our confidence in the estimate of effect.
- **Moderate quality**: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- **Low quality**: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
- **Very low quality**: We are very uncertain about the estimate.

1. Unclear risk of selection bias (Boulet 1994); unclear risk of reporting bias (Ponsioen 2005; Ribeiro 2007).
2. Dichotomous outcome data based on less than 300 events.
4. Unclear risk of selection bias (Boulet 1994); unclear risk of reporting bias (Ponsioen 2005; Ribeiro 2007; Rytilä 2008).
5. Unclear risk of selection bias (Boulet 1994; Pornsuriyasak 2005) and other bias (Pornsuriyasak 2005).
6 Continuous outcome data based on total population size less than 400.
7 Wide 95% CI.
BACKGROUND

Cough, defined as “a forced expulsive manoeuvre, usually against a closed glottis and . . . associated with a characteristic sound” (Morice 2006), constitutes both a vital protective reflex and a common symptom of many pulmonary and several extra-pulmonary conditions.

In adults, subacute cough is defined as a cough of three to eight weeks duration, with a large proportion of cases due to postinfectious cough (Irwin 2006a; Kwon 2006). Chronic cough persists for more than eight weeks (Gibson 2010; Irwin 2006a; Morice 2004; Morice 2006), and, in most cases, is attributed to asthma, rhinitis or gastro-oesophageal reflux disease (GORD) (Irwin 1998; Gibson 2010), as well as other pulmonary conditions. While the prevalence of subacute cough is less clear, up to 40% of the population report having chronic cough (Morice 2004).

Description of the condition

Cough of intrapulmonary aetiology (lung-based cause) occurs when mechanical or chemical irritants activate vagal sensory nerve fibres in the airways and lungs, triggering a reflex arc that results in a co-ordinated motor output (Canning 2007). While failure of this defence mechanism can be life-threatening, increased sensitivity to the cough reflex is associated with excessive, persistent cough (Howden 2010). Chronic cough is associated with a substantial deterioration in quality of life, comparable to severe chronic obstructive pulmonary disease (COPD; French 1998), with diverse effects on all aspects of health, for example, causing musculoskeletal chest pain, sleep disturbance, anxiety and impaired social functioning (Birring 2003; French 2002; French 2004; McGarvey 2006). Given this associated morbidity, it is not surprising that cough is the most common symptom prompting presentation to general practice in Australia (Britt 2011). Cough is also associated with high rates of secondary care consultations, with chronic cough accounting for up to 38% of all referrals to respiratory physicians (Irwin 1990; McGarvey 1998a). In addition to these significant time costs, annual expenditure on over-the-counter and prescription treatments is probably in the order of billions of US dollars (Irwin 1998). While clinical guidelines recommend treatment of the underlying condition (Gibson 2010; Irwin 2006a; Morice 2007), a specific cause is not established in up to 46% of people who are described as having idiopathic or unexplained cough (Haque 2005; Irwin 2006b; Levine 2008; McGarvey 1998b; O’Connell 1994; Poe 1989).

Description of the intervention

Inhaled corticosteroids (ICS) may be administered via a metered dose inhaler (MDI), dry powder inhaler (DPI) or nebuliser (Bateman 2009). Through altering transcription of inflammatory mediators and direct actions on inflammatory cells, ICS suppress airway inflammation (Barnes 2006). Inhaled steroids are first-line therapy for many inflammatory airway diseases, including asthma (Bateman 2009), and are indicated for people with severe COPD and frequent exacerbations (Yang 2012). ICS may cause local side effects including oropharyngeal candidiasis (thrush in mouth/throat), dysphonia (hoarseness) and cough (Roland 2004). While direct delivery into the airways by inhalation significantly reduces the risk of systemic (whole body) side effects, absorption from the lungs can lead to complications including easy bruising, reduced bone mineral density and adrenal suppression (Bateman 2009; Lipworth 1999).

Why it is important to do this review

Clinical guidelines recommend empirical ICS treatment for non-specific and refractory cough (Gibson 2010), suspected cough-variant asthma (CVA) (Irwin 2006a; Morice 2004; Morice 2006), non-asthmatic eosinophilic bronchitis (Irwin 2006a; Morice 2004; Morice 2006), and atopic cough (Morice 2006). The efficacy of this intervention, however, remains contentious, with randomised controlled trials yielding conflicting results. A recent Cochrane systematic review has questioned the efficacy of ICS for non-specific cough in children (Tomerak 2009), and a Cochrane systematic review of ICS for subacute cough in children has recently been published (Anderson-James 2013). By providing systematic evidence relating to this intervention, this review aims to clarify uncertainty in current clinical practice and to elucidate the potential benefits of ICS in reducing the significant burden associated with subacute and chronic cough in adults. Through evaluating the strengths and limitations of current research evidence, this review also aims to inform future research directions.

OBJECTIVES

To assess the effects of ICS for subacute and chronic cough in adults.
METHODS

Criteria for considering studies for this review

Types of studies
As defined a priori in a published protocol (Johnstone 2011), we reviewed all published and unpublished randomised controlled trials (RCTs) comparing ICS with placebo for treatment of subacute and chronic cough in adults.

Types of participants
We considered all studies that included adults (over 18 years) with subacute or chronic cough, defined respectively as three to eight weeks, or more than eight weeks duration, respectively. We excluded participants with other known chronic respiratory diseases including asthma, chronic obstructive pulmonary disease and bronchiectasis, however, we included people with CVA (without demonstrated bronchodilator reversibility) and eosinophilic bronchitis (eosinophilic airway inflammation with sputum eosinophilia of greater than 2.5%; Gibson 2002).

Types of interventions
We included all randomised controlled comparisons of ICS versus placebo. ICS could be administered by MDI, DPI or nebuliser. Where trials included the use of other medications, all participants had to have equal access to such medications. We excluded trials without a placebo comparison group.

Types of outcome measures

Primary outcomes
1. Proportion of participants with clinical cure or significant improvement (over 70% reduction in cough severity measure) at follow up (clinical success).

Secondary outcomes
1. Proportion of participants with clinical cure or over 50% reduction in cough severity measure at follow up.
2. Proportion of participants with clinical cure at follow up.
3. Mean change in objective and subjective cough severity measures - cough frequency, cough receptor sensitivity, quality of life, Likert scale, visual analogue scale (VAS), level of interference of cough, cough diary.
4. Mean change in pulmonary function measures - bronchial hyper-responsiveness (BHR), spirometry, peak expiratory flow (PEF).
5. Complications of cough - requirement for medication change, time off work.
6. Biomarkers of inflammation - sputum biomarkers (total and differential cell counts, inflammatory mediators), exhaled gases.
7. Adverse effects of intervention - local side effects (oropharyngeal candidiasis, dysphonia, cough) and systemic side effects (easy bruising, reduced bone mineral density, adrenal suppression).

Where a study reported two or more cough severity measures, we used the first from the following hierarchy of cough severity measures:
1. Objective cough indices - cough frequency, cough receptor sensitivity.
2. Symptomatic measures as assessed by the participant - quality of life, Likert scale, VAS, level of interference of cough, cough diary.
3. Symptomatic measures as assessed by clinicians - Likert scale, VAS, level of interference of cough.

Where a study reported two or more cough severity measures of equal ranking on the hierarchy, we used the measure most comparable to those reported by other studies in the meta-analysis.

Search methods for identification of studies

Electronic searches
Trials were identified by searches of the following databases:
- The Cochrane Airways Group Trials Register (CAGR) (December 2012);
- Cochrane Central Register of Controlled Trials (CENTRAL), (The Cochrane Library 2012, Issue 12);
- MEDLINE (Ovid): (1948 to November week 3, 2012);
- EMBASE (Ovid): (1980 to week 49, 2012);
- ClinicalTrials.gov (December 2012).

The searches were conducted in December 2012, with no restriction on language of publication. Search strategies are listed in Appendix 1, Appendix 2, Appendix 3, Appendix 4 and Appendix 5.

Searching other resources
We reviewed the reference lists of all primary studies and review articles for additional references. We asked contact authors of included trials to identify other published and unpublished studies. We also searched manufacturers’ clinical trial registries.

Data collection and analysis

Selection of studies

Inhaled corticosteroids for subacute and chronic cough in adults (Review)

Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
Two reviewers (KJ, IY) independently assessed all potentially relevant studies identified through the search strategy for inclusion in the review. We resolved any disagreements through discussion.

**Data extraction and management**

For each trial that satisfied the inclusion criteria, we recorded the following information using a data collection form:

- **Design and methodology:** year of study, source of funding, design, randomisation, blinding (including allocation concealment and blinding of participants, care providers and outcome assessors) and statement of withdrawals.
- **Participants:** study setting, inclusion and exclusion criteria, participant recruitment details including number eligible, number enrolled, characteristics of study population (including age range, sex, ethnicity, diagnosis, other symptoms), number in treatment and control groups, baseline characteristics of treatment and control groups, number completing trial and number of withdrawals including reasons for withdrawal (e.g. clinical, side-effects, refusal) and whether intention-to-treat analysis is possible.
- **Interventions:** drug, dose, type of administration (i.e. MDI, DPI, nebulised), duration of intervention and co-interventions.
- **Outcomes:** primary and secondary outcomes as described above.

Two reviewers (KJ, IY) independently extracted data from included studies. Where required, we requested missing information from the study authors.

We calculated budesonide equivalent doses from the ranges outlined in the *Global strategy for asthma management and prevention* guidelines (Bateman 2009).

**Assessment of risk of bias in included studies**

Two reviewers (KJ, IY) independently assessed risk of bias for each study using the criteria outlined in *The Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We resolved any disagreement by consensus.

We assessed the risk of bias according to the following domains:

1. Random sequence generation (selection bias).
3. Blinding of participants and personnel (performance bias).
5. Incomplete outcome data (attrition bias).
6. Selective reporting (reporting bias).
7. Other bias.

We graded each potential source of bias as being high, low or at unclear risk of bias.

**Measures of treatment effect**

For all dichotomous outcomes, we calculated the odds ratio (OR) and 95% confidence interval (CI). We calculated the mean difference (MD) and 95% CI for all continuous outcomes. Where studies used different measurement scales, we calculated the standardised mean difference (SMD). In the case of missing data due to drop-outs, we performed a modified intention-to-treat analysis.

We compared the characteristics of each included study to determine whether meta-analysis of results was possible. We included all studies that satisfied the inclusion criteria and reported one or more outcomes of interest in the meta-analysis. We determined numbers needed to treat to benefit (NNTB) using an online calculator (Cates 2008).

We performed all statistical analysis using Review Manager 5 (RevMan) software.

**Unit of analysis issues**

We described all cross-over trials, and included them in meta-analysis where first period data was available. Where this was not available, we analysed data using the generic inverse variance method. Handling of cross-over trials in these ways differed from our protocol (see Differences between protocol and review).

**Dealing with missing data**

We contacted investigators or study sponsors in order to verify key study characteristics and obtain missing numerical outcome data, where possible.

**Assessment of heterogeneity**

We used the $I^2$ statistic to measure heterogeneity among the trials in each analysis. We assessed the importance of the level of heterogeneity identified as described in *The Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). Where possible, we explored substantial heterogeneity by pre-specified subgroup analysis.

**Assessment of reporting biases**

We assessed selective reporting within each trial by comparing the protocol and final published study, or otherwise the methods and results sections. Where we suspected reporting bias, we attempted to contact study authors asking them to provide missing outcome data. Where this was not possible, and the missing data were thought to introduce serious bias, we explored the impact of including such studies in the overall assessment of results by means of a sensitivity analysis. We explored publication bias using a funnel plot where meta-analysis with at least ten studies was possible.
Data synthesis

We used a fixed-effect model to calculate the summary ORs, MDs and their 95% CIs. Where there were concerns about statistical heterogeneity, we used a random-effects model. Where meta-analysis was not possible, or appropriate, we undertook a narrative review of the findings. We determined the quality of evidence for each pooled outcome based on the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach using GRADEprofiler 3.2.

Subgroup analysis and investigation of heterogeneity

Where possible, we carried out the following subgroup analyses for the primary outcome:
1. Dose of ICS (low to medium defined as < (less than) 800 µg/day budesonide equivalent and high defined as ≥ (more than or equal to) 800 µg/day budesonide equivalent).
2. Final diagnosis such as eosinophilic bronchitis, CVA, unexplained cough.
3. Duration of treatment (up to 4 weeks and > (more than) 4 weeks).
4. Duration of cough (3 to 8 weeks and > 8 weeks).

Sensitivity analysis

Where possible, we used sensitivity analysis to assess the robustness of the overall outcomes to the following factors:
1. Variation in inclusion criteria.
2. Risk of bias.
3. Study size.
4. Analysis using random-effects model.
5. Analysis by treatment received versus intention-to-treat.
6. Method of inhalation (e.g. MDI, DPI, nebuliser).

Results

Description of studies

Results of the search

A total of 1051 records were retrieved from electronic and hand-searches. Of these, eight studies reported in 16 references met the inclusion criteria. For full details of the study selection process, please see Figure 1. We excluded 12 studies; reasons for their exclusion are provided in the Characteristics of excluded studies table.
Figure 1. Study flow diagram showing study selection process.

1031 records identified through database searching

20 additional records identified through other sources

1051 records screened

1021 records excluded

30 full-text articles and abstracts assessed for eligibility

12 full-text articles and abstracts excluded, with reasons, and 2 studies awaiting classification

8 studies reported in 16 references included in qualitative synthesis

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

Key study characteristics and outcomes of the included studies are summarised in Characteristics of included studies and Figure 2.

Figure 2. Summary of included studies. Abbreviations BHR = bronchial hyper-responsiveness; cells = sputum total and/or differential cell count; dose = budesonide equivalent daily dose; DPI = dry powder inhaler; ECP = eosinophilic cationic protein; FEF25-75% = forced expiratory flow 25%-75%; FEV1 = forced expiratory volume in 1 second; FVC = forced vital capacity; gases = exhaled gases (CO, NO); meds = additional medication use; MDI = metered dose inhaler; night = nocturnal awakening; other = other sputum biomarkers (MPO, PGE2, LTB4, Cys-LT, IL-8, TNF-alpha, fibrinogen, albumin, substance P); PEF = peak expiratory flow; URTI = upper respiratory tract infection.

<table>
<thead>
<tr>
<th>Study ID</th>
<th>Cough duration</th>
<th>Participants (inclusion/exclusion criteria)</th>
<th>Intervention</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boulet 1994*</td>
<td>&gt; 4 weeks</td>
<td>×</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>Chaudhuri 2004</td>
<td>&gt; 1 year</td>
<td>×</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>Eivald 1999*</td>
<td>1 hour in half of the last 30 days</td>
<td>?</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>Pizzichini 1999</td>
<td>&gt; 1 year</td>
<td>×</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>Subacute chronic cough only</td>
<td>&gt; 3 weeks</td>
<td>?</td>
<td>?</td>
<td>×</td>
</tr>
<tr>
<td>Pomorskyaske 2005</td>
<td>≥ 3 weeks (post-URTI)</td>
<td>×</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>Ribello 2007</td>
<td>≥ 8 weeks</td>
<td>×</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>Rybka 2006</td>
<td>&gt; 8 weeks + 1 additional symptom</td>
<td>×</td>
<td>×</td>
<td>×</td>
</tr>
</tbody>
</table>

Key:
- Crossover (2 wks washout) = Mostly subacute
- Excluded
- Included
- Treated permitted
- Unclear
- No significant treatment effect
- Significant treatment effect
- Not reported

Parallel group: ×50 Mostly chronic

---

Inhaled corticosteroids for subacute and chronic cough in adults (Review)
Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
**Study design**

Of the eight randomised controlled trials identified, five were parallel group trials (Pizzichini 1999; Ponsioen 2005; Pornsuriyasak 2005; Ribeiro 2007; Rytilä 2008), two were cross-over trials that described first period results and were, therefore, able to be treated as parallel group trials (Boulet 1994; Evald 1989), and one was a cross-over trial with a two-week washout period between treatments (Chaudhuri 2004).

**Sample sizes**

The number of participants enrolled in each study ranged from 14 to 144 (Boulet 1994; Rytilä 2008).

**Setting**

Study locations included Brazil (Ribeiro 2007), Canada (Boulet 1994; Pizzichini 1999; Rytilä 2008), Denmark (Evald 1989), Finland, Greece, Hungary, Norway, Sweden (Rytilä 2008), Thailand (Pornsuriyasak 2005), the Netherlands (Ponsioen 2005), and the United Kingdom (Chaudhuri 2004; Rytilä 2008). Participants were recruited from hospital or specialist clinics (Boulet 1994; Evald 1989; Pornsuriyasak 2005; Rytilä 2008), primary care practices (Ponsioen 2005), or a combination of community and hospital settings (Chaudhuri 2004; Pizzichini 1999; Ribeiro 2007).

**Participants**

A total of 570 participants with cough were randomly allocated to receive ICS or placebo. All studies included adults only, with the exception of one smaller study that included participants aged 15 to 65 years (Evald 1989). While children aged over 15 years were also eligible for inclusion in Pornsuriyasak 2005, the mean ± SD age of participants (40.6 years ± 11.8 years for ICS, 38.8 years ± 13.0 years for placebo) suggests that this is less likely to have influenced the results of this study. Inclusion criteria for duration of cough ranged from at least two weeks (Ponsioen 2005) through to more than one year (Chaudhuri 2004; Pizzichini 1999). Unpublished data including only those participants with cough for at least three weeks was obtained for the study that included people with acute cough of two weeks (Ponsioen 2005). Four studies included people with both subacute and chronic cough. Participants with subacute cough predominated in two studies (Ponsioen 2005; Pornsuriyasak 2005), and participants with chronic cough predominated in two (Boulet 1994; Evald 1989). Four studies examined participants with chronic cough only (Chaudhuri 2004; Pizzichini 1999; Ribeiro 2007; Rytilä 2008). Most participants had nonspecific cough, except for those in a study specifically examining post-upper respiratory tract infection (URTI) cough (Pornsuriyasak 2005).

All studies excluded people with asthma, on the basis of history or spirometry, and those with known respiratory disease through history, examination, spirometry or chest X-ray. There was significant heterogeneity in terms of other eligibility criteria. Five studies excluded smokers (Boulet 1994; Chaudhuri 2004; Pizzichini 1999; Pornsuriyasak 2005; Rytilä 2008). People with demonstrated BHR were excluded from two studies (Boulet 1994; Pizzichini 1999), and only three people with a mildly positive bronchial provocation test were included in one study of 30 participants (Pornsuriyasak 2005). Four studies excluded people taking medications that potentially contribute to cough (Boulet 1994; Chaudhuri 2004; Pizzichini 1999; Pornsuriyasak 2005). Five studies excluded people with recent inhaled or oral steroid use (Boulet 1994; Chaudhuri 2004; Evald 1989; Pizzichini 1999; Ribeiro 2007). Six studies excluded people with recent respiratory infection (Boulet 1994; Chaudhuri 2004; Evald 1989; Pizzichini 1999; Ribeiro 2007; Rytilä 2008). It was unclear whether medications causing cough, recent steroid use or recent respiratory infection were among the exclusion criteria for Ponsioen 2005. People with GORD were excluded from three studies (Pornsuriyasak 2005; Ribeiro 2007; Rytilä 2008), and were specifically permitted in one study where this condition was treated (Pizzichini 1999). People with postnasal drip (PND), sinusitis or both were excluded from three studies (Ponsioen 2005; Ribeiro 2007; Rytilä 2008), and permitted in two studies where treated (Pizzichini 1999; Rytilä 2008).

**Interventions**

All trials compared ICS with placebo. The ICS used were beclomethasone, budesonide, fluticasone and mometasone. Daily budesonide equivalent doses of ICS ranged from 320 µg/day (Evald 1989), to 1600 µg/day (Boulet 1994; Chaudhuri 2004; Ponsioen 2005), with two trials investigating low to medium dose ICS and six trials investigating high dose ICS. Dose frequency ranged from one puff per day (Rytilä 2008) to two puffs four times daily (Boulet 1994). Treatment duration was two weeks in five studies (Chaudhuri 2004; Evald 1989; Pizzichini 1999; Ponsioen 2005; Ribeiro 2007), four weeks in two studies (Boulet 1994; Pornsuriyasak 2005) and eight weeks in the remaining study (Rytilä 2008). ICS was administered by DPI or MDI in four studies each. Two studies of MDI reported use of a spacer (Boulet 1994; Ponsioen 2005).

**Outcomes**

**Cough severity**

All studies reported some form of symptomatic measure assessed by the participant. This included cough symptom scores, VAS and cough diaries. Statistical reporting varied between studies. Changes in cough severity were reported as mean change from baseline, pre- and post-intervention means and differences of differences. Primary outcome data were available for only three studies (Boulet 1994; Ponsioen 2005; Ribeiro 2007). None of the studies used cough meters to quantify cough objectively.
Pulmonary function

Four studies assessed BHR (Boulet 1994; Ponsioen 2005; Pornsuriyasak 2005; Rytilä 2008), and three studies reported this outcome (Boulet 1994; Ponsioen 2005; Pornsuriyasak 2005). Four studies stated that they investigated forced expiratory volume in one second (FEV₁) (Evald 1989; Pizzichini 1999; Ponsioen 2005; Pornsuriyasak 2005), though this outcome was only reported in two (Ponsioen 2005; Pornsuriyasak 2005). Three investigated forced vital capacity (FVC) (Evald 1989; Pizzichini 1999; Pornsuriyasak 2005), although data was reported in only one of these trials (Pornsuriyasak 2005). One study measured change in forced expiratory flow 25% to 75% (FEF₂₅₋₇₅%) (Pornsuriyasak 2005). Two studies examined changes in peak expiratory flow (Evald 1989; Rytilä 2008).

Complications of cough


Sputum biomarkers of inflammation

Several studies measured changes in sputum biomarkers of inflammation, including total and differential cell counts (Chaudhuri 2004; Pizzichini 1999), eosinophils (Chaudhuri 2004; Pizzichini 1999; Rytilä 2008), and eosinophilic cationic protein (ECP) (Chaudhuri 2004; Pizzichini 1999; Rytilä 2008). Two studies also investigated a range of other inflammatory mediators (Chaudhuri 2004; Pizzichini 1999). Chaudhuri 2004 also assessed the effect of ICS on exhaled nitric oxide (eNO) and carbon monoxide (CO).

Adverse effects of intervention

Four studies investigated adverse effects, which were variably defined in terms of local side effects (Pizzichini 1999), and adverse effects that were considered to be related to treatment (Ponsioen 2005), or might be related to treatment (Rytilä 2008). In Ribeiro 2007, adverse effects was not defined as an outcome of interest, but was reported in the results.

Excluded studies

We excluded 12 studies; the reasons for their exclusions are detailed in the Characteristics of excluded studies table. The most common reason for exclusion was lack of a placebo comparison group.

Risk of bias in included studies

The overall quality of included studies was generally good, with several studies having a low risk of bias in nearly all categories, as shown in Figure 3. Unpublished data was sought for all studies. For full details, please see Characteristics of included studies.
Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

<table>
<thead>
<tr>
<th>Study</th>
<th>Random sequence generation (selection bias)</th>
<th>Allocation concealment (selection bias)</th>
<th>Blinding of participants and personnel (performance bias)</th>
<th>Blinding of outcome assessment (detection bias)</th>
<th>Incomplete outcome data (attrition bias)</th>
<th>Selective reporting (reporting bias)</th>
<th>Other bias</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boulet 1994</td>
<td>?</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Chaudhuri 2004</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>?</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Pizzichini 1999</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>?</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Ponsioen 2005</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>?</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Ribeiro 2007</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>?</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Rytilä 2008</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>?</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>
Allocation
Reports of five studies described the method of randomisation used (Chaudhuri 2004; Pizzichini 1999; Ponsioen 2005; Ribeiro 2007; Rytilä 2008). The risk of bias resulting from random sequence generation was unclear in the remaining three studies. The method of allocation concealment was adequately described in five studies (Chaudhuri 2004; Pizzichini 1999; Ponsioen 2005; Ribeiro 2007; Rytilä 2008), and was unclear in the remaining three.

Blinding
All studies were described as double blind, resulting in a low risk of performance bias and detection bias. Five studies described the use of an identical or matching placebo inhaler (Chaudhuri 2004; Pizzichini 1999; Ponsioen 2005; Ribeiro 2007; Rytilä 2008).

Incomplete outcome data
Two studies reported no drop-outs (Boulet 1994; Ribeiro 2007), and, in the four studies that reported the groups from which drop-outs occurred, attrition rates were comparable for the treatment and control groups, which resulted in an assessment of a low risk of attrition bias for those trials (Pizzichini 1999; Ponsioen 2005; Pornsuriyasak 2005; Rytilä 2008). The remaining two studies did not state from which treatment group participants withdrew, resulting in an assessment of an unclear risk of attrition bias (Chaudhuri 2004; Evald 1989).

Selective reporting
Protocols were not available for any of the included studies, therefore, we compared the methods and results sections of all included studies. Risk of reporting bias was unclear or high in the majority of studies, with only two receiving a grading of low risk (Boulet 1994; Chaudhuri 2004). Reasons for a grading of unclear or high risk of reporting bias included inadequate reporting of number of people screened and eligible (Evald 1989; Pornsuriyasak 2005) and outcomes in either the methods or results section (Evald 1989; Pizzichini 1999; Ponsioen 2005; Ribeiro 2007; Rytilä 2008).

Other potential sources of bias
Chaudhuri 2004 noted no carry-over effect in the active treatment group, resulting in a low risk of bias. The two cross-over trials without a washout period described first period results, and thus were able to be treated as parallel group trials, resulting in a low risk of bias (Boulet 1994; Evald 1989). The inclusion criterion of post-URTI cough was inadequately defined in Pornsuriyasak 2005, which led to an unclear risk of bias grading. Compliance was reasonably high in the two studies in which it was monitored (Ponsioen 2005; Rytilä 2008).

Effects of interventions
See: Summary of findings for the main comparison Inhaled corticosteroids (ICS) compared to placebo for adults with subacute and chronic cough

Cough severity

Cross-over trials

Studies showing significant improvement
Chaudhuri 2004 investigated the efficacy of fluticasone 1000 µg/day or placebo for two weeks among 93 non-smoking participants who had had chronic cough for over a year, using a two-week washout period between treatments. A significant reduction in cough VAS score was noted after treatment with ICS compared with placebo (mean ± standard error of the mean (SEM) 1.0 cm ± 0.3 cm). People taking medications causing cough as well as those with recent steroid use and recent respiratory infection were not eligible. Cough in these people was attributable to PND (34%), GORD (20%), CVA (15%), bronchiectasis (10%), eosinophilic bronchitis (6%), habitual cough (2%) and bronchitis (1%). Ten people (11%) had idiopathic cough.

Parallel group trials

Studies showing no significant improvement
Two studies were designed as cross-over trials but reported first period results (Boulet 1994; Evald 1989), which allowed them to be treated as parallel group trials. In a trial of 14 non-smokers with cough for more than four weeks, Boulet 1994 found no significant reduction in overall mean cough scores after participants had received either beclomethasone 2000 µg/day or placebo for four weeks (MD -0.91 on a cough score scale of 0 to 10; 95% CI -2.24 to 0.42). Participants had normal airway responsiveness, were not taking medications causing cough and had no history of recent oral, or inhaled, steroid use or recent respiratory infection. Cough was attributable to GORD (36%), PND (21%), both (31%) or no specific cause (21%). The Evald 1989 trial of 40 non-smokers with daily dry cough for at least one hour in more than half of the previous 30 days, found no significant reduction in cough symptom scores for participants.
who received either beclomethasone 400 µg/day or placebo for two weeks (data not reported). Smokers, people treated with anti-asthmatic drugs, and those with recent respiratory infection were excluded.

Four parallel group trials found that ICS did not significantly reduce cough severity (Pizzichini 1999; Ponsioen 2005; Pornsuriyasak 2005; Rytilä 2008). In a study of 50 non-smokers who had had chronic cough for over a year, Pizzichini 1999 found no statistically significant reduction in proportion with over 50% reduction in VAS score (OR 0.53; 95% CI 0.06 to 4.91) following treatment with budesonide 800 µg/day for two weeks versus placebo. People with BHR, medications causing cough, recent steroid use, recent respiratory infection and untreated GORD, PND and sinusitis were excluded.

A trial of inhaled fluticasone 1000 µg/day for two weeks versus placebo in 135 participants with cough for at least two weeks demonstrated a significant reduction in cough score among participants (Ponsioen 2005). Smokers, people with BHR and those with GORD were included. The eligibility of people taking medications causing cough or those with recent steroid use or recent respiratory infection was unclear. Unpublished data, however, that excluded the subgroup participants with acute cough demonstrated that ICS treatment did not significantly increase the odds of achieving clinical cure or more than 70% improvement in cough severity measure. ICS treatment did not produce a significant reduction in cough severity measure (MD -0.38 on a cough score scale of 0 to 3; 95% CI -1.05 to 0.28). Pornsuriyasak 2005 studied the effect of inhaled budesonide 800 µg/day versus placebo for four weeks among 30 participants with persistent post-URTI cough for more than three weeks. Oral corticosteroids were terminated one week prior to entry into the study. Exclusion criteria included smoking, BHR, use of medications causing cough, GORD and untreated sinusitis. Symptom scores were not significantly different between the ICS and placebo groups after two (MD -0.33; 95% CI -1.66 to 1.00) or four weeks (MD -0.36; 95% CI -1.52 to 0.80) of treatment.

Rytilä 2008 investigated the efficacy of mometasone 400 µg/day versus placebo for eight weeks in a trial with 144 participants who had had cough for at least two months with at least one of the following additional symptoms: chest tightness, wheezing, shortness of breath, or exercise-induced cough or wheezing. Smokers and participants with recent respiratory infection, GORD, PND and sinusitis were excluded. At follow up, there was no significant difference between the two groups in the proportion of participants who were symptom free (29% ICS, 26% placebo, P value 0.7). This study demonstrated no significant change in morning and evening cough scores after four or eight weeks, despite a significant improvement in total morning symptom score after eight weeks.

One parallel group trial showed evidence of significant improvement in cough severity with ICS treatment. Ribeiro 2007 examined the effect of beclomethasone 1500 µg/day versus placebo for two weeks among 64 participants who had had chronic cough for at least eight weeks. Extrapulmonary causes of cough, including GORD and PND, were excluded. Treatment with ICS caused complete resolution in 82% of participants, compared with 15% in the placebo group (P value < 0.05), and resulted in a significantly greater mean reduction in cough diary score (1.37 ± 1.21 ICS, 0.54 ± 0.7 placebo) and VAS (79% ± 29.3% ICS, 15.1% ± 31.1% placebo).

Combined results

Proportion of participants with clinical cure or significant improvement (over 70% reduction in cough severity measure) at follow up (clinical success)

Only three studies contributed data for the primary outcome measure (Proportion of participants with clinical cure or significant improvement (over 70% reduction in cough severity measure) at follow up) in a form suitable for meta-analysis, although ultimately we were unable to pool it due to heterogeneity. Each of these studies examined high dose ICS via MDI for a maximum of four weeks (Analysis 1.1). One of these studies found a significant treatment effect (Ribeiro 2007). Participants had cough for at least three weeks (Ponsioen 2005), more than four weeks (Boulet 1994), or at least eight weeks (Ribeiro 2007). These studies differed in their eligibility criteria relating to exclusion of smokers, BHR, medications causing cough (Boulet 1994), and those with GORD or PND/sinusitis (Ribeiro 2007). While people with recent steroid use and recent respiratory infection were excluded from both Boulet 1994 and Ribeiro 2007, it was unclear whether or not these people and those taking medications causing cough were excluded from Ponsioen 2005. Substantial to considerable clinical heterogeneity was reflected in statistical heterogeneity (I² = 85%). Therefore, pooling of results was deemed inappropriate. The quality of evidence was low because of the unclear risk of selection (Boulet 1994) and reporting bias in included studies (Ponsioen 2005; Ribeiro 2007), and the small number of recorded events.

Proportion of participants with clinical cure or over 50% reduction in cough severity measure at follow up

Data pertaining to the proportion of participants who achieved a more than 50% reduction in cough severity measure at follow up were available from four studies, however heterogeneity in study characteristics limited the validity of meta-analysis (Analysis 1.2). This did not include the cross-over study that found a significant treatment effect (Chaudhuri 2004), or one of the participants with BHR, medications causing cough, recent steroid use, recent respiratory infection and untreated GORD, PND and sinusitis were excluded. At follow up, there was no significant difference between the two groups in the proportion of participants who were symptom free (29% ICS, 26% placebo, P value 0.7). This study demonstrated no significant change in morning and evening cough scores after four or eight weeks, despite a significant improvement in total morning symptom score after eight weeks.

Studies showing significant improvement

Inhaled corticosteroids for subacute and chronic cough in adults (Review)

Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
larger higher quality studies which found no significant treatment effect (Rytilä 2008). Eligibility criteria for the studies were quite heterogeneous. Cough duration ranged from at least three weeks (Ponsioen 2005 subacute and chronic cough participants only), to more than one year (Pizzichini 1999), and studies varied in terms of whether smoking, BHR, use of medications causing cough (Boulet 1994; Pizzichini 1999), recent steroid use (Boulet 1994; Pizzichini 1999; Ribeiro 2007), recent respiratory infection (Boulet 1994; Pizzichini 1999; Ribeiro 2007), GORD (Pizzichini 1999 (untreated); Ribeiro 2007) and PND/sinusitis (Pizzichini 1999 (untreated); Ponsioen 2005; Ribeiro 2007) were among the exclusion criteria. Each of these studies used high dose ICS for up to four weeks duration via either DPI or MDI. Cough severity was measured in terms of cough symptom scores rated between 0 and 10 (Boulet 1994), or 0 and 3 (Ponsioen 2005), cough diary scores taking into account frequency, severity, duration and sleep interruption (Ribeiro 2007), and cough discomfort VAS scores (Pizzichini 1999; Ribeiro 2007). The quality of each of these studies was generally good. Among this diverse population using high dose ICS for up to four weeks, the calculated I² statistic was 81%, which may represent substantial to considerable heterogeneity. The quality of evidence was low, with evidence downgraded due to the unclear risk of selection bias in Boulet 1994 and small number of events.

**Proportion of participants with clinical cure at follow up**

Data for the proportion of participants with clinical cure were available from four studies (Analysis 1.3), one of which found a significant difference in cure rates (Ribeiro 2007). Of these four studies, one examined predominantly subacute cough (Ponsioen 2005), and three predominantly chronic cough (Boulet 1994; Ribeiro 2007; Rytilä 2008). All studies had a low risk of bias in nearly all categories. These studies were considered too heterogeneous in their participants, interventions and outcome measures to pool. Studies varied as to whether smoking (Boulet 1994; Rytilä 2008), BHR (Boulet 1994), taking medications causing cough (Boulet 1994), recent steroid use (Boulet 1994; Ribeiro 2007), recent respiratory infection (Boulet 1994, Rytilä 2008), and extrapulmonary causes were among the exclusion criteria (Ribeiro 2007; Rytilä 2008). Several of these eligibility criteria were unclear in Ponsioen 2005. Interventions included high dose ICS administered via MDI for up to four weeks (Boulet 1994; Ponsioen 2005; Ribeiro 2007), and low dose ICS administered by DPI for eight weeks (Rytilä 2008). Clinical cure was defined as a cough symptom score of zero (Boulet 1994; Ponsioen 2005), resolution of cough (Ribeiro 2007), or a total symptom score of 0 to 2 out of 36 (Rytilä 2008). For these reasons, pooling of these studies was deemed inappropriate. Substantial to considerable heterogeneity was reflected by the calculated I² statistic of 82%. The overall quality of this narrative evidence was low due to the unclear risk of selection bias (Boulet 1994), and reporting bias in included studies (Ponsioen 2005; Ribeiro 2007; Rytilä 2008), and the small number of events.

**Mean reduction in cough severity measure**

**Cough score**

For mean change in cough symptom score, data were available from five studies that included 364 participants; one of the studies found ICS to be beneficial (Ribeiro 2007). Interventions were high dose ICS for two weeks (Ponsioen 2005; Ribeiro 2007), or four weeks (Boulet 1994; Pornsuriyasak 2005), or low dose ICS for eight weeks (Rytilä 2008). These studies differed in their eligibility criteria. Participants had cough for at least three weeks (Ponsioen 2005 subacute and chronic cough participants only), more than four weeks following an URTI (Pornsuriyasak 2005), more than four weeks (Boulet 1994), or at least eight weeks (Ribeiro 2007; Rytilä 2008). Participants in Rytilä 2008 also had at least one additional symptom suggestive of asthma. Smokers were excluded from three studies (Boulet 1994; Pornsuriyasak 2005; Rytilä 2008). People with BHR and those taking medications potentially causing cough were excluded from Boulet 1994 and Pornsuriyasak 2005. Recent steroid use was excluded in Boulet 1994 and Ribeiro 2007, whereas oral corticosteroids were terminated at least one week prior to entry in Pornsuriyasak 2005. Recent respiratory infection was excluded in Boulet 1994, Ribeiro 2007 and Rytilä 2008, whereas Pornsuriyasak 2005 specifically examined post-URTI cough. Three studies also excluded GORD (Pornsuriyasak 2005; Ribeiro 2007; Rytilä 2008), and four excluded PND/sinusitis (Ponsioen 2005; Pornsuriyasak 2005 (untreated); Ribeiro 2007; Rytilä 2008). Cough symptom scores were measured on scales from 0 to 3 (Rytilä 2008 (morning cough score)), 0 to 6 (Boulet 2005 calculated from sum of daytime and nighttime scores), and 0 to 10 (Boulet 1994), and, in the other two studies, were calculated from scores for factors including frequency and sleep interruption (Pornsuriyasak 2005; Ribeiro 2007). Boulet 1994 and Pornsuriyasak 2005 were small studies with a low or unclear risk of bias in most categories. Ponsioen 2005, Ribeiro 2007 and Rytilä 2008 were larger studies that achieved a low risk of bias rating in six of seven domains. The pooled study data demonstrated a significant standardised mean reduction in cough score of -0.34 (95% CI -0.56 to -0.13; 346 participants; Analysis 1.4; Figure 4). This result is unlikely to be clinically significant given that this improvement correlates to less than the minimal response defined by Ribeiro 2007 (a reduction of at least two points for each question) in each of the five included studies. The calculated I² statistic was 0%. The overall quality of evidence was graded as low due to the unclear risk of selection bias (Boulet 1994; Pornsuriyasak 2005) and other bias (Pornsuriyasak 2005) in two included studies, and a total study population size less than 400.
Figure 4. Forest plot of comparison: ICS versus Placebo, outcome: Mean change in cough score.

<table>
<thead>
<tr>
<th>Study or Subgroup</th>
<th>ICS</th>
<th>Placebo</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boulet 1994</td>
<td>-1.15</td>
<td>1.60</td>
<td>7</td>
</tr>
<tr>
<td>Ponsioen 2005</td>
<td>-2.398</td>
<td>1.764</td>
<td>52</td>
</tr>
<tr>
<td>Pornsuriyasak 2005</td>
<td>-1.07</td>
<td>1.74</td>
<td>14</td>
</tr>
<tr>
<td>Ribeiro 2007</td>
<td>-1.16</td>
<td>1.24</td>
<td>2</td>
</tr>
<tr>
<td>Rytilä 2008</td>
<td>-0.44</td>
<td>0.699</td>
<td>78</td>
</tr>
</tbody>
</table>

Total: 159

VAS for cough

Data for mean change in VAS score were available for only two studies. Both examined the efficacy of high dose ICS for two weeks. Exclusions, of smokers and people with use of medications causing cough (Chaudhuri 2004), recent steroid use, recent respiratory infection, GORD and PND/sinusitis (Ribeiro 2007), varied between the two studies. The overall risk of bias was low in both studies. The I² statistic of 86% calculated from these pooled data indicated substantial to considerable heterogeneity. For these reasons, pooling was deemed inappropriate for this outcome.

Pulmonary function

Pulmonary function was generally not affected by ICS treatment, with the exception of one study that detected a significant improvement in PEF. None of the three studies that measured BHR found a significant difference after treatment with ICS versus placebo (Boulet 1994; Ponsioen 2005; Pornsuriyasak 2005). BHR data were available as dichotomous data for Ponsioen 2005 (Analysis 1.12), and continuous data for Boulet 1994 (Analysis 1.13), which prevented pooling of data for this outcome.

Two studies found no significant improvement in FEV₁ following treatment with ICS (Ponsioen 2005; Pornsuriyasak 2005), with data available for only one of these (Ponsioen 2005; Analysis 1.14). The one study that reported FVC and FEF25–75% demonstrated no significant improvement in either outcome (Pornsuriyasak 2005). Improvement in PEF varied, with one study finding a significant treatment effect for morning PEF after four weeks and eight weeks, and evening PEF after eight weeks (Rytilä 2008), and the other study finding no significant effect (Evald 1989). Insufficient data were available to produce a meta-analysis for change in pulmonary function.

Complications of cough

Requirement for additional use of medication was the only complication of cough shown to be reduced by ICS treatment. Specifically, requirement for additional medication after the treatment period (Ponsioen 2005; Analysis 1.15), and use of reliever medication (Rytilä 2008), were significantly decreased. Three studies that examined frequency of nocturnal awakenings noted no significant reduction with ICS treatment (Evald 1989; Ponsioen 2005; Ribeiro 2007), and data were reported for only one study (Ribeiro 2007; Analysis 1.16). Ponsioen 2005 found no significant decrease in days off work. Pooling was not possible due to inadequate reporting of data.

Biomarkers of inflammation

Sputum total and differential cell counts

Sputum total and differential cell counts were not significantly affected by ICS (Chaudhuri 2004; Pizzichini 1999), with the exception of eosinophils, which were significantly reduced in one study (Rytilä 2008), but not in two others (Chaudhuri 2004; Pizzichini 1999). Where the final diagnosis was known (Chaudhuri 2004), subgroup analysis showed a significant improvement in sputum eosinophilia among participants with cough attributable to CVA (-4.60%; 95% CI -7.10 to -2.10; Analysis 1.21).

Inflammatory mediators

ECP was the only inflammatory mediator significantly reduced by ICS (Chaudhuri 2004; Rytilä 2008), but this was not seen in all studies (Pizzichini 1999). Subgroup analysis by cough aetiology in Chaudhuri 2004 (Analysis 1.18) revealed no significant effects. No study showed significant improvement in levels of other sputum biomarkers of inflammation including interleukin-8 (IL-8) (Chaudhuri 2004; Pizzichini 1999), cysteinyl leukotriene (Cys-LT), leukotriene B4 (LTB₄), myeloperoxidase (MPO) and prostaglandin E₂ (PGE₂), tumour necrosis factor alpha (TNF-
α) (Chaudhuri 2004), or fibrinogen, albumin, substance P and bronchial epithelial cells (Pizzichini 1999). When analysed by final diagnosis in Chaudhuri 2004, participants with bronchiectasis showed a significant increase in MPO (133.5 µg/mL; 95% CI 27.0 to 239.9; Analysis 1.23) and a significant reduction in IL-8 (-74.7 ng/mL; 95% CI -146.3 to -3.1; Analysis 1.27). Insufficient data prevented meta-analysis for these outcomes.

Exhaled gases

In Chaudhuri 2004, ICS treatment resulted in significant reductions in eNO overall (-2.1 ppb; 95% CI -3.6 to -0.6), and specifically among participants with cough attributable to GORD (-3.1 ppb; 95% CI -5.8 to -0.5) and CVA (-3.3 ppb; 95% CI -6.5 to -0.2) (Analysis 1.29). Carbon monoxide was significantly reduced overall (-0.3 ppm; 95% CI -0.6 to -0.0), but not for any individual subgroup of cough aetiology (Analysis 1.30).

Adverse effects of intervention

Treatment with ICS was not associated with a significant increase in total adverse effects compared with placebo in the studies that reported this outcome (Pizzichini 1999; Ponsioen 2005; Ribeiro 2007; Rytilä 2008). In terms of specific adverse effects, no significant differences were found in hoarseness, sore throat, oral candidiasis (Ponsioen 2005; Ribeiro 2007), or severe adverse effects (Ponsioen 2005; Ribeiro 2007; Rytilä 2008). More commonly reported adverse effects included hoarseness, sore throat, dry mouth and headache.

Combined results

The proportion of participants with adverse effects was reported in four studies (Pizzichini 1999; Ponsioen 2005; Ribeiro 2007; Rytilä 2008), allowing meta-analysis for this outcome. These studies all had a low risk of bias in six of the seven domains assessed. These trials examined a heterogeneous population comprising participants with cough for at least two weeks (Pornsuriyasak 2005), to more than one year (Pizzichini 1999). Studies also varied in terms of exclusion of smokers (Pizzichini 1999; Rytilä 2008), people with BHR (Pizzichini 1999), medications causing cough (Pizzichini 1999; Ponsioen 2005), recent steroid use (Ponsioen 2005; Ribeiro 2007), recent respiratory infection (Pizzichini 1999; Ribeiro 2007; Rytilä 2008), and people with GORD (Pizzichini 1999 (untreated); Ribeiro 2007; Rytilä 2008). People with PND/sinusitis were excluded from three of these studies (Ponsioen 2005; Ribeiro 2007; Rytilä 2008), with adequately treated participants only being permitted in one study (Pizzichini 1999). It was unclear whether use of medications causing cough, recent steroid use or recent respiratory infection were among the exclusion criteria for Ponsioen 2005. Three studies were of high dose ICS for two weeks (Pizzichini 1999; Ponsioen 2005; Ribeiro 2007), with one study examining low dose ICS for eight weeks via either MDI or DPI (Rytilä 2008). No individual study reported a significant increase in adverse effects, and the pooled effect estimate was also not statistically significant (OR 1.67; 95% CI 0.92 to 3.04; 381 participants; Analysis 1.31; Figure 5). The calculated I² statistic of 0% indicated that heterogeneity in study populations, interventions and measured outcomes was probably not important. The quality of evidence was downgraded to moderate due to the wide confidence interval.

Figure 5. Forest plot of comparison: I ICS versus Placebo, outcome: 1.31 Proportion with adverse effects.

<table>
<thead>
<tr>
<th>Study or Subgroup</th>
<th>ICS Events</th>
<th>Placebo Events</th>
<th>Total</th>
<th>Weight</th>
<th>Odds Ratio M-H, Fixed, 95% CI</th>
<th>Odds Ratio M-H, Fixed, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pizzichini 1999</td>
<td>3</td>
<td>21</td>
<td>24</td>
<td>4.8%</td>
<td>3.67 [0.35, 38.34]</td>
<td></td>
</tr>
<tr>
<td>Ponsioen 2005</td>
<td>10</td>
<td>55</td>
<td>65</td>
<td>43.8%</td>
<td>1.19 [0.45, 3.16]</td>
<td></td>
</tr>
<tr>
<td>Ribeiro 2007</td>
<td>2</td>
<td>44</td>
<td>46</td>
<td>3.3%</td>
<td>2.41 [0.11, 52.57]</td>
<td></td>
</tr>
<tr>
<td>Rytilä 2008</td>
<td>18</td>
<td>70</td>
<td>88</td>
<td>47.9%</td>
<td>1.88 [0.85, 4.29]</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>200</td>
<td>181</td>
<td>381</td>
<td>100.0%</td>
<td>1.67 [0.92, 3.04]</td>
<td></td>
</tr>
<tr>
<td>Total events</td>
<td>33</td>
<td>21</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heterogeneity Chi²</td>
<td>1.01</td>
<td>df = 3 (P = 0.80); P = 0%</td>
<td>Test for overall effect: Z = 1.76 (P = 0.09)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(1) Local side effects only
Pooling of results was not appropriate for the outcomes of specific and severe adverse effects due to these events being infrequent, or not present in studies (Analysis 1.32; Analysis 1.33).

**DISCUSSION**

**Summary of main results**

Eight studies examined the efficacy of ICS in people with subacute and chronic cough associated with frequently undiagnosed, as well as unexplained, causes. Studies were heterogeneous in their populations, interventions, outcome measures, reporting and results (Figure 2). A significant reduction in cough severity was observed in one cross-over trial and one parallel group trial. Please see [Summary of findings for the main comparison](#) for details of the main findings.

For the primary outcome of clinical cure or significant improvement (more than 70% reduction in cough severity measure) at follow up (clinical success), data were available for only three studies that were too heterogeneous in their study characteristics to allow meta-analysis.

Meta-analysis for the outcomes of proportion with more than 50% improvement and clinical cure was not possible due to heterogeneity in study design.

The mean decrease in cough score following ICS treatment was 0.34 standard deviations lower compared to placebo (95% CI -0.56 to -0.13; 346 participants) following ICS, though the quality of this evidence was low. Meta-analysis was not possible for the outcome of mean change in VAS score due to heterogeneity in study characteristics.

In terms of pulmonary function, there was no improvement in BHR in the three studies of high dose short duration ICS that examined this outcome. Two studies demonstrated no significant improvement in spirometry. PEF significantly improved in one study but not in another.

Treatment with ICS was associated with a reduction in additional medication requirement. No effect on nocturnal awakenings, or days off work, was found in three studies, and one study, respectively.

Significant improvements in sputum eosinophils were demonstrated in one study but not in two others. ECP was significantly reduced by ICS in two out of three studies. Sputum total cell counts, neutrophils and lymphocytes were not improved in two studies. No study showed improvement in levels of other sputum biomarkers. Exhaled NO and CO significantly decreased in one study but not in another.

While the moderate quality evidence demonstrated no statistically significant increase in adverse effects, the potential benefits of ICS therapy need to be considered in the light of these possible harms.

**Impact of heterogeneity of studies**

The conflicting findings probably result from significant diversity in study design, participants, interventions and outcome measures between studies.

**Study design**

Cross-over trials of persistent cough increase risk of bias due to the potential for a prolonged carry-over effect, which is difficult to control for with a washout period. For instance, Boulet 1994 attributed the reduction in cough symptoms in four participants who received placebo during the second treatment period to the persistent effect of beclomethasone from the first period. However, Chaudhuri 2004 found no carry-over effect.

**Participants**

Inconsistent study findings are most likely largely attributable to heterogeneity in study populations. This was described as a limitation in Rytilä 2008. Heterogeneity among review participants is explained by the diversity in aetiology of cough due to the inherently unexplained nature of the condition, as well as differences in sample sizes, study settings and eligibility criteria.

**Sample sizes**

None of the smaller studies, i.e. with 50 participants or fewer, found ICS to be effective (Boulet 1994; Evald 1989; Pizzichini 1999; Pornsuriyasak 2005). This is likely to reflect the fact that these had inadequate statistical power to detect a significant treatment effect. Inadequate sample size was recognised as a limitation in two study reports (Pornsuriyasak 2005; Rytilä 2008).

**Setting**

Differences in study settings may have influenced the types of participants recruited, which would have contributed to heterogeneity in study populations, and, in turn, to inconsistencies in the efficacy of ICS. None of the studies that recruited participants exclusively from hospital or specialty clinics found ICS to be beneficial (Boulet 1994; Evald 1989; Pornsuriyasak 2005; Rytilä 2008). As Rytilä 2008 recognised, only participants with severe symptoms were likely to be included in such a setting. While Ponsioen 2005 found no correlation between baseline cough severity and treatment effect, it is possible that people presenting to specialist clinics may have a cough that is more refractory to treatment for other reasons, for example if previous primary care interventions have been unsuccessful. Ponsioen 2005, however, who recruited exclusively from primary care practices, did not find a significant treatment effect once people with acute cough were excluded. Studies of participants recruited from a combination of community and hospital settings found ICS to be effective in some instances (Chaudhuri 2004; Ribeiro 2007), but not others (Pizzichini 1999).

**Cough duration**

---

Inhaled corticosteroids for subacute and chronic cough in adults (Review)

Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
Some authors suggested that people with cough of shorter duration may be more responsive to ICS (Chaudhuri 2004), and a correlation between cough duration and treatment response was seen in Ribeiro 2007. While this does not appear to be supported by the results of this review, confounding factors are probably present. ICS was not effective in reducing cough among trials of people with predominantly subacute cough. While Ponsioen 2005 noted a significant improvement following ICS among participants with cough for at least two weeks duration, the treatment effect was not significant once 31 participants with acute cough were excluded from the analysis (mean cough duration 5.5 weeks (SD 3.2)).

Furthermore, Ponsioen 2005 found no significant treatment effect among participants with post-URTI cough of greater than three weeks duration (mean 5.93 weeks (SD 1.94)), however this may be more of a reflection of the differing aetiology of subacute cough. None of the studies of people with predominantly chronic cough found a significant treatment effect (Boulet 1994; Evald 1989), however, confounding factors were probably present. For instance, Evald 1989 suggested that the significant period effect observed suggested that some participants may have had infectious cough, which is not responsive to ICS (Frank 1985). There was no clear relationship between cough duration and ICS efficacy in the studies that included only chronic cough participants. ICS reduced cough severity in studies of people with cough for a mean 44.6 weeks (SD 86.2) (Ribeiro 2007), and mean 16.2 years (SD 16.1) (Chaudhuri 2004), but not in two other studies where mean cough duration was 9.8 years (95% CI 5.3 to 14.2) and 11.8 years (95% CI 4.6 to 19.2) for ICS and placebo groups respectively (Pizzichini 1999), and not specified (Rytilä 2008). Rytilä 2008 suggested the inconsistency in response to ICS observed in their study might be partially attributable to “high variability of symptoms with asymptomatic periods”.

**Age**

Cough aetiology and management differs between children and adults (Chang 2005). Hence, the inclusion of children older than 15 years in Evald 1989 may have contributed to the poor response to ICS in this study, given that ICS may not be effective in children who have had cough for more than three weeks (Tomerak 2009).

**Bronchial hyper-responsiveness**

The relationship between BHR and the steroid-responsive condition of CVA may explain why ICS was effective only in studies that did not exclude people with BHR. Bronchial provocation testing is recommended for the evaluation of cough in people with suspected CVA who have had a non-diagnostic physical examination and spirometry (Irwin 2006a). Due to its negative predictive value of nearly 100%, a negative test effectively excludes CVA (Madison 2010). The positive predictive value of bronchial provocation testing is lower (60% to 80%), and hence the presence of BHR does not necessarily mean the patient has asthma. Therefore, response to anti-asthma treatment is required to diagnose CVA (Madison 2010).

Both studies that found ICS to be effective did not exclude participants with BHR. Ribeiro 2007 noted a significant treatment effect among participants with baseline mean ± SD PD20 measurements of 5.35 mg ± 3.2 mg and 4.56 mg ± 3.7 mg for treatment and control groups, respectively. It should be noted that the significant difference between these groups at baseline was considered to be unimportant given that the PD20 values of participants with BHR were higher than those typically seen in asthma (usually less than 4 mg/mL), and, therefore, could be consistent with hyper-responsiveness found in healthy people (Ribeiro 2007). While Chaudhuri 2004 did not report baseline BHR data, participants with CVA showed the greatest improvement in VAS. CVA was diagnosed based primarily on response to ICS and short-acting β2-agonist after the treatment period, however, bronchial challenge testing was also utilised to help establish a diagnosis. Therefore, this subgroup probably also had BHR.

Furthermore, ICS were not effective in both Boulet 1994 and Pizzichini 1999, where people with baseline BHR were excluded, and in Ponsioen 2005, where only four people with a positive bronchial provocation test were included (4/30), three of whom had only mild BHR. In contrast, three studies that included participants with BHR found no effect (Evald 1989; Ponsioen 2005; Rytilä 2008). BHR was identified in 30% of participants (3/10 tested) in Evald 1989, 36% of participants (34/94 tested) in Ponsioen 2005, and 50% (35/70 tested) of participants in Rytilä 2008. Failure to identify a response in these studies could be due to other confounding factors, for example low dose of ICS (Evald 1989; Rytilä 2008). Also, given that the methacholine inhalation challenge has been shown to be falsely positive in 22% of people with chronic cough (Irwin 1990), it is possible that these studies included people with BHR, but without CVA.

Two studies specifically investigated the relationship between baseline BHR and response to ICS, with neither finding a correlation (Ribeiro 2007; Rytilä 2008). This, however, was assessed in terms of total symptom score (not necessarily cough) in Rytilä 2008. Ribeiro 2007 noted that BHR increased the odds of resolution of cough with ICS treatment (OR 9.8, 95% CI 1.09 to 88.23). The trialists attributed the higher number of participants with both BHR and response to ICS than in other studies to different study eligibility criteria and the high dose of ICS used.

It should be recognised that the absence of BHR does not exclude a steroid-responsive cough (Morice 2006), and this is consistent with the finding that improvement in symptom scores was not always restricted to people with BHR (Rytilä 2008). For example, these people may have had eosinophilic bronchitis which is responsive to ICS but does not involve BHR (Irwin 2006a).

**Airway inflammation**

Differences in airway inflammation among participants may have contributed to inconsistencies in responses to ICS. While there was no clear evidence as to whether sputum eosinophilia improved...
response, participants without sputum eosinophilia showed a poor response.
In other airway inflammatory conditions, sputum eosinophilia seems to predict steroid responsiveness (Chanez 1997; Kitaguchi 2012; Pizzichini 1998), whereas participants with neutrophilic inflammation may be less likely to respond (Green 2002; Pavord 1999). Previous studies have noted a relationship between eosinophilia and steroid responsiveness among people with chronic cough (Gibson 1989; Gibson 1998), and symptoms suggestive of asthma (Rytilä 2000). Further, ICS have been shown to reduce eosinophilic airway inflammation in eosinophilic bronchitis (Gibson 1995; Xu 2011).

Among the included studies that performed baseline sputum analysis, populations without sputum eosinophilia showed a poor response to ICS. Boulet 1994 observed no response among participants with predominantly mononuclear inflammation. Non-eosinophilic, predominantly neutrophilic airway inflammation was suggested as a cause for the very modest response achieved in Chaudhuri 2004 and the lack of effect in Pizzichini 1999. ICS did not significantly reduce sputum eosinophil counts in either study, whereas ECP, an indirect marker of eosinophil activity, was reduced in Chaudhuri 2004, but not in Pizzichini 1999. Rytilä 2008 also found no significant improvements in cough scores among a group of participants where approximately one-fifth had sputum eosinophilia, however, the authors suggested that higher rates of eosinophilic inflammation may have been detected with repeated sputum measurements. While Evald 1989 did not investigate sputum inflammation, participants in this trial did not respond to ICS and had a low incidence of blood eosinophilia, which is a less sensitive marker of airway eosinophilia (Pizzichini 1997).

Despite this, no individual included study found that sputum eosinophilia predicted the response to ICS. Rytilä 2008 found no correlation between participants with baseline eosinophilia and a response to ICS. In one report of a subset of included patients, Chaudhuri 2004 found that neither sputum eosinophils nor neutrophils predicted ICS responsiveness, however, a normal sputum eosinophil count was associated with a poorer response. While bronchial biopsies were not histologically different between the steroid responders and non-responders in Boulet 1994, the authors suggested that insufficient dose or duration of treatment may have limited the trial’s ability to detect an effect. ICS showed mixed effects on sputum eosinophils and ECP among included studies, and this was perhaps also due to differences in inflammatory patterns of participants.

Sputum inflammation was not investigated in three studies (Ponsioen 2005; Pornsuriyasak 2005; Ribeiro 2007).

**Smoking**

There was no clear trend in response to ICS among studies that did and did not exclude smokers; however, other evidence suggests that smoking reduces efficacy of ICS in cough. For example, ICS were more effective in non-smokers than smokers when directly compared in Ponsioen 2005, with the authors suggesting that this was because non-smokers have a greater baseline cough-reflex sensitivity in comparison to smokers (Dicpinigaitis 2003). Furthermore, studies have shown that ICS are ineffective in reducing smoking-related neutrophilic airway inflammation (Cox 1999).

**Recent respiratory infection**

ICS may not be effective for the treatment of post-URTI cough. Both studies that found ICS to be effective excluded people with recent respiratory infection (Chaudhuri 2004; Ribeiro 2007). Further, ICS were not effective in the one study that specifically examined post-URTI cough (Pornsuriyasak 2005), nor in the other study of predominantly subacute cough (Ponsioen 2005), however, this perceived relationship may have been confounded by other factors.

**Recent steroid use**

The impact of recent steroid use on efficacy of ICS remains uncertain. People with recent steroid use were excluded from both of the studies that found ICS to be successful (Chaudhuri 2004; Ribeiro 2007). In the studies that did not exclude people with previous steroid use, ICS were not effective (Pornsuriyasak 2005; Rytilä 2008). This result may, however, have been confounded by other study design factors. In contrast, several studies that excluded people with recent steroid use also found no effect (Boulet 1994; Evald 1989; Pizzichini 1999). It was unclear whether these people were excluded from the Ponsioen 2005 trial, which did not find ICS to be beneficial.

Several authors specifically investigated this relationship. Chaudhuri 2004 excluded people with inhaled or oral steroid use in the previous three weeks, but found no difference in response among participants who had received a prior course of ICS. Ribeiro 2007 also noted that participants who had previously received corticosteroids showed no significant difference in response to ICS. In an earlier report, however, Ribeiro 2007 noted that “drug use in recent weeks or months” was significantly lower in responders than non-responders to ICS, although whether this related to steroid use was not clear.

**Medications causing cough**

There were no clear trends in the results of studies that did and did not exclude people taking medications potentially causing cough.

**Extrapulmonary causes of cough**

There were no notable differences in the efficacy of ICS amongst trials that did and did not exclude cough potentially attributable to extrapulmonary causes such as GORD and PND.

**Interventions**

Variation in the interventions used may also have contributed to the observed inconsistencies between studies.
Dose

Both of the studies that showed a significant treatment effect used a budesonide equivalent dose of at least 1200 µg/day (Chaudhuri 2004; Ribeiro 2007), whereas ICS was not effective in reducing cough scores in either of the studies that used a low dose (Evald 1989; Rytilä 2008). In a recent Cochrane systematic review of ICS for non-specific chronic cough in children, Tomerak 2009 also found a significant improvement in the one study that used high dose ICS, but not in the study using low dose ICS. The apparent reduced efficacy of low dose ICS, however, may be an artefact of confounding factors. For instance, Evald 1989 used only a short duration of treatment, and participants in Rytilä 2008 were required to have an additional symptom in addition to cough. Furthermore, Ponsioen 2005 described no dose-effect relationship.

Treatment duration

While longer duration of treatment may increase the observed efficacy of ICS, the two studies that found ICS to be effective used a treatment period of only two weeks (Chaudhuri 2004; Ribeiro 2007). Other studies using high dose ICS for four weeks did not necessarily produce a response to ICS (Boulet 1994; Pornsuriyasak 2005).

Dosage regimen

Twice-daily dosing was most common, and was used by four studies that found no treatment effect (Evald 1989; Pizzichini 1999; Ponsioen 2005; Pornsuriyasak 2005), and one of the studies that found ICS to be effective (Chaudhuri 2004). The second study that identified a treatment effect used thrice (three times) daily dosing (Ribeiro 2007). Once-daily dosing was not effective in reducing cough scores in Rytilä 2008. Studies of the efficacy of once-daily dosing of ICS in asthma have given conflicting results (Boulet 2004). Dosing four-times daily did not produce a treatment effect (Boulet 1994), perhaps because of the inverse relationship between dose frequency and compliance (Boulet 2004; Claxton 2001). Compliance was reasonably high in studies that used twice-daily and once-daily dosing (Ponsioen 2005; Rytilä 2008). Studies that found a treatment effect used two or six puffs a day (Chaudhuri 2004; Ribeiro 2007). Studies with eight puffs a day were not effective (Boulet 1994; Evald 1989; Pornsuriyasak 2005). This may also have been related to compliance or inhalation technique.

Type of ICS

While differences in pharmacodynamic and pharmacokinetic properties of different types of ICS can lead to differences in efficacy and safety (Derendorf 2006), there were no clear differences between the types of ICS used in studies that did, and did not, demonstrate a significant treatment effect.

Administration

Although the use of different inhalation devices can have implications for inhalation technique and compliance, and in turn efficacy (Barnes 1998), the type of inhalation device used did not seem to influence whether or not a treatment effect was observed.

Outcomes

The use of different outcome measures between studies also limited the validity of comparing the studies, and may have confounded the observed relationships between ICS and change in cough severity. No study reported objective cough severity measures, which is common in cough treatment research (Leconte 2011). While all studies used some form of cough score assessed by the participant, these subjective scores have largely not been validated (Leconte 2011), as recognised in one report (Pornsuriyasak 2005). Only one study report described validation of the cough score used against objective measures of cough frequency and intensity (Ponsioen 2005). Additionally, the types of cough scores used lack consistency between trials, thereby limiting the generalisability of results. For these reasons, the clinical importance of a reduction in a subjective cough severity measure that has not been validated, or used in other studies, is unclear.

Overall completeness and applicability of evidence

While we identified many relevant, high quality RCTs of ICS for subacute and chronic cough, our ability to address all primary and secondary outcomes was limited by differences in study design and reporting. Variation in eligibility criteria, interventions and outcome measures made pooling of data inappropriate for several outcomes where data were available. Additionally, data pertaining to the primary outcome were available for only three studies, and most secondary outcomes were assessed by only a few studies. For these reasons, a narrative review of findings, rather than statistical pooling and meta-analysis, was necessary for most outcomes. Conversely, this substantial heterogeneity probably increases the external validity of the conclusions drawn. The wide variety of people who participated in these studies most probably reflects the diversity of people with this common, and inherently difficult to define, clinical problem. Similarly, the varied interventions used probably replicate variations in clinical practice. Additionally, while data for some of the outcomes were limited, a range of clinically important outcomes were assessed across the studies. Therefore, it is likely that all relevant types of participants, interventions and outcomes have been investigated.

Quality of the evidence

We assessed the effects of ICS in people with subacute and chronic cough through narrative review and limited meta-analysis of eight
studies including 570 participants. The robustness of the conclusions we could draw was restricted by methodological limitations, including the cross-over design of one trial, heterogeneity in study characteristics, and inadequate reporting of data. These factors limited statistical pooling of data, and probably also contributed to the inconsistencies in study findings, with two trials finding ICS to be beneficial, and six trials finding ICS to be largely ineffective. While comparing the number of positive studies with the number of negative studies through narrative review allowed us to determine whether there is any evidence for an effect of ICS, accurate quantification of the extent and direction of effect was not possible with the limited data available. The quality of evidence for each pooled outcome measure was determined using GRADE criteria. Narrative evidence for the primary outcome was of low quality, with evidence downgraded on the basis of the unclear risk of selection bias (Boulet 1994) and reporting bias (Ponsioen 2005; Ribeiro 2007) in included studies, and the small number of recorded events. Evidence for the finding that ICS increased the proportion achieving a greater than 50% improvement in cough severity measure was downgraded to low quality due to the unclear risk of selection bias in one included study (Boulet 1994) and the small number of recorded events. The unclear risk of selection (Boulet 1994) and reporting bias in included studies (Ponsioen 2005; Ribeiro 2007; Ryttilä 2008), and the small number of events for the proportion of participants with clinical cure at follow up resulted in a low quality grading for this narrative evidence. Evidence for mean change in cough score was deemed to be low quality because of unclear risk of selection bias (Boulet 1994; Porksnuriyasak 2005) and other bias (Pornsuriyasak 2005) in included studies, and a total population size that was less than 400. Moderate quality evidence was used to assess the proportion that experienced adverse effects of treatment, with evidence quality downgraded due to the wide confidence interval.

Potential biases in the review process

Several methodological strengths minimised the risk of bias in the review process. Explicit methodology was defined a priori in a published protocol. Comprehensive, systematic search strategies and independent study selection by two authors maximised the likelihood of identifying all relevant studies. Independent data extraction by two authors also reduced the risk of error in data collection.

Limitations of the review that may have introduced bias arose from inconsistent reporting of data in the trial reports. Despite attempts to contact study authors, not all the required data could be obtained. Inconsistent reporting and missing data limited our ability to pool data through meta-analysis for the majority of outcomes. Hence, it is possible that the effects of ICS could have been underestimated, since the results of individual studies that lacked statistical power could not be combined. Narrative review in the place of statistical pooling of data can introduce bias where statistical significance is used to define studies showing an effect, and does not take into account the weighting of studies (Higgins 2011). This post hoc approach was also a limitation of subgroup analysis, for which we were unable to perform tests for interaction.

Agreements and disagreements with other studies or reviews

This is the first systematic review of ICS for subacute and chronic cough in adults. While there are numerous narrative reviews of management strategies for subacute and chronic cough in general, many of these focus primarily on cause-directed treatment (e.g. Chummun 2011), or antitussive therapy (e.g. Bolser 2010). Previous reviews of ICS for cough have been largely limited to people with CVA (Antoniu 2007; Cazzola 2008). A recent systematic review of pharmacological and non-pharmacological interventions for cough in adults identified four of the same trials as our review (Chaudhuri 2004; Evald 1989; Molassiotis 2010; Pizzichini 1999; Ribeiro 2007), but did not draw specific conclusions about ICS for adults with subacute and chronic cough. This Cochrane review provides weak evidence in support of some current clinical guidelines. These guidelines recommend the systematic exclusion and treatment of common causes of subacute and chronic cough, which, in some instances, includes treatment with ICS.

Subacute cough is most often attributed to postinfectious cough, which can be treated with the inhaled anticholinergic drug ipratropium or failing this, ICS (Irwin 2006a). While no clear conclusions could be drawn, the results of this review suggest that ICS may not be beneficial in these people, especially given that one study that specifically examined post-URTI cough showed that ICS were not beneficial (Pornsuriyasak 2005). Subacute cough in the absence of an obvious infectious cause is managed in the same way as chronic cough (Irwin 2006a). After examination, chest X-ray and spirometry, people with chronic cough with no identifiable cause should undergo investigation and treatment for the most common causes of chronic cough - asthma, PND and GORD (Irwin 2006a). ICS are specifically indicated in several instances.

Current Australian cough guidelines recommend that adults with non-specific or refractory cough receive an empirical trial of ICS therapy (Gibson 2010). The strength of this recommendation is classified as strong according to the GRADE approach, which considers not only quality of evidence but also balance between desirable and undesirable effects, values and preferences and costs (Guyatt 2008), however, no studies are specifically cited in these guidelines. Given that ICS seem to benefit some people, the findings of our review provide weak evidence in support of this recommendation.

Several international guidelines also recommend ICS for CVA (Irwin 2006a; Morice 2004; Morice 2006), and people with a positive bronchial provocation test (Morice 2006). Our review
identified only one study that specifically addressed CVA, which noted a significant mean reduction in VAS score of 1.4 cm (95% CI -0.0 to 2.7) after two weeks (Chaudhuri 2004). Given that all the studies that showed ICS to be effective included participants with baseline BHR, it is possible that many participants who responded did have CVA.

ICS are also recommended for non-asthmatic eosinophilic bronchitis (Irwin 2006a; Morice 2004; Morice 2006), and atopic cough (Morice 2006). No randomised controlled trials that specifically examined these conditions met the inclusion criteria for this review, and subgroup analysis could not be performed for the small number of participants with eosinophilic bronchitis who were identified in one study (Chaudhuri 2004).

AUTHORS’ CONCLUSIONS

Implications for practice

This Cochrane review provides the first comprehensive systematic review of the evidence for inhaled corticosteroids (ICS) for subacute and chronic cough in adults. Overall, the studies were highly heterogeneous and the results were inconsistent. One parallel group trial of chronic cough which identified a significant treatment effect contributed the majority of statistical heterogeneity for several outcomes. The factors that predict response to ICS could not be fully determined. A trial of ICS should only be considered in adults after thorough work-up including chest X-ray and consideration of spirometry and other appropriate investigations, in accordance with international cough guidelines (Gibson 2010; Irwin 2006a; Morice 2004; Morice 2006).

Implications for research

This systematic review has demonstrated the need for greater consistency in study design, participants, interventions, outcome measures and reporting in future trials of ICS for subacute and chronic cough in adults.

Study design

Cross-over trials should not be used because of the risk of a significant carry-over effect. Assessing time taken to achieve clinical cure or significant improvement as well as factors that predict response to treatment would also be useful. This should include further evaluation of eNO, which has shown promise as a surrogate marker for airway eosinophilia and steroid responsive cough (Taylor 2006). A follow up period sufficient to investigate possible carry-over effects, as well as cough recurrence, after cessation of ICS therapy has also been recommended previously (Chang 2011).

Participants

More clearly defined study populations are required for future studies (Chang 2011). Consistency in eligibility criteria is required to minimise heterogeneity in study populations and, in turn, to allow more meaningful comparisons to be made (Molassiotis 2010). This could include using standardised guidelines requiring history and examination, spirometry and chest X-ray to exclude people with more readily identifiable and treatable causes in order to focus on those with unexplained cough. It would be most useful for this to reflect diagnostic protocols used in clinical practice in order to select a clinically relevant patient population. Medications causing cough and untreated extrapulmonary disease (e.g. gastrooesophageal reflux disease, postnasal drip) should be excluded as causes for cough. Smoking status, recent steroid use and recent respiratory infection should also be recorded. Determining the final cause for cough, as done in some included studies (Boulet 1994; Chaudhuri 2004), would also help elucidate the people most likely to respond to ICS.

Interventions

The results of this review suggest that high dose ICS for two weeks may be an appropriate intervention where a trial of ICS is considered, however, further investigation of the optimal duration of therapy would be useful (Chang 2011). Prolonged duration of ICS treatment for more than eight weeks could be studied. Using comparable interventions across trials would also facilitate more meaningful pooling of study data.

Outcomes

Consistency in cough severity measures is integral for allowing comparison of results between studies. Cough severity should be the primary outcome, and should be assessed by validated subjective and objective measures (Chang 2011; Molassiotis 2010). While objective cough measures provide evidence of frequency and intensity, subjective cough scores better reflect participants’ perceptions of the cough and quality of life. Assessment of both types of outcome measures would, therefore, be very useful in determining whether ICS treatment yields clinically important outcomes (Chang 2011; Leconte 2011; Molassiotis 2010). A recent systematic review on cough assessment endorsed the Leicester Cough Questionnaire (LCQ) and Cough Quality of Life Questionnaire (CQLQ) instruments as validated quality of life scores (Leconte 2011). Visual analogue scales are also useful for assessing people’s perception of cough severity, despite not necessarily correlating with either objective or quality-of-life scores. While cough frequency can also be used to monitor treatment response, further validation of available techniques has been recommended (Leconte 2011). Determining the minimal clinically important difference in these outcome measures is also integral to defining the practical benefit of any identified treatment effects.

Reporting

Inconsistent reporting of outcomes needs be addressed in future research. Thresholds for defining clinically significant improve-
ments in cough severity measurements (e.g., improvement of more than 70%) should also be determined a priori and reported (Chang 2011). For continuous outcomes, reporting data, even if they are not statistically significant, would be useful for future comparison between studies, as would including data for all time points measured. Ideally, dichotomous outcomes should be reported as the number of events, non-events and participants in each group; and continuous outcomes should be reported in terms of the mean difference, standard deviation and number of participants for each group (Higgins 2011).

ACKNOWLEDGEMENTS

We thank Dr Chris Cates, Dr Emma Welsh and Elizabeth Stovold from the Cochrane Airways Group for their invaluable advice and support in writing this review. We thank Elizabeth Stovold for performing the electronic database searches. We also thank Dr Wim Hop (Ponsioen 2005), and Dr Marcos Ribeiro (Ribeiro 2007), for providing additional study information and data. KJ was supported by a Cochrane scholarship from the Australian Satellite of the Cochrane Airways Group and Asthma Foundation of Queensland. IY was supported by an NHMRC Career Development Fellowship. KF and AC were supported by NHMRC Practitioner Fellowships.

REFERENCES

References to studies included in this review

Boulet 1994 [published data only]


Chaudhuri 2004 [published data only]


Evald 1989 [published data only]


Pizzichini 1999 [published data only]


Ponsioen 2005 [published and unpublished data]


Pornsuriyasak 2005 [published data only]

Inhaled corticosteroids for subacute and chronic cough in adults (Review)

References to studies excluded from this review

Brightling 2000 [published data only]

Cheriyan 1994 [published data only]

Fujimoto 2003 [published data only]

Park 2007 [published data only]

Rytilä 2000 [published data only]

Stankovic 2010 [published data only]

Stankovic 2004 [published data only]

Wei 2011 [published data only]

Xu 2011 [published data only]

References to studies awaiting assessment

Tagaya 2009 [published data only]


Tagaya 2011 [published data only]
Additional references

Anderson-James 2013

Antoniu 2007

Barnes 1998

Barnes 2006

Birring 2011

Bolser 2010

Boulet 2004

Britt 2011

Canning 2007

Cates 2008

Cazzola 2008

Chanez 1997

Chang 2005

Chang 2011

Chummun 2011

Claxton 2001

Cox 1999

Derendorf 2006

Dicpinigaitis 2003

Frank 1985

French 1998

French 2002

French 2004

Gibson 1989

Gibson 1995
Gibson PG, Hargrave FE, Girgis-Gabardo A, Morris M, Denburg JA, Dolovich J. Chronic cough with eosinophilic

Gibson 1998

Gibson 2002

Gibson 2010

GRADeprofiler 3.2

Green 2002

Guyatt 2008

Haque 2005

Higgins 2011

Howden 2010

Irwin 1990

Irwin 1998

Irwin 2006a

Irwin 2006b

Jatakanon 1999

Johnstone 2011

Kitaguchi 2012

Kwon 2006

Leconte 2011

Lee 2001

Levine 2008

Lipworth 1999

Madison 2010

McGarvey 1998a
McGarvey 1998b

McGarvey 1999

McGarvey 2006

Molassiotis 2010

Morice 2004

Morice 2006

Morice 2007

Morice 2010

Nair 2010

Niimi 1998

O’Connell 1994

Pavord 1999

Pizzichini 1997
Pizzichini E, Pizzichini MM, Efthimiadis A, Dolovich J, Hargreave FE. Measuring airway inflammation in asthma: eosinophils and eosinophilic cationic protein in induced sputum compared with peripheral blood. *Journal of Allergy and Clinical Immunology* 1997;99(4):539–44.

Pizzichini 1998

Poe 1989

Review Manager 5

Roland 2004

Taylor 2006

Tomerak 2009

Yang 2012

* Indicates the major publication for the study
### Characteristics of included studies  
*ordered by study ID*

**Boulet 1994**

| Methods | Design: cross-over, no washout, first period data available.  
Randomisation: yes, method not reported.  
Blinding: double blind.  
Withdrawals: none. |
|----------|---------------------------------------------------------------|
| Participants | Setting: single-centre study, Laval Hospital clinic (Canada)  
Number screened: not reported.  
Number eligible: 19.  
Number randomised: 14.  
Number in treatment group: 14 (cross-over).  
Number in control group: 14 (cross-over).  
Number of withdrawals: 0.  
Number completing trial: 14.  
Sex: 4 M, 15 F (eligible population).  
Age: 25–58 years (eligible population).  
Cough duration: mean 3.8 years.  
Inclusion criteria: dry cough for > 4 weeks, normal airway response to methacholine \( (PC_{20}) \), the provocative concentration of methacholine inducing a 20% decrease in \( FEV_1 \) being 20 mg/mL or more), normal chest examination and radiograph  
Exclusion criteria: use of a medication known to induce chronic cough (e.g. angiotensin-converting enzyme (ACE) inhibitors, beta-blockers), inhaled or oral steroid intake within the preceding 3 months, evidence of respiratory infection in the previous 4 weeks, smoking within the preceding 2 years, past or present history of asthma, chronic bronchitis, any other chest or systemic disease  
Baseline characteristics of treatment: control groups: comparable (cross-over) |
| Interventions | ICS: beclomethasone dipropionate 500 \( \mu g \), 4 times daily (2000 \( \mu g/d \) day)  
Control: placebo.  
Administration method: MDI with Aerochamber.  
Treatment duration: 4 weeks, no washout.  
Co-interventions: none. |
| Outcomes | Symptom score.  
Other respiratory symptoms.  
BHR (methacholine challenge, citric acid challenge). |
| Notes | Main outcome: Mean daily cough scores not significantly different between the two treatment groups |

### 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>Randomised, method not reported.</td>
</tr>
</tbody>
</table>
### Boulet 1994 (Continued)

<table>
<thead>
<tr>
<th>Source</th>
<th>Risk Assessment</th>
<th>Method</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Method not reported.</td>
<td></td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Double blind.</td>
<td>All outcomes</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Double blind.</td>
<td>All outcomes</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>No drop-outs.</td>
<td>All outcomes</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>Number eligible reported, all outcomes reported.</td>
<td></td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>First period data available.</td>
<td></td>
</tr>
</tbody>
</table>

### Chaudhuri 2004

<table>
<thead>
<tr>
<th>Source</th>
<th>Method</th>
<th>Participants</th>
<th>Interventions</th>
</tr>
</thead>
</table>
### Outcomes

Co-interventions: none.

<table>
<thead>
<tr>
<th>Outcomes</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>VAS (cough severity).</td>
<td></td>
</tr>
<tr>
<td>Sputum total and differential cell counts.</td>
<td></td>
</tr>
<tr>
<td>Eosinophilic cationic protein (ECP).</td>
<td></td>
</tr>
<tr>
<td>Myeloperoxidase (MPO).</td>
<td></td>
</tr>
<tr>
<td>Prostaglandin E₂ (PGE₂).</td>
<td></td>
</tr>
<tr>
<td>Leukotriene B₄ (LTB₄).</td>
<td></td>
</tr>
<tr>
<td>Cys-leukotrienes (Cys-LT).</td>
<td></td>
</tr>
<tr>
<td>Interleukin-8 (IL-8).</td>
<td></td>
</tr>
<tr>
<td>Tumour necrosis factor-alpha (TNF-α).</td>
<td></td>
</tr>
<tr>
<td>Exhaled nitric oxide (eNO).</td>
<td></td>
</tr>
<tr>
<td>Carbon monoxide (CO).</td>
<td></td>
</tr>
</tbody>
</table>

### Notes

Main outcome: cough severity and sputum ECP levels were modestly reduced by ICS. Other sputum biomarker levels were unaltered.

### 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>Computer-generated randomisation.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Randomisation code withheld from investigators until completion of the study</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Double blind, study medication packed by central pharmacy and was identical in appearance and taste</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>Five withdrawals with reasons reported, treatment received by drop-outs not reported</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>Number screened and eligible reported, reasons for withdrawal reported, all outcomes reported</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No carry-over effect observed despite cross-over trial design with 2 weeks washout</td>
</tr>
</tbody>
</table>
**Evald 1989**

### Methods

- **Design:** cross-over, no washout, first period data available.
- **Randomisation:** method not reported.
- **Blinding:** double blind.
- **Withdrawals:** stated.

### Participants

- **Setting:** Bispebjerg Hospital chest clinic (Denmark).
- **Number screened:** not reported.
- **Number eligible:** not reported.
- **Number randomised:** 40.
- **Number in treatment group:** N/A (cross-over).
- **Number in control group:** N/A (cross-over).
- **Number of withdrawals:** 13 (7 run-in, 2 first period, 4 second period)
- **Number completing trial:** 31 first period, 27 second period.
- **Sex:** 7 M, 24 F.
- **Age:** 15-64 years.
- **Cough duration:** 3 < 15 days; 6 > 1 month; 10 > 3 months; 9 > 1 year 3 > 5 years
- **Inclusion criteria:** daily dry cough of at least 1 hour duration in more than half of the last 30 days
- **Exclusion criteria:** obstructive lung function (ratio between the FEV<sub>1</sub> and FVC no more than 70%), significant reversibility after bronchodilating treatment (increase in FEV<sub>1</sub> 30 min after three inhalations of salbutamol 300 µg and three inhalations ipratropium 60 µg not exceeding 20% or 500 mL), diurnal variation above 20% in morning and evening PEF during home monitoring for 1 week (calculated at visit 2 after the run-in period), treatment with anti-asthmatic drugs, abnormal chest X-ray, recent respiratory infection, other chest disease, pregnancy
- **Baseline characteristics of treatment:control groups:** comparable (cross-over)

### Interventions

- **ICS:** beclomethasone dipropionate 50 µg 4 puffs twice daily (400 µg/day)
- **Control:** placebo.
- **Administration method:** MDI.
- **Treatment duration:** 2 weeks, no washout.
- **Co-interventions:** none.

### Outcomes

- **Patient's subjective effect of treatment (score).**
- **Degree of cough (including number of days with cough, number of cough attacks a day).**
- **Duration of cough attacks estimated for the whole day.**
- **Spirometry.**
- **PEF.**
- **Number of awakenings each night because of cough.**
- **Duration of insomnia at night because of cough.**

### Notes

- **Main outcome:** no significant treatment effect found for any of the measured variables
- **Baseline investigations:** spirometry, bronchodilator reversibility, bloods, skin prick test, bronchial provocation test
- **Run-in period:** 1 week

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>

---

Inhaled corticosteroids for subacute and chronic cough in adults (Review)  
Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
Evald 1989  (Continued)

<table>
<thead>
<tr>
<th>Evaluation Area</th>
<th>Risk of Bias</th>
<th>Reporting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Randomised, method not reported.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Method not reported.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>2 withdrawals during period one, 4 during period two; treatment received by drop-outs not reported</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Number screened and eligible not reported, scores for participants’ subjective effect of treatment not reported, reasons for withdrawal reported</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>First period results described separately; run-in period.</td>
</tr>
</tbody>
</table>

Pizzichini 1999

| Methods | Design: parallel group.  
|         | Randomisation: yes, computer-generated.  
|         | Blinding: double blind.  
|         | Withdrawals: stated.   |
| Participants | Setting: Firestone Regional Chest and Allergy Clinic and recruited by advertisement (Canada)  
|         | Number screened: 84.  
|         | Number eligible: 50.  
|         | Number randomised: 50.  
|         | Number in treatment group: 21 (completed).  
|         | Number in control group: 23 (completed).  
|         | Number of withdrawals (treatment:control): 6 (4:2).  
|         | Number completing trial (treatment:control): 44 (21:23).  
|         | Sex: 16 M, 28 F (completed).  
|         | Age: 20-75 years.  
|         | Cough duration: 6-19.2 years (completed).  
|         | Inclusion criteria: daily bothersome cough for at least 1 year with no other respiratory symptoms, no evidence of asthma (FEV₁ > 70% predicted, FEV₁/FVC > 70%, little response to a bronchodilator), normal methacholine airway responsiveness (a provocation concentration of methacholine to cause a fall in FEV₁ of 20% (PC₂₀) of > 8 mg/mL) |
People with symptomatic gastroesophageal reflux not improved with treatment and PND who had been previously investigated and treated, and who had no sinusitis on sinus X-rays were included.

Exclusion criteria: smokers, ex-smokers of 6 months or less, indication of respiratory infection in previous month, history of chronic bronchitis, radiological evidence of chest disease, other recognised condition or drugs to account for the cough, cardiovascular or renal disease requiring regular medication, pregnant, received corticosteroids within the month.

Baseline characteristics of treatment: control groups: comparable.

## Interventions

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>ICS:</td>
<td>budesonide 400 µg twice daily (800 µg/day).</td>
</tr>
<tr>
<td>Control:</td>
<td>placebo.</td>
</tr>
<tr>
<td>Administration method:</td>
<td>DPI (Turbuhaler).</td>
</tr>
<tr>
<td>Treatment duration:</td>
<td>2 weeks.</td>
</tr>
<tr>
<td>Co-interventions:</td>
<td>none.</td>
</tr>
</tbody>
</table>

## Outcomes

|                                | Questionnaire of cough frequency, cough discomfort (9-point Likert scale) |
|                                | VAS (cough discomfort in the previous two days).                         |
|                                | Spirometry.                                                              |
|                                | Sputum total and differential cell counts, ECP, IL-8, fibrinogen, albumin, substance P |
|                                | Local side effects (structured questionnaire, oropharyngeal inspection) |

## Notes

Main outcome: treatment did not affect cough or sputum measurements, perhaps because the cause was not associated with sputum eosinophilia. Baseline investigations: bloods, allergy skin prick tests. Study period followed by 2 weeks of open label budesonide.

## 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>Computer-generated randomisation.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Randomisation generated off-site, concealed from investigators, administered by research nurse</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Double blind, identical placebo.</td>
</tr>
<tr>
<td>All outcomes</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Similar withdrawal rates for treatment and control groups.</td>
</tr>
<tr>
<td>All outcomes</td>
<td>Low risk</td>
<td></td>
</tr>
</tbody>
</table>
### Selective reporting (reporting bias)

<table>
<thead>
<tr>
<th>Study</th>
<th>Risk</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pizzichini 1999</td>
<td>Unclear</td>
<td>Several outcomes not reported (change in cough questionnaire, BHR, spirometry, albumin), number screened and eligible reported, reasons for withdrawals stated</td>
</tr>
</tbody>
</table>

### Other bias

<table>
<thead>
<tr>
<th>Study</th>
<th>Risk</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pizzichini 1999</td>
<td>Low</td>
<td>Parallel group trial.</td>
</tr>
</tbody>
</table>

### Ponsioen 2005

#### Methods

Design: parallel group.
Randomisation: yes, computer-generated.
Blinding: double blind.
Withdrawals: 2.

#### Participants

Setting: community-based primary healthcare centre (6 practices; The Netherlands)
Number screened: 162.
Number eligible: 135.
Number randomised: 135.

**Total study population (published data):**
- Number in treatment group: 67 randomised (GSK report), 65 analysed
- Number in control group: 68 randomised (GSK report), 68 analysed
- Number of withdrawals (treatment:control): 2 (2:0).
- Number completing trial (treatment:control): 133 (65:68).
- Sex: 47 M, 86 F (analysed population).
- Age (years): mean 47.0 (SD 10.1) treatment, mean 43.4 (SD 11.2) control
- Cough duration (weeks): mean 4.2 (SD 2.5) treatment, mean 5 (SD 3.7) control; 31 acute cough (< 3 weeks), 89 subacute cough (3-8 weeks), 13 chronic cough (8-17 weeks) (analysed population)

**Subacute and chronic cough participants only (unpublished data):**
- Number in treatment group: 52.
- Number in control group: 50.
- Number of withdrawals (treatment:control): 0.
- Number completing trial (treatment:control): 102 (52:50).
- Sex: 34 M, 68 F.
- Age (years): mean 46.7 (SD 10.5) treatment, mean 44.5 (SD 11.2) control
- Cough duration (weeks): mean 4.8 (SD 2.5) treatment, mean 6.1 (SD 3.7) control

Inclusion criteria: aged 18-65 years with cough of ≥ 2 weeks duration. Participants completed a daily diary card for cough (score 0 = absent, 1 = mild, 2 = moderate, 3 = severe) and other LRT symptoms regarding the previous day and night. Only people with a night score of ≥1 point and a combined day plus night score of ≥3 points were included.

Exclusion criteria: history of asthma; incidences of self-reported wheeze, pharmacy data indicating asthma-like symptoms or variability in lung function in the previous year; current treatment that might influence the cough; FEV<sub>1</sub> < 60% predicted; any concurrent airway disease (e.g. pneumonia, cancer, tuberculosis, tonsillitis, sinusitis); uncontrolled systemic disease or pregnancy; people previously randomised for the study

Baseline characteristics of treatment:control groups: comparable
### Interventions

ICS: fluticasone propionate 500 µg twice daily (1000 µg/day)
Control: placebo.
Administration method: MDI via Volumatic spacer.
Treatment duration: 2 weeks.
Co-interventions: no rescue medication allowed, concurrent medication for lower respiratory tract symptoms was registered in the diary

### Outcomes

Symptom score (cough, sputum production, wheezing, shortness of breath and chest tightness)
Perception of whether coughing had strongly improved, improved, not changed or increased
Spirometry (FEV₁, FVC).
BHR (histamine challenge).
Number of awakenings at night.
Number of days off work.
Requirement for additional medication after the treatment period
Hoarseness.
Other adverse events.
(Number cigarettes smoked).

### Notes

Main outcome: among the total study population, cough score decreased significantly more in the treatment group, however a favourable effect was only detectable in non-smokers. ICS was not effective after exclusion of participants with acute cough. Allergy, FEV₁ and BHR at baseline did not predict the efficacy of ICS.
Baseline investigations: bloods (CRP, Phadiatop).

### 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>Computer-generated randomisation.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Study medication provided by GlaxoSmithKline (GSK) according to randomisation list</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Double blind, matching placebo inhaler (GSK report).</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>Only 2 withdrawals (treatment group).</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Ponsioen 2005 (Continued)

<table>
<thead>
<tr>
<th>Bias Type</th>
<th>Risk Level</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>FEF&lt;sub&gt;25&lt;/sub&gt;−&lt;sub&gt;75&lt;/sub&gt;% reported in results but is not described as an outcome in methods section, number screened and eligible reported, reasons for withdrawals stated</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>Compliance measured; parallel group trial.</td>
</tr>
</tbody>
</table>

### Pornsuriyasak 2005

**Methods**

- Design: parallel group.
- Randomisation: yes, method not reported.
- Blinding: double blind.
- Withdrawals: stated.

**Participants**

- Setting: 1200-bed university hospital (Thailand).
- Number screened: not reported.
- Number eligible: not reported.
- Number randomised: 30.
- Number in treatment group: 15.
- Number in control group: 15.
- Number of withdrawals (treatment:control): 4 (1:3).
- Number completing trial (treatment:control): 26 (14:12).
- Sex: 6 M, 24 F.
- Age (years): mean 40.6 (SD 11.8) treatment, mean 38.8 (SD 13.0) control
- Cough duration (weeks): mean 5.93 (SD 1.94) treatment, mean 4.66 (SD 2.05) control
- Inclusion criteria: consenting, non-smoking adults with persistent post-URTI cough of > 3 weeks duration; aged > 15 years; normal physical examination; normal CXR; spirometry at baseline: FVC ≥ 80% predicted, FEV<sub>1</sub> ≥ 80% predicted, and FEV<sub>1</sub>/FVC (≥ 70% predicted); sinusitis treated appropriately by an Ear, Nose, Throat physician before entry into the study, if the patient had physical signs and/or an abnormal sinus radiography. People with a negative methacholine challenge test were randomised, however three people with a mildly positive bronchial provocation test were included
- Exclusion criteria: medical history suggesting asthma; symptoms suggesting gastroesophageal reflux prior to or during post-URTI coughing bouts; history of taking medication that induces coughing; contraindication to methacholine challenge testing. Initial treatment with beta<sub>2</sub> agonists (inhaled and oral), theophyllines, corticosteroids (oral), and inhaled anticholinergics were terminated at least 1 week prior to entry into the study.
- Baseline characteristics of treatment:control groups: comparable

**Interventions**

- ICS: budesonide 100 µg, 4 puffs twice daily (800 µg daily).
- Control: placebo, 4 puffs twice daily.
- Administration method: DPI.
- Treatment duration: 4 weeks.
- Co-interventions: other medications were allowed.

**Outcomes**

- Symptom score (frequency of cough, frequency of coughing bouts, symptoms associated with cough, night-time cough)
- BHR (methacholine challenge test)
<table>
<thead>
<tr>
<th>Notes</th>
<th>Main outcome: ICS ineffective in treating persistent post-URT I cough in previously healthy individuals</th>
</tr>
</thead>
</table>

### 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>Randomised, method not reported.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Method not reported.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>Low risk</td>
<td>Similar attrition rates for treatment and control groups.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Number of participants screened and eligible not reported, reasons for withdrawals stated, all outcomes reported</td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>Post-URT I cough not defined; parallel group trial.</td>
</tr>
</tbody>
</table>

### Ribeiro 2007

<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>Setting: Outpatient Respiratory Clinic of the Hospital São Paulo, general practitioners and hospital clinics (Brazil) Number screened: 147. Number eligible: 64. Number randomised: 64. Number in treatment group: 44. Number in control group: 20.</td>
</tr>
</tbody>
</table>
Number of withdrawals (treatment:control): 0 (0:0).
Number completing trial (treatment:control): 64 (44:20).
Sex: 22 M, 42 F.
Age (years): mean 46.4 (SD 17.4) treatment, mean 50.1 (SD 18.1) control
Cough duration (weeks): mean 48.2 (SD 99.6) treatment, mean 36.7 (SD 45.5) control
Inclusion criteria: cough for at least 8 weeks with normal chest radiograph, plain sinus radiographs in four positions, and ears, nose and throat examination
Exclusion criteria: previous gastroesophageal reflux disease diagnosis, positive 24-hour oesophageal pH measurement, concurrent respiratory tract infections, and a history or medical diagnosis of asthma, chronic obstructive pulmonary disease, or chronic rhinosinusitis (PND syndrome) and evidence of airflow limitation with a FEV₁/FVC of $\leq 70\%$, no use of medications for cough in the 4 weeks leading up to entry into the study
Baseline characteristics of treatment:control groups: PD$_{20}$ (provocation dose causing a decline in FEV₁ of 20%) greater in treatment group (5.35 mg/mL ± 3.2 mg/mL versus 4.56 mg/mL ± 3.7 mg/mL; P value 0.01), otherwise comparable

### Interventions

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Treatment</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICS: chlorofluorocarbon-beclomethasone 250 µg, 2 puffs 3 times daily (1500 µg/day)</td>
<td>Studies a change in FEV₁</td>
<td>Placebo, 2 puffs 3 times daily</td>
</tr>
<tr>
<td>Administration method: MDI</td>
<td>Control and treatment received similar packaging</td>
<td></td>
</tr>
<tr>
<td>Treatment duration: 2 weeks.</td>
<td>Study participants were double blind</td>
<td></td>
</tr>
<tr>
<td>Co-interventions: none.</td>
<td>Study participants were double blind</td>
<td></td>
</tr>
</tbody>
</table>

### Outcomes

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Symptom diary: frequency (throughout the day), severity (on arising and throughout the day), duration of coughing (on arising and throughout the day), sleep interruption (throughout the night)</th>
<th>VAS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adverse events reported by participants.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Notes

Main outcome: ICS provided an excellent response in a subgroup of participants with chronic cough that did not correlate with atopy or airway hyper-responsiveness

### 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>Simple randomisation in an unbalanced design using a random number table at a ratio of 2:1 (treatment versus control)</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Randomisation performed by an outside observer.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>Double blind, packaging of the study and placebo was identical in appearance and taste and identically marked</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Low risk</td>
<td>Double blind.</td>
</tr>
</tbody>
</table>
Incomplete outcome data (attrition bias)  | Low risk  | No withdrawals.  
Selective reporting (reporting bias)  | Unclear risk  | Adverse effects not reported as an outcome of interest in methods section but is reported in results, number screened and eligible reported, all outcomes reported  
Other bias  | Low risk  | Parallel group trial.  

**Rytilä 2008**

**Methods**
- Design: parallel group.
- Randomisation: yes, computer-generated.
- Blinding: double blind.
- Withdrawals: stated.

**Participants**
- Setting: 23 study centres in Finland, Sweden, Norway, Greece, Hungary, UK and Canada; participants were referred to specialists, who were the investigators in the study.
- Number screened: 229.
- Number eligible: not reported.
- Number randomised: 144.
- Number in treatment group: 71.
- Number in control group: 73.
- Number of withdrawals (treatment:control): 23 (10:13).
- Number completing trial (treatment:control): 121 (61:60).
- Sex: 42 M, 99 F (of 141 with sex recorded).
- Age: 20-67 years.
- Cough duration: not reported, inclusion criteria ≥ 2 months.
- Inclusion criteria: FEV₁ ≥ 80% predicted; cough (with or without sputum production) plus at least one additional symptom from chest tightness, wheezing, shortness of breath, or exercise-induced cough or wheezing for ≥ 2 months but < 2 years; average symptom score of ≥ 1 (scale 0-3) for cough and for sputum production during 7 days of the run-in period (1-2 weeks).
- Exclusion criteria: physician-diagnosed asthma; ≥ 12% increase in absolute FEV₁ during reversibility testing at screening; average daily morning/evening peak expiratory flow (PEF) variability ≥ 20% for the week prior to baseline; history of smoking within 12 months prior to screening or a smoking history > 10 pack-years; evidence of chronic obstructive pulmonary disease, chronic cough due to PND, asthma, chronic bronchitis, sinusitis or gastro-oesophageal reflux (careful medical history and radiographs of the chest and paranasal sinuses were obtained); an URTI within 4 weeks prior to screening. People with symptoms of allergic/nonallergic rhinitis were treated with nasal corticosteroids and/or antihistamines before the study; such treatment could not be changed during the study.
- Baseline characteristics of treatment:control groups: comparable.
**Interventions**

ICS: mometasone furoate 400 μg daily.  
Control: placebo.  
Administration method: DPI (Twisthaler).  
Treatment duration: 8 weeks.  
Co-interventions: salbutamol inhaler could be used as a reliever medication, no other medications were allowed

**Outcomes**

Symptom scores: cough, sputum production, wheeze, shortness of breath, chest tightness and exercise-induced cough/wheeze  
BHR (histamine or methacholine).  
PEF.  
Requirement for supplemental salbutamol use.  
Sputum eosinophils, ECP.  
Adverse events reported by patient.

**Notes**

Main outcomes: ICS improved total morning symptom scores but not total evening symptom scores. ICS improved all individual symptom scores, although this was not always statistically significant. ICS improved morning and evening PEF and reduced sputum eosinophils and ECP

**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>Computer-generated randomisation</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Randomisation code maintained in sealed envelope</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Double blind, identical-looking placebo inhaler</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>Similar withdrawal rates for treatment and control groups; reasons for withdrawals reported</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Some outcomes not reported (BHR at 8 weeks, individual symptom scores other than wheeze and cough), PEF monitoring not part of study design section but is reported in methods, number eligible not reported, number screened reported, reasons for withdrawals stated</td>
</tr>
</tbody>
</table>
Other bias  |  Low risk

Abbreviations

> = more than; < = less than; ≥ = greater than or equal to; BHR = bronchial hyper-responsiveness; CRP = CXR = ; DPI = dry powder inhaler; ECP = eosinophilic cationic protein; F = female; FEF = forced expiratory flow; FEV\textsubscript{1} = forced expiratory volume in one second; FVC = forced vital capacity; GSK = GlaxoSmithKline; ICS = inhaled corticosteroids; M = male; MDI = metered dose inhaler; N/A = not applicable; PC\textsubscript{20} = ; PEF = peak; expiratory flow; PND = postnasal drip; URTI = upper respiratory tract infection.

Characteristics of excluded studies  [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brightling 2000</td>
<td>Open label uncontrolled trial.</td>
</tr>
<tr>
<td>Cheriyan 1994</td>
<td>Retrospective uncontrolled trial.</td>
</tr>
<tr>
<td>Fujimoto 2003</td>
<td>Prospective uncontrolled trial of bronchodilator, antiallergic and inhaled or oral glucocorticoid therapy</td>
</tr>
<tr>
<td>Gillissen 2007</td>
<td>Inclusion criteria of post-infectious cough of between 3 and 14 days duration (i.e. acute cough)</td>
</tr>
<tr>
<td>Han 2009</td>
<td>Randomised trial of inhaled fluticasone versus oral codeine plus levodropropizine</td>
</tr>
<tr>
<td>Matsuoka 2010</td>
<td>Retrospective uncontrolled study.</td>
</tr>
<tr>
<td>Park 2007</td>
<td>Uncontrolled trial assessing roles of the capsaicin cough sensitivity test, methacholine bronchial provocation test and induced sputum test in evaluation of chronic nonproductive cough</td>
</tr>
<tr>
<td>Rytilä 2000</td>
<td>Inclusion criteria of at least 2 of 6 respiratory symptoms (cough, chest tightness with wheezing, shortness of breath, sputum production, wheezing or cough at exercise, and disturbed sleep), not necessarily always including cough</td>
</tr>
<tr>
<td>Stankovic 2010</td>
<td>Open-label, non-randomised trial of inhaled corticosteroid and β\textsubscript{2} agonist versus oral β\textsubscript{2} agonist.</td>
</tr>
<tr>
<td>Stankovic 2004</td>
<td>Open-label, non-randomised, uncontrolled trial of inhaled β\textsubscript{2} agonist followed by ICS treatment.</td>
</tr>
<tr>
<td>Wei 2011</td>
<td>Prospective observational study of inhaled and oral bronchodilator therapy versus inhaled budesonide and oral bronchodilator therapy for cough-variant asthma</td>
</tr>
<tr>
<td>Xu 2011</td>
<td>Comparison of 4, 8 and 16 weeks inhaled budesonide therapy for eosinophilic bronchitis; no placebo comparison</td>
</tr>
</tbody>
</table>

Inhaled corticosteroids for subacute and chronic cough in adults (Review)  
Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
**Characteristics of studies awaiting assessment** [ordered by study ID]

### Tagaya 2009

<table>
<thead>
<tr>
<th>Methods</th>
</tr>
</thead>
</table>
| Design: parallel group.  
Randomisation: yes, method not reported.  
Blinding: not reported.  
Withdrawals: not reported. |

<table>
<thead>
<tr>
<th>Participants</th>
</tr>
</thead>
</table>
| Setting: not reported (Japan).  
Number screened: not reported.  
Number eligible: not reported.  
Number randomised: 25.  
Number in treatment group: not reported.  
Number in control group: not reported.  
Number of withdrawals (treatment:control): not reported.  
Number completing trial (treatment:control): not reported.  
Sex: not reported.  
Age range: not reported.  
Cough duration: not reported.  
Inclusion criteria: CVA.  
Exclusion criteria: not reported.  
Baseline characteristics of treatment:control groups: not reported |

<table>
<thead>
<tr>
<th>Interventions</th>
</tr>
</thead>
</table>
| ICS: salmeterol/fluticasone propionate combination 50/100 µg, once daily (100 µg/day)  
Control: salmeterol 50 µg twice daily - different dose of salmeterol  
Administration method: not reported.  
Treatment duration: 12 weeks.  
Co-interventions: long acting beta-adrenoreceptor agonist (LABA) |

<table>
<thead>
<tr>
<th>Outcomes</th>
</tr>
</thead>
</table>
| Cough score.  
FEV₁.  
PEF.  
Sputum eosinophils, ECP. |

<table>
<thead>
<tr>
<th>Notes</th>
</tr>
</thead>
</table>
| Main outcome: maintenance therapy with SFC improved cough symptoms, pulmonary function and airway inflammation. Discontinuation caused worsening of the disease.  
Reported as abstract. Contacted Dr Tagaya but unpublished data not available. Awaiting publication of full paper |

### Tagaya 2011

<table>
<thead>
<tr>
<th>Methods</th>
</tr>
</thead>
</table>
| Design: parallel group.  
Randomisation: yes, method not reported.  
Blinding: not reported.  
Withdrawals: not reported. |

<table>
<thead>
<tr>
<th>Participants</th>
</tr>
</thead>
</table>
| Setting: multicentre trial (Japan).  
Number screened: not reported.  
Number eligible: not reported.  
Number randomised: 27. |

---

Inhaled corticosteroids for subacute and chronic cough in adults (Review)  
Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
Number in treatment group: 14.
Number in control group: 13.
Number of withdrawals (treatment:control): not reported.
Number completing trial (treatment:control): not reported.
Sex: not reported.
Age range: not reported.
Cough duration: not reported.
Inclusion criteria: CVA according to Japanese cough guidelines
Exclusion criteria: not reported.
Baseline characteristics of treatment:control groups: comparable

**Interventions**

ICS: budesonide/formoterol combination 160/4.5 μg twice daily (total budesonide dose 320 μg/day)
Control: salmeterol 50 μg twice daily.
Administration method: DPI.
Treatment duration: 8 weeks.
Co-interventions: LABA, supplemental procaterol (SABA).

**Outcomes**

Cough symptom score.
Cough and sputum assessment questionnaire (CASA-Q).
FEV₁.
PEF.
Supplemental use of inhaled procaterol (SABA).
Sputum eosinophils, ECP.

**Notes**

Main outcome: treatment decreased cough symptom scores, CASA-Q scores, diurnal variation of PEF, eosinophil counts and ECP.
Reported as abstract. Contacted Dr Tagaya but unpublished data not available. Awaiting publication of full paper.

**Abbreviations**

CVA = cough-variant asthma; DPI = dry powder inhaler; ECP = eosinophilic cationic protein; FEV₁ = forced expiratory volume in one second; ICS = inhaled corticosteroids; PEF = peak expiratory flow; SFC = salmeterol/fluticasone propionate combination.
## Data and Analyses

### Comparison 1. ICS 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 Proportion of participants with clinical cure or significant improvement (&gt; 70% reduction in cough severity measure) at follow up (clinical success)</td>
<td>3</td>
<td></td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>1.1 Cough score</td>
<td>3</td>
<td></td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>1.2 VAS</td>
<td>1</td>
<td></td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>2 Proportion of participants with clinical cure or &gt; 50% reduction in cough severity measure at follow up</td>
<td>4</td>
<td></td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>3 Proportion of participants with clinical cure at follow up</td>
<td>4</td>
<td></td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>4 Mean change in cough score</td>
<td>5</td>
<td>346</td>
<td>Std. Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.34 [-0.56, -0.13]</td>
</tr>
<tr>
<td>5 Mean change in cough measures on VAS</td>
<td>2</td>
<td></td>
<td>Std. Mean Difference (Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>6 Mean change in VAS after 2 weeks by final diagnosis [cm]</td>
<td>1</td>
<td></td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>6.1 PNDS</td>
<td>1</td>
<td></td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>6.2 GORD</td>
<td>1</td>
<td></td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>6.3 CVA</td>
<td>1</td>
<td></td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>6.4 Bronchiectasis</td>
<td>1</td>
<td></td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>6.5 Idiopathic cough</td>
<td>1</td>
<td></td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>7 Mean change in morning and evening cough score</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>7.1 Morning cough score 4 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>7.2 Morning cough score 8 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>7.3 Evening cough score 4 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>7.4 Evening cough score 8 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>8 Mean change in morning and evening total symptom score</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>8.1 Morning total symptom score 4 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>8.2 Morning total symptom score 8 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>8.3 Evening total symptom score 4 weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
</tbody>
</table>
### 8.4 Evening total symptom score 8 weeks

Mean Difference (IV, Fixed, 95% CI) 0.0 [0.0, 0.0]

### 9 Mean change in cough frequency

| 9.1 After 1 week | Mean Difference (IV, Fixed, 95% CI) Totals not selected |
| 9.2 After 2 weeks | Mean Difference (IV, Fixed, 95% CI) Totals not selected |

### 10 Mean change in cough severity

| 10.1 After 1 week | Mean Difference (IV, Fixed, 95% CI) Totals not selected |
| 10.2 After 2 weeks | Mean Difference (IV, Fixed, 95% CI) Totals not selected |

### 11 Mean change in cough time of day

| 11.1 After 1 week | Mean Difference (IV, Fixed, 95% CI) Totals not selected |
| 11.2 After 2 weeks | Mean Difference (IV, Fixed, 95% CI) Totals not selected |

### 12 Proportion without BHR after treatment

| 12.1 Baseline BHR | Odds Ratio (M-H, Fixed, 95% CI) 1.28 [0.38, 4.34] |
| 12.2 No baseline BHR | Odds Ratio (M-H, Fixed, 95% CI) 1.32 [0.08, 22.15] |

### 13 Mean change in BHR

| 13.1 Measured by methacholine challenge | Mean Difference (IV, Fixed, 95% CI) Totals not selected |
| 13.2 Measured by citric acid challenge | Mean Difference (IV, Fixed, 95% CI) Totals not selected |

### 14 Change in FEV₁

| 14.1 Non-smokers | Mean Difference (IV, Fixed, 95% CI) Totals not selected |
| 14.2 Smokers | Mean Difference (IV, Fixed, 95% CI) Totals not selected |

### 15 Proportion requiring additional medication

| 15.1 Non-smokers | Odds Ratio (M-H, Fixed, 95% CI) 0.31 [0.13, 0.76] |
| 15.2 Smokers | Odds Ratio (M-H, Fixed, 95% CI) 0.85 [0.27, 2.63] |

### 16 Mean change in sleep interruption

| 16.1 After 1 week | Mean Difference (IV, Fixed, 95% CI) Totals not selected |
| 16.2 After 2 weeks | Mean Difference (IV, Fixed, 95% CI) Totals not selected |

### 17 Mean change in sputum ECP (ng/mL)

| 18.1 PNDS | Mean Difference (Fixed, 95% CI) -396.0 [-791.99, -0.01] |
| 18.2 GORD | Mean Difference (Fixed, 95% CI) Totals not selected |
| 18.3 CVA | Mean Difference (Fixed, 95% CI) Totals not selected |
| 18.4 Bronchiectasis | Mean Difference (Fixed, 95% CI) Totals not selected |
| 18.5 Idiopathic cough | Mean Difference (Fixed, 95% CI) Totals not selected |

### 19 Change in sputum total cells (×10⁶)

| 19.1 PNDS | Mean Difference (Fixed, 95% CI) Subtotals only |
| 19.2 GORD | Mean Difference (Fixed, 95% CI) Subtotals only |
| 19.3 CVA | Mean Difference (Fixed, 95% CI) Subtotals only |
| 19.4 Bronchiectasis | Mean Difference (Fixed, 95% CI) Subtotals only |
| 19.5 Idiopathic cough | Mean Difference (Fixed, 95% CI) Subtotals only |
| 19.6 All causes | Mean Difference (Fixed, 95% CI) Subtotals only |

### 20 Change in sputum neutrophils [%]

| 20.1 PNDS | Mean Difference (Fixed, 95% CI) Subtotals only |

---

**Inhaled corticosteroids for subacute and chronic cough in adults (Review)**

Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
<table>
<thead>
<tr>
<th></th>
<th>Study Count</th>
<th>Effect Measure</th>
<th>Mean Difference (Fixed, 95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20.2 GORD</td>
<td>1</td>
<td>Mean Difference</td>
<td>-13.8 [-29.75, 2.15]</td>
</tr>
<tr>
<td>20.3 CVA</td>
<td>1</td>
<td>Mean Difference</td>
<td>0.9 [-19.90, 21.70]</td>
</tr>
<tr>
<td>20.4 Bronchiectasis</td>
<td>1</td>
<td>Mean Difference</td>
<td>0.6 [-20.20, 21.40]</td>
</tr>
<tr>
<td>20.5 Idiopathic cough</td>
<td>1</td>
<td>Mean Difference</td>
<td>5.3 [-14.20, 24.80]</td>
</tr>
<tr>
<td>20.6 All causes</td>
<td>1</td>
<td>Mean Difference</td>
<td>1.1 [-7.50, 9.70]</td>
</tr>
<tr>
<td>21 Change in sputum eosinophils [%]</td>
<td>1</td>
<td>Mean Difference</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>21.1 PNDS</td>
<td>1</td>
<td>Mean Difference</td>
<td>0.0 [-2.05, 2.05]</td>
</tr>
<tr>
<td>21.2 GORD</td>
<td>1</td>
<td>Mean Difference</td>
<td>-0.1 [0.00, 1.80]</td>
</tr>
<tr>
<td>21.3 CVA</td>
<td>1</td>
<td>Mean Difference</td>
<td>-4.6 [-7.10, -2.10]</td>
</tr>
<tr>
<td>21.4 Bronchiectasis</td>
<td>1</td>
<td>Mean Difference</td>
<td>1.5 [-0.95, 3.95]</td>
</tr>
<tr>
<td>21.5 Idiopathic cough</td>
<td>1</td>
<td>Mean Difference</td>
<td>0.1 [-2.20, 2.40]</td>
</tr>
<tr>
<td>21.6 All causes</td>
<td>1</td>
<td>Mean Difference</td>
<td>-0.7 [-1.75, 0.35]</td>
</tr>
<tr>
<td>22 Change in sputum lymphocytes [%]</td>
<td>1</td>
<td>Mean Difference</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>22.1 PNDS</td>
<td>1</td>
<td>Mean Difference</td>
<td>0.0 [-0.35, 0.35]</td>
</tr>
<tr>
<td>22.2 GORD</td>
<td>1</td>
<td>Mean Difference</td>
<td>0.1 [-0.25, 0.45]</td>
</tr>
<tr>
<td>22.3 CVA</td>
<td>1</td>
<td>Mean Difference</td>
<td>-0.3 [-0.70, 0.10]</td>
</tr>
<tr>
<td>22.4 Bronchiectasis</td>
<td>1</td>
<td>Mean Difference</td>
<td>-0.1 [-0.50, 0.30]</td>
</tr>
<tr>
<td>22.5 Idiopathic cough</td>
<td>1</td>
<td>Mean Difference</td>
<td>-0.1 [-0.50, 0.30]</td>
</tr>
<tr>
<td>22.6 All causes</td>
<td>1</td>
<td>Mean Difference</td>
<td>-0.1 [-0.25, 0.05]</td>
</tr>
<tr>
<td>23 Change in sputum MPO [µg/mL]</td>
<td>1</td>
<td>Mean Difference</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>23.1 PNDS</td>
<td>1</td>
<td>Mean Difference</td>
<td>-34.0 [-113.30, 45.30]</td>
</tr>
<tr>
<td>23.2 GORD</td>
<td>1</td>
<td>Mean Difference</td>
<td>-17.7 [-97.05, 61.65]</td>
</tr>
<tr>
<td>23.3 CVA</td>
<td>1</td>
<td>Mean Difference</td>
<td>70.0 [-98.30, 238.30]</td>
</tr>
<tr>
<td>23.4 Bronchiectasis</td>
<td>1</td>
<td>Mean Difference</td>
<td>133.5 [27.05, 239.95]</td>
</tr>
<tr>
<td>23.5 Idiopathic cough</td>
<td>1</td>
<td>Mean Difference</td>
<td>9.1 [-128.30, 146.50]</td>
</tr>
<tr>
<td>23.6 All causes</td>
<td>1</td>
<td>Mean Difference</td>
<td>10.7 [-29.60, 51.00]</td>
</tr>
<tr>
<td>24 Change in sputum PGE2 [ng/mL]</td>
<td>1</td>
<td>Mean Difference</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>24.1 PNDS</td>
<td>1</td>
<td>Mean Difference</td>
<td>4.9 [-10.30, 20.10]</td>
</tr>
<tr>
<td>24.2 GORD</td>
<td>1</td>
<td>Mean Difference</td>
<td>-4.7 [-20.85, 11.45]</td>
</tr>
<tr>
<td>24.3 CVA</td>
<td>1</td>
<td>Mean Difference</td>
<td>12.1 [-10.70, 34.90]</td>
</tr>
<tr>
<td>24.4 Bronchiectasis</td>
<td>1</td>
<td>Mean Difference</td>
<td>-11.6 [-30.25, 7.05]</td>
</tr>
<tr>
<td>24.5 Idiopathic cough</td>
<td>1</td>
<td>Mean Difference</td>
<td>-9.8 [-32.65, 13.05]</td>
</tr>
<tr>
<td>24.6 All causes</td>
<td>1</td>
<td>Mean Difference</td>
<td>-1.9 [-9.05, 5.25]</td>
</tr>
<tr>
<td>25 Change in sputum LTB4 [ng/mL]</td>
<td>1</td>
<td>Mean Difference</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>25.1 PNDS</td>
<td>1</td>
<td>Mean Difference</td>
<td>5.1 [-32.85, 43.05]</td>
</tr>
<tr>
<td>25.2 GORD</td>
<td>1</td>
<td>Mean Difference</td>
<td>1.70 [-36.30, 39.70]</td>
</tr>
<tr>
<td>25.3 CVA</td>
<td>1</td>
<td>Mean Difference</td>
<td>25.6 [-18.25, 69.45]</td>
</tr>
<tr>
<td>25.4 Bronchiectasis</td>
<td>1</td>
<td>Mean Difference</td>
<td>-15.7 [-49.65, 18.25]</td>
</tr>
<tr>
<td>25.5 Idiopathic cough</td>
<td>1</td>
<td>Mean Difference</td>
<td>-9.2 [-53.05, 34.65]</td>
</tr>
<tr>
<td>25.6 All causes</td>
<td>1</td>
<td>Mean Difference</td>
<td>2.3 [-13.20, 17.80]</td>
</tr>
<tr>
<td>Section</td>
<td>Description</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>---------</td>
<td>-------------</td>
<td>----------------------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>26</td>
<td>Change in sputum Cys-LT [ng/mL]</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.1 [-1.50, 1.70]</td>
</tr>
<tr>
<td>26.1</td>
<td>PNDS</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.2 [-1.85, 1.45]</td>
</tr>
<tr>
<td>26.2</td>
<td>GORD</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.6 [-2.85, 1.65]</td>
</tr>
<tr>
<td>26.3</td>
<td>CVA</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-1.4 [-3.25, 0.45]</td>
</tr>
<tr>
<td>26.4</td>
<td>Bronchiectasis</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.3 [-2.30, 2.90]</td>
</tr>
<tr>
<td>26.5</td>
<td>Idiopathic cough</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.4 [-1.20, 0.40]</td>
</tr>
<tr>
<td>26.6</td>
<td>All causes</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>27</td>
<td>Change in sputum IL-8 [ng/mL]</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-3.5 [-61.95, 54.95]</td>
</tr>
<tr>
<td>27.1</td>
<td>PNDS</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-27.0 [-88.95, 34.95]</td>
</tr>
<tr>
<td>27.2</td>
<td>GORD</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>1.5 [-86.10, 89.10]</td>
</tr>
<tr>
<td>27.3</td>
<td>CVA</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-74.7 [-146.30, -310]</td>
</tr>
<tr>
<td>27.4</td>
<td>Bronchiectasis</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-20.9 [-122.10, 80.30]</td>
</tr>
<tr>
<td>27.5</td>
<td>Idiopathic cough</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-21.5 [-48.25, 5.25]</td>
</tr>
<tr>
<td>27.6</td>
<td>All causes</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>28</td>
<td>Change in sputum TNF-α [ng/mL]</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-3.9 [-17.70, 9.90]</td>
</tr>
<tr>
<td>28.1</td>
<td>PNDS</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>3.6 [-8.40, 15.60]</td>
</tr>
<tr>
<td>28.2</td>
<td>GORD</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [-1.30, 13.80]</td>
</tr>
<tr>
<td>28.3</td>
<td>CVA</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.4 [-10.30, 11.10]</td>
</tr>
<tr>
<td>28.4</td>
<td>Bronchiectasis</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>28.5</td>
<td>Idiopathic cough</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.3 [-4.85, 5.45]</td>
</tr>
<tr>
<td>28.6</td>
<td>All causes</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>29</td>
<td>Change in eNO [ppb]</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-2.1 [-3.56, -0.64]</td>
</tr>
<tr>
<td>29.1</td>
<td>PNDS</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-3.1 [-5.75, -0.45]</td>
</tr>
<tr>
<td>29.2</td>
<td>GORD</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-3.3 [-6.45, -0.15]</td>
</tr>
<tr>
<td>29.3</td>
<td>CVA</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.90 [-4.50, 2.70]</td>
</tr>
<tr>
<td>29.4</td>
<td>Bronchiectasis</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-1.4 [-4.85, 2.05]</td>
</tr>
<tr>
<td>29.5</td>
<td>Idiopathic cough</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-2.1 [-3.56, -0.64]</td>
</tr>
<tr>
<td>29.6</td>
<td>All causes</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>30</td>
<td>Change in exhaled CO [ppm]</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.1 [-0.60, 0.40]</td>
</tr>
<tr>
<td>30.1</td>
<td>PNDS</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.3 [-1.00, 0.40]</td>
</tr>
<tr>
<td>30.2</td>
<td>GORD</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.0 [-1.10, 0.50]</td>
</tr>
<tr>
<td>30.3</td>
<td>CVA</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.7 [-1.60, 0.20]</td>
</tr>
<tr>
<td>30.4</td>
<td>Bronchiectasis</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>0.3 [-0.95, 0.75]</td>
</tr>
<tr>
<td>30.5</td>
<td>Idiopathic cough</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>-0.34 [-0.66, -0.02]</td>
</tr>
<tr>
<td>30.6</td>
<td>All causes</td>
<td>Mean Difference (Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>31</td>
<td>Proportion with adverse effects</td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>1.67 [0.92, 3.04]</td>
</tr>
<tr>
<td>32</td>
<td>Proportion with specific adverse effects</td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>32.1</td>
<td>Hoarseness</td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>32.2</td>
<td>Sore throat</td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>32.3</td>
<td>Oral candidiasis</td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>0.0 [0.0, 0.0]</td>
</tr>
<tr>
<td>33</td>
<td>Proportion with severe adverse effects</td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
</tbody>
</table>
WHAT'S NEW
Last assessed as up-to-date: 13 December 2012.

<table>
<thead>
<tr>
<th>Date</th>
<th>Event</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 March 2013</td>
<td>Amended</td>
<td>Minor amendments to the text. Conclusions not changed.</td>
</tr>
</tbody>
</table>

CONTRIBUTIONS OF AUTHORS
The protocol was written by KJ and IY, based on the protocol by Anderson-James 2013, with input from all co-reviewers. Studies were independently assessed and data extracted by KJ and IY. Initial analysis was undertaken by KJ, with IY and AC. All authors contributed to the analysis and final review.

DECLARATIONS OF INTEREST
KJ, AC and RB declare no conflicts of interest.

Professor Fong declares that he has received travel and accommodation sponsorship several times to speak at or participate in educational meetings, which have been organised by an independent organising committee and sponsored by industry. Professor Fong is involved with the Lung Cancer Consultative Group of the Australian Lung Foundation (not-for-profit, public benevolent institution) and attends professional scientific meetings including those organised by the Thoracic Society of Australia and New Zealand, where some unrestricted sponsorship is usually provided by industry.

Dr Yang declares that he has received travel and accommodation sponsorship several times to speak at or participate in educational meetings, which have been organised by an independent organising committee and sponsored by industry. Dr Yang is involved with the National COPD Executive of the Australian Lung Foundation (not-for-profit, public benevolent institution) and attends professional scientific meetings including those organised by the Thoracic Society of Australia and New Zealand, where some unrestricted sponsorship is usually provided by industry.

SOURCES OF SUPPORT
Internal sources
  • School of Medicine, The University of Queensland, Australia.
  
  Funding towards KJ’s MBBS Honours project to the value of $1000
**External sources**

- Australian Cochrane Airways Group Network Student Scholarship, supported by the Asthma Foundation Queensland, Australia. For KJ to the value of $2000
- NHMRC Practitioner Fellowship (AC, KF) and NHMRC Career Development Fellowship (IY), Australia.

**DIFFERENCES BETWEEN PROTOCOL AND REVIEW**

**Criteria for considering studies for this review**

**Types of participants**

- BHR was removed as an exclusion criterion for types of participants.
- We included one study where people as young as 15 (years) were eligible for inclusion (Evald 1989).
- We included one study of cough for at least two weeks (Ponsioen 2005). Unpublished data excluding people with acute cough was used when available.

**Types of outcome measures**

The following additional secondary outcomes were recorded and analysed:

- Proportion of participants with a greater than 50% reduction in cough severity measure at follow up.
- Proportion of participants with clinical cure at follow up.
- Mean change in pulmonary function measures (spirometry, peak expiratory flow (PEF)).
- Biomarkers of inflammation - sputum biomarkers (total and differential cell counts, inflammatory mediators), exhaled gases.

Where a study reported two or more cough severity measures of equal ranking on the hierarchy of cough severity measures, the measure most comparable to those used by other studies for the same outcome comparison was used in meta-analysis.

**Search methods for identification of studies**

With regard to contacting experts in the field, we contacted only the authors of identified trials, and we searched manufacturers’ online clinical trial registries, rather than contact manufacturers directly.

**Data collection and analysis**

Cross-over trials were included in pooled data where first period data were available. Where these were not available, data were analysed using the generic inverse variance method.

We explored publication bias using a funnel plot when meta-analysis with at least ten studies was possible.
INDEX TERMS

Medical Subject Headings (MeSH)
Acute Disease; Administration, Inhalation; Adrenal Cortex Hormones [*administration & dosage]; Antitussive Agents [*administration & dosage]; Chronic Disease; Cough [*drug therapy]; Randomized Controlled Trials as Topic; Treatment Outcome

MeSH check words
Adult; Humans