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Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care (Review)

Coxeter P, Del Mar CB, McGregor L, Beller EM, Hoffmann TC

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[Intervention Review]

Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Peter Coxeter¹, Chris B Del Mar¹, Leanne McGregor², Elaine M Beller¹, Tammy C Hoffmann^{1,3}

¹Centre for Research in Evidence-Based Practice (CREBP), Bond University, Gold Coast, Australia. ²Centre of National Research on Disability and Rehabilitation (CONROD), Menzies Health Institute Queensland | School of Allied Health, Griffith University, Southport, Australia. ³School of Health and Rehabilitation Sciences, The University of Queensland, Brisbane, Australia

Contact address: Tammy C Hoffmann, Centre for Research in Evidence-Based Practice (CREBP), Bond University, Gold Coast, Queensland, 4229, Australia. thoffmann@bond.edu.au.

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ABSTRACT

Background

Shared decision making is an important component of patient-centred care. It is a set of communication and evidence-based practice skills that elicits patients' expectations, clarifies any misperceptions and discusses the best available evidence for benefits and harms of treatment. Acute respiratory infections (ARIs) are one of the most common reasons for consulting in primary care and obtaining prescriptions for antibiotics. However, antibiotics offer few benefits for ARIs, and their excessive use contributes to antibiotic resistance - an evolving public health crisis. Greater explicit consideration of the benefit-harm trade-off within shared decision making may reduce antibiotic prescribing for ARIs in primary care.

Objectives

To assess whether interventions that aim to facilitate shared decision making increase or reduce antibiotic prescribing for ARIs in primary care.

Search methods

We searched CENTRAL (2014, Issue 11), MEDLINE (1946 to November week 3, 2014), EMBASE (2010 to December 2014) and Web of Science (1985 to December 2014). We searched for other published, unpublished or ongoing trials by searching bibliographies of published articles, personal communication with key trial authors and content experts, and by searching trial registries at the National Institutes of Health and the World Health Organization.

Selection criteria

Randomised controlled trials (RCTs) (individual level or cluster-randomised), which evaluated the effectiveness of interventions that promote shared decision making (as the focus or a component of the intervention) about antibiotic prescribing for ARIs in primary care.

Data collection and analysis

Two review authors independently extracted and collected data. Antibiotic prescribing was the primary outcome, and secondary outcomes included clinically important adverse endpoints (e.g. re-consultations, hospital admissions, mortality) and process measures (e.g. patient satisfaction). We assessed the risk of bias of all included trials and the quality of evidence. We contacted trial authors to obtain missing information where available.

Main results

We identified 10 published reports of nine original RCTs (one report was a long-term follow-up of the original trial) in over 1100 primary care doctors and around 492,000 patients.

The main risk of bias came from participants in most studies knowing whether they had received the intervention or not, and we downgraded the rating of the quality of evidence because of this.

We meta-analysed data using a random-effects model on the primary and key secondary outcomes and formally assessed heterogeneity. Remaining outcomes are presented narratively.

There is moderate quality evidence that interventions that aim to facilitate shared decision making reduce antibiotic use for ARIs in primary care (immediately after or within six weeks of the consultation), compared with usual care, from 47% to 29%: risk ratio (RR) 0.61, 95% confidence interval (CI) 0.55 to 0.68. Reduction in antibiotic prescribing occurred without an increase in patient-initiated re-consultations (RR 0.87, 95% CI 0.74 to 1.03, moderate quality evidence) or a decrease in patient satisfaction with the consultation (OR 0.86, 95% CI 0.57 to 1.30, low quality evidence). There were insufficient data to assess the effects of the intervention on sustained reduction in antibiotic prescribing, adverse clinical outcomes (such as hospital admission, incidence of pneumonia and mortality), or measures of patient and caregiver involvement in shared decision making (such as satisfaction with the consultation; regret or conflict with the decision made; or treatment compliance following the decision). No studies assessed antibiotic resistance in colonising or infective organisms.

Authors' conclusions

Interventions that aim to facilitate shared decision making reduce antibiotic prescribing in primary care in the short term. Effects on longer-term rates of prescribing are uncertain and more evidence is needed to determine how any sustained reduction in antibiotic prescribing affects hospital admission, pneumonia and death.

PLAIN LANGUAGE SUMMARY

Interventions that facilitate shared decisions between primary care clinicians and patients about antibiotic use for acute respiratory infections

Review question

We wanted to see if shared decision making was better or worse than usual care in reducing antibiotic prescribing for an acute respiratory infection in primary care.

Background

Shared decision making enables health decisions to be made jointly by a clinician and patient. The decision making occurs after the options and their benefits and harms have been discussed together with the patient's values and preferences.

Acute respiratory infections (such as an acute cough, middle ear infection or sore throat) are one of the most common reasons to see a health professional, and antibiotics are commonly prescribed despite good evidence that they have little benefit for these conditions. Any decision to prescribe an antibiotic should be balanced by any benefits against the risk of common harms (such as rash and stomach upset) and the contribution to antibiotic resistance - now a major threat to human health.

Shared decision making provides an ideal opportunity within a primary care consultation for greater consideration about the trade-off between benefit and harm of antibiotics for acute respiratory illnesses. Antibiotic prescribing may decrease as a result.

Study characteristics

We identified 10 studies (nine trials and one follow-up study) up to December 2014. In total, the studies involved over 1100 primary care doctors and around 492,000 patients. The intervention was different in each study. Six of the studies involved training clinicians (mostly primary care doctors) in communication skills that are needed to facilitate shared decision making. In three studies, as well as training doctors in these skills, patients were also given written information about antibiotics for acute respiratory infections. All included trials received funding from government sources. No studies declared a conflict of interest.

Key results

Interventions that aim to facilitate shared decision making significantly reduce antibiotic prescribing for acute respiratory infections in primary care, without a decrease in patients' satisfaction with the consultation, or an increase in repeat consultations for the same illness. There was not enough information to decide whether shared decision making affects other clinically adverse secondary outcomes, measures of clinician and patient involvement in sharing decision making, or antibiotic resistance.

Quality of the evidence

We rated the quality of the evidence as moderate or low for all outcomes.

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SUMMARY OF FINDINGS FOR THE MAIN COMPARISON [Explanation]

Shared decision making compared to usual care for acute respiratory infections in primary care

Patient or population: antibiotic use in acute respiratory infections Setting: primary care Intervention: interventions to facilitate shared decision making

Comparison: usual care

Outcomes	Anticipated absolute e	ffects* (95% CI)	Relative effect (95% CI)	∾ of participants (studies)	Quality of the evidence Comments (GRADE)	
	Risk with usual care	Risk with Interventions to facilitate shared de- cision making				
Antibiotics prescribed or dispensed (6 weeks			RR 0.61 (0.55 to 0.68)	10172 (8 RCTs)	⊕⊕⊕⊖ MODERATE ¹	
or less) assessed with: risk ra- tio	47 per 100	29 per 100 (26 to 32)				
Antibiotics prescribed or dispensed (12			RR 0.74 (0.49 to 1.11)	481588 (3 RCTs) ³	$\oplus \oplus \bigcirc \bigcirc$ LOW ¹²	
months or greater) assessed with: risk ra- tio	47 per 100	35 per 100 (23 to 52)				
Patient initiated re-con-	Moderate		RR 0.87	1861	$\oplus \oplus \oplus \bigcirc$	
sultations for the same	40 per 100	35 per 100 (30 to 41)	(0.74 to 1.03)	(4 RCTs)	MODERATE ¹	
Patient satisfaction I with the consultation	Moderate		OR 0.86	1052	⊕⊕⊖⊖ LOW ¹⁴	
	71 per 100	68 per 100 (58 to 76)	(0.57 to 1.30)	(2 RCTs)		

*The risk in the intervention group (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% Cl).

Cl: confidence interval; OR: Odds ratio; RCT: randomised controlled trial; RR: Risk ratio

GRADE Working Group grades of evidence

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

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

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

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

Downgraded one level because of risk of bias: participants in most studies were aware of whether they had received the intervention or not.

 2 Downgraded one level because of imprecision: confidence interval includes reduction and possible increase in use of antibiotics. There was considerable heterogeneity in the rates of antibiotic prescribing during longer-term follow-up (12 months or greater).

³ Sample numbers in one trial, Butler 2012, were calculated from mean list size at baseline multiplied by the number of participating practices in each group (practice list sizes vary over time and no denominator data were available).

⁴ Downgraded one level due to imprecision: confidence interval includes both satisfaction and lack of satisfaction of patients with the consultation.

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BACKGROUND

Description of the condition

Acute respiratory infections (ARIs) are one of the most common reasons for consulting in primary care. Antibiotics are often prescribed (Gill 2006; Gonzales 1997; Gonzales 2001), often unnecessarily as systematic reviews conclude that antibiotics have little benefit for reducing symptom duration or complications in acute otitis media (Venekamp 2015), sore throat (Spinks 2013), bronchitis (Smith 2014), and sinusitis (Ahovuo-Saloranta 2014), and no benefit for laryngitis (Gonzales 2001) or colds (Kenealy 2013). The limited benefits of antibiotics for ARIs may be outweighed by unnecessary exposure to common adverse reactions (such as diarrhoea, candidiasis, rash, abdominal pain and/or diarrhoea and nausea and/or vomiting (Gillies 2015)), increased healthcare costs and contribution to antibiotic resistance (Chung 2007; Costelloe 2010).

Several doctor- and patient-related factors influence clinicians' prescribing behaviour. They include: clinical uncertainty and fear of disease progression; inadequate physician knowledge (Altiner 2007); underestimates of the contribution of prescribing antibiotics to the development of resistance (Wood 2013); and perceived patient expectations for an antibiotic and the subsequent pressure felt to meet this expectation (Arroll 2002). Antibiotic prescribing for ARIs also creates a 'vicious cycle' through the medicalisation of otherwise uncomplicated and self limiting illnesses, encouraging patients to re-consult with similar expectations for an antibiotic for similar illness episodes in the future (Butler 1998).

Antibiotic use exerts a selection pressure on bacteria to develop resistance (WHO 2012). Patients prescribed an antibiotic for respiratory tract infections develop measurable bacterial resistance in their commensal bacteria to that antibiotic for up to 12 months (Costelloe 2010). Although the development of individual resistance is transient, and decays after about a year in the absence of antibiotic use, it is sufficient to sustain high levels of population resistance (Chung 2007). Persistent prescribing of antibiotics, and excessive use of broad spectrum antibiotics in place of narrower spectrum ones, are modifiable factors that contribute to resistance (WHO 2012). Antibiotic resistance is now an evolving global threat to public health (WHO 2012). The rational use of antibiotics is therefore one of the most important strategies for preserving the therapeutic benefit of antibiotic treatment (WHO 2001; WHO 2012).

Description of the intervention

Shared decision making is the process of enabling a health professional and patient to make a joint treatment or management decision based on the best available evidence and the patient's values and preferences (Charles 1997; Makoul 2006). It consists of eliciting patients' expectations and clarifying any misperceptions, discussing treatment options, and communicating the benefits and harms of each option and their likelihood. Shared decision making supports the principle of patient autonomy and the right to self determination (Elwyn 2012), and has been shown to improve patients' satisfaction with decisions and concordance of decisions with their values (Spatz 2012). Some of the skills required of clinicians to facilitate shared decision making include proficient communication and rapport building skills as well as access to the best available evidence. It is one of the most important ways of bringing evidence to the point of clinical decisions and a potential strategy for reducing the overuse of ineffective treatments (Elwyn 2012).

How the intervention might work

The diagnostic uncertainty associated with ARIs and the trade-off between the benefits and harms of antibiotics mean that shared decision making may provide an ideal opportunity for clinicians and their patients to choose appropriate treatment or management options, including the decision to not use an antibiotic (Butler 2001). By engaging the patient and clinician to explicitly discuss the benefits and harms of antibiotics against a background of evidence demonstrating that it is less effective than most patients expect, there is high potential for it to be effective. Many patients elect for conservative treatment options after participating in shared decision making (Elwyn 2012).

Why it is important to do this review

Concern about antibiotic resistance is now an international public health crisis (WHO 2012), and finding ways to minimise unnecessary antibiotic prescribing in primary care is imperative. Shared decision making may be an important process to achieve this. Several related Cochrane systematic reviews have been undertaken. Arnold 2005 reviewed the effectiveness of interventions to improve antibiotic stewardship in outpatient care (including the decision to prescribe an antibiotic, and the type, dose and duration of antibiotic therapy). However, broad inclusion criteria and subsequent heterogeneity of the identified interventions limited the generalisability of practice recommendations. Importantly, this review also did not focus on, or explicitly consider, shared decision making interventions for inclusion.

The review by Stacey assessed the effectiveness of decision aids for people facing any treatment or screening decision (Stacey 2014). Decision aids are only one tool used to facilitate shared decision making in clinical care, and it may be enabled through methods other than, or in addition to, decision aids. Similarly, the review by Kinnersley evaluated the effect of interventions to encourage patient health communication and information seeking prior to the primary care consultation that shared some but not all compo-

nents necessary for shared decision making to occur (Kinnersley 2007). Légaré 2014 assessed the effectiveness of interventions to facilitate clinicians' uptake of shared decision making but not the use or effect of shared decision making in a particular condition. The growing interest in shared decision making for potential improvement in treatment decisions and patient outcomes is evident from Cochrane systematic reviews in other clinically important areas including mental health (Duncan 2010) and paediatric oncology (Coyne 2013). If shared decision making is shown to reduce prescribing among primary care doctors, then steps can be taken to incorporate it into primary care consultations for ARI across many countries.

OBJECTIVES

To assess whether interventions that aim to facilitate shared decision making increase or reduce antibiotic prescribing for ARIs in primary care.

METHODS

Criteria for considering studies for this review

Types of studies

Randomised controlled trials (RCTs) (individual level or cluster-RCTs), which evaluated the effectiveness of shared decision making in reducing antibiotic prescribing in primary care. Quasi-RCTs, quasi-experimental studies (controlled clinical trials), controlled before and after studies and interrupted time series analyses were not eligible.

Types of participants

As interventions that aim to facilitate shared decision making may be directed at clinicians, patients, or both, participants eligible for this review could be:

1. clinicians who provide primary care (community practices, hospital-affiliated or government-run outpatient clinics); or

2. patients who present with any combination of symptoms of acute (less than four weeks' duration) respiratory infection (or the parents of similarly affected children).

Types of interventions

There is no one accepted definition of shared decision making (Makoul 2006); nor is there consensus on the core skills that shared decision making training should address (Légaré 2013). Therefore, we considered interventions eligible if the trial explicitly stated that

the intervention was aimed at facilitating shared decision making or if the intervention explicitly addressed more than one of the essential elements of shared decision making that are described by Makoul 2006. These include: explaining the problem to be addressed; discussing options; communicating benefits and risks of each option; eliciting patient expectations, values, preferences or concerns; discussing patients' ability/self efficacy; and checking or clarifying understanding.

These elements may have been addressed by providing training in specific skills or providing decision support information or tools (such as decision aids (Stacey 2014), option grids (Elwyn 2013), or decision boxes (Giguere 2012)), which provide information about relevant issues (such as options, benefits, harms, questions to ask, etc). The skills training and information/tools could be provided to either clinicians, patients, or both. Interventions may have been delivered in any primary care environment and we imposed no restriction on the training and/or information mode, format or intensity of delivery.

We did not include interventions that consisted solely of the passive provision of patient information without the two-way sharing of information necessary for shared decision making, or which aimed to enhance clinicians' and/or patients' general communication skills.

Types of outcome measures

Primary outcomes

1. Prescription of antibiotics (for example, antibiotics prescribed per consultation, or a change in the population rate of antibiotic prescriptions per unit of time).

Secondary outcomes

1. Number or rate of patient-initiated re-consultations for unresolved ARI (i.e. same illness episode).

2. Incidence of colonisation with, or infection due to, antibiotic-resistant organisms.

3. Incidence of hospital admission.

4. Incidence of pneumonia (clinical with radiological confirmation).

5. Incidence of acute otitis media complications (for example, tympanic membrane perforation, contralateral otitis (in unilateral cases), mastoiditis, meningitis).

6. Mortality due to respiratory illness or similar.

- 7. All-cause mortality.
- 8. Measures of patient and caregiver satisfaction.

9. Measures of patient and caregiver satisfaction with the decision reached, decisional conflict and decisional regret.

10. Measures of extent of patient involvement in the decision making process (for example, consultations analysed using tools such as the OPTION instrument; Elwyn 2003).

11. Measures of treatment compliance or adherence to decision reached.

Search methods for identification of studies

Electronic searches

We searched the Cochrane Central Register of Controlled Trials (CENTRAL 2014, Issue 11), which includes the Cochrane Acute Respiratory Infections Group's Specialised Register, MEDLINE (1946 to November week 3, 2014), EMBASE (2010 to December 2014) and Web of Science (1985 to December 2014).

We searched MEDLINE using the search terms described in Appendix 1. We combined the MEDLINE search with the Cochrane Highly Sensitive Search Strategy for identifying randomised trials in MEDLINE: sensitivity- and precision-maximising version (2008 revision); Ovid format. We used the MEDLINE search strategy to search CENTRAL and adapted it to search EM-BASE (Appendix 2) and Web of Science (Appendix 3). We imposed no language, publication date or publication status restrictions on the electronic database searches.

We searched the National Institutes of Health registry of clinical trials (www.clinicaltrials.gov) and the World Health Organization's (WHO) clinical trials registry (www.who.int/ictrp/en/) for completed and ongoing studies eligible for inclusion. We searched Web of Science and EMBASE to identify potentially relevant conference abstracts and proceedings.

Searching other resources

We searched the bibliographies of retrieved articles and published reviews for additional studies. We personally communicated with trial authors of significant publications and content experts (Professor Paul Little, Professor Christopher Butler and Professor France Légaré) to identify further published, unpublished or ongoing trials.

Data collection and analysis

Selection of studies

We merged search results into reference management software (Endnote X6) and removed duplicate references. Two review authors (PC, LM) independently screened the titles and abstracts of retrieved records. We attempted to identify multiple reports of single studies following the criteria recommended in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We retrieved full-text copies of all potentially relevant articles for full-text evaluation. The final list of eligible trials was confirmed

following discussion and consensus among review authors (PC, TH, LM, CDM).

Data extraction and management

Two review authors (PC, LM) independently extracted data from each included trial using a specifically designed electronic data extraction form. We resolved disagreements by discussion and consensus, with one review author (CDM) acting as arbitrator where required. Data extraction was blind to names of authors, institutions and publication title. We extracted the following key study features where available:

1. Trial characteristics and methodological quality - risk of bias (see below); trial design, including unit of randomisation and number of comparator arms; blinding; generation of allocation sequence; allocation concealment; number of participants; theoretical or conceptual basis of the intervention; number of intervention components; description of intervention and comparator arms; length of follow-up; sample size estimate (power calculation); number of patients randomised to each intervention arm; number of patients completing the trial; reasons for withdrawal; and intention-to-treat (ITT) or per protocol analysis.

2. Patient (and/or caregiver) characteristics - age, gender and sociodemographic variables; types of ARI; duration of ARI prior to study recruitment; co-morbidities.

3. Healthcare professional characteristics - age; gender; experience; primary care setting type.

4. Outcome measures - all primary and secondary outcomes.

Assessment of risk of bias in included studies

Two authors (PC, LM) independently assessed the risk of bias of included studies and two acted as arbitrators (TH, CDM). We assessed risk of bias using the 'Risk of bias' tool available in RevMan 2014 and the criteria explained in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We assessed the reliability of the sequence generation, allocation concealment, blinding (participants, personnel and outcome assessors), incomplete outcome data and selective outcome reporting bias, as well as other sources of bias. We ranked studies as high, low or unclear risk of bias as described in the *Cochrane Handbook for Systematic Reviews of Interventions* and present our assessments in a 'Risk of bias' summary figure (Higgins 2011). As all included studies were cluster-RCTs, we assessed additional sources of bias including recruitment bias, baseline imbalance between clusters, loss of clusters and incorrect analysis (Higgins 2011).

Measures of treatment effect

Measures of treatment effect included dichotomous (binary), rate and continuous primary or secondary outcome data. Some studies calculated mean difference (MD) for continuous outcomes (median difference or median and interquartile range where data are

not normally distributed) and for dichotomous outcomes, risk ratio (RR), odds ratio (OR) or rate ratio (RaR) were reported. In accordance with our protocol we have based the primary analysis on data reported as adjusted risk ratios. Additional analyses of the prescribing outcomes also present adjusted odds ratios and risk differences to incorporate additional information as analysed in the included studies.

Unit of analysis issues

Studies presented effect measures adjusted for clustering effects (at practice, provider and/or patient hierarchies) or potential confounders in multilevel analysis, and/or applied generalised linear mixed models or generalised estimating equations. Intraclass correlation coefficients were estimated in sample size calculations (Briel 2006; Cals 2009; Francis 2009; Légaré 2012; Little 2013; Welschen 2004), or reported (Briel 2006; Francis 2009), to account for clustering effects. Where intraclass correlation coefficients were not reported (Butler 2012; Cals 2013), we imputed them from another similar included study.

Dealing with missing data

The majority of studies performed ITT analyses (Briel 2006; Butler 2012; Cals 2009; Francis 2009; Légaré 2011; Little 2013; Welschen 2004). One study presented data only from practices with complete follow-up (Altiner 2007). The long-term followup study of Cals 2009 included data only where medical records could be accessed for the follow-up period (87.9% of original trial cohort) (Cals 2013). The principle of analysis was not stated in one study (Légaré 2012). Drop-out rates and contributing reasons were sufficiently disclosed in all studies, and one study reporting relatively high attrition performed a sensitivity analysis to explore effects from differential missing values (Altiner 2007).

Assessment of heterogeneity

We used a random-effects model for all meta-analyses due the observed methodological diversity and used the I² statistic to measure heterogeneity as recommended in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011).

Assessment of reporting biases

We minimised reporting bias by conducting a comprehensive search for studies that met the eligibility criteria, including grey literature and unpublished trials; and by contacting trials authors for missing information. There were insufficient studies to test for publication bias using a funnel plot.

Data synthesis

Meta-analyses of studies were limited to studies reporting a comparable effect estimate. Therefore, the test for overall effect is limited to analysis in each subgroup. Studies reporting data that could not be combined for meta-analysis are reported narratively. Forest plots were also not generated for data reported by a single study, or where the synthesis of available pilot data to the substantive study (for example, Légaré 2011) would not meaningfully increase the power or precision of observed effects. Similarly, meta-analyses of secondary outcomes were limited to studies reporting comparable measures, those providing similar effect estimates, or where there were sufficient trials for comparison (such as patient satisfaction with the consultation). Caution is warranted for conclusions for each outcome due to the low numbers of trials for each comparison. We used RevMan 2014 to enter and analyse data to estimate a weighted treatment effect (with 95% confidence intervals (CIs). We analysed data using the random-effects model due to the expected heterogeneity in combining diverse shared decision making interventions.

We created Summary of findings for the main comparison using the following outcomes: antibiotic prescribing in the short term (less than six weeks), longer-term antibiotic prescribing (12 months or longer), re-consultation for the same illness episode and patient satisfaction with the consultation. We used the five GRADE considerations (study limitations, consistency of effect, imprecision, indirectness and publication bias) to assess the quality of evidence of the studies contributing data for meta-analyses of prespecified outcomes (GRADE Working Group 2004). We used the methods and recommendations described in Section 8.5 and Chapter 12 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011), using GRADEpro GDT software (GRADEproGDT 2015). We justified decisions to downgrade or upgrade the quality of studies using footnotes and comments to aid the reader's understanding of the review.

Subgroup analysis and investigation of heterogeneity

There were insufficient studies to conduct a subgroup analysis of trials that incorporate shared decision making as part of a multifaceted intervention compared with trials in which shared decision making was the standalone intervention. Subgroup analysis of interventions targeted at clinicians versus patients/parents was also not conducted due to a lack of studies. We did not conduct planned subgroup analyses of children versus adult trial populations, trials with low risk of bias versus high risk, and cluster-RCTs versus individually randomised studies due to insufficient studies.

Sensitivity analysis

Insufficient studies prevented a planned sensitivity analysis excluding trials found to have a higher versus low risk of bias to examine the effect of trial quality on the magnitude and direction of effect.

RESULTS

Description of studies

Results of the search

We retrieved a total of 3272 studies from the searches of the electronic databases after duplicates were removed. Two review authors (PC, LM) independently screened record titles and abstracts and, following consensus, 3256 records did not meet our inclusion criteria and were excluded. A recent published study protocol was identified and we contacted the lead author to confirm the study was ongoing and study results would not be available in time for this review (Altiner 2012). We retrieved full-text reports of the remaining 16 records and two review authors (PC, LM) screened these independently. We excluded six studies after they did not meet the a priori eligibility criteria for shared decision making interventions. All review authors (PC, TH, LM, CDM) considered the provisional list of 10 studies for inclusion. We contacted trial authors of two of these studies for further elaboration on respective study interventions to determine that both were eligible for inclusion (Altiner 2007; Briel 2006). We included 10 published reports of nine original studies: one publication reports long-term follow-up outcome data of an earlier study (Cals 2013), and another published report, Légaré 2011, presents pilot data for relevant outcomes distinct from the subsequent substantive cluster-RCT (Légaré 2012). See Figure 1.

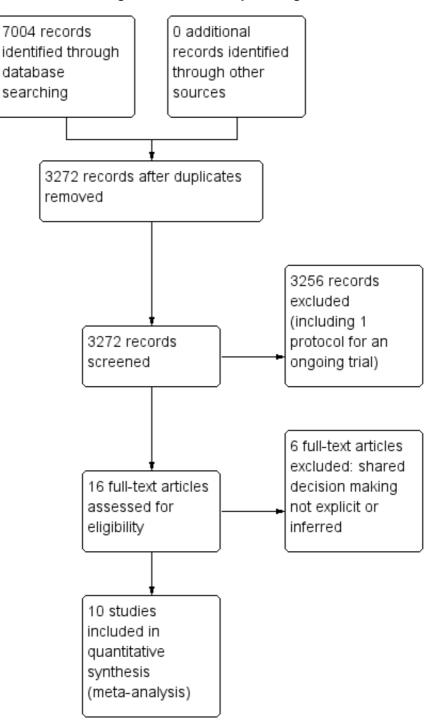


Figure I. PRISMA study flow diagram.

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Included studies

Study design

Six studies used a two-arm randomised group design: experimental versus control (usual care) (Altiner 2007; Butler 2012; Francis 2009; Légaré 2011; Légaré 2012; Welschen 2004). In one study the control group received the intervention after the experimental group had been exposed to the programme (Légaré 2011). Briel 2006 compared three arms: full intervention versus limited intervention versus non-randomised controls that acted as distractors to the intention of the real comparison and were not analysed). Two studies compared four parallel study arms: intervention (a) versus intervention (b) versus intervention (a + b) versus control (Cals 2009; Little 2013). These two cluster-RCTs incorporated a pre-specified factorial analysis plan (Cals 2009; Little 2013). Trial data for interventions not relevant to the present review (such as C-reactive protein point of care testing (Cals 2009; Little 2013), or costs (Butler 2012)) are not presented.

All nine original studies included were cluster-RCTs. The unit of randomisation in studies was the general practitioner (GP) (Altiner 2007; Briel 2006), general practice (Butler 2012; Cals 2009; Francis 2009; Little 2013), GP peer review group (Welschen 2004), family practice teaching unit (Légaré 2012), and family medicine group (Légaré 2011).

All trials received funding. None disclosed conflicts of interest except Cals 2013 (one study author received travel/lecture funds from a point of care test device manufacturer being evaluated in the study, for which data were not relevant to this review). Ethical approval was documented in all studies.

Characteristics of settings and participants

The studies were conducted in Germany (Altiner 2007), Switzerland (Briel 2006), the Netherlands (Cals 2009; Cals 2013; Welschen 2004); England (Francis 2009), Wales (Butler 2012; Francis 2009), and Canada (Légaré 2011; Légaré 2012). One multinational trial was conducted across six European countries (Netherlands, Belgium, Spain, Poland, England and Wales) (Little 2013).

Recruitment of clinicians

Participating general practitioners (GPs) were recruited directly (Altiner 2007; Briel 2006), or through participating general practices (Butler 2012; Cals 2009; Francis 2009; Little 2013), peer review groups (Welschen 2004), family practice teaching units (Légaré 2012), or family medicine groups (Légaré 2011). The existing nationwide structure of GP peer review groups in the Netherlands comprise GPs and collaborating pharmacists that aim to promote rational prescribing through audit and feedback (Welschen 2004). UK general practices comprise GPs and nurse prescribers (Little 2013). GPs within Family Medicine Groups in Canada (Quebec) also work closely with nurses for care of registered individuals (Légaré 2011). Family Practice Teaching Units in Quebec include both physician teachers and residents (Légaré 2012).

Recruitment of patients

Specific ARI diagnoses and participant eligibility varied a little across studies. In several studies GPs recruited all patients (adults and children accompanied by a legal guardian) (Légaré 2011; Légaré 2012; Welschen 2004), or only adult patients (Briel 2006), consulting with symptoms of ARI. One study included adult patients presenting predominately with acute lower respiratory tract infections (LRTIs) and upper respiratory tract infections (URTIs) (Little 2013). Cals 2009 included adult patients only with suspected LRTI. Altiner 2007 restricted patient eligibility to patients over 16 years of age consulting for acute cough. Conversely, Butler 2012 included patients with any condition registered with participating practices. Francis 2009 included only children (six months to 14 years) and their parents consulting for a respiratory tract infection.

Study exclusion criteria also differed a little among studies. Asthma was an explicit exclusion criterion in two studies (Altiner 2007; Francis 2009), and was not reason for exclusion in another (Welschen 2004). Patients with chronic obstructive pulmonary disease (COPD) were ineligible in one study (Altiner 2007), although were eligible for inclusion in two trials (Briel 2006; Welschen 2004). The proportion of patients diagnosed with asthma/COPD ranged from ~ 2% to ~ 3.5% (Briel 2006; Welschen 2004) up to ~ 18.5% (Little 2013). Patients with pneumonia were excluded from participation in one study (Briel 2006). However, they were eligible in two studies (Little 2013; Welschen 2004), and this was diagnosed in ~ 3.5% of participants in Welschen 2004.

Characteristics of interventions and comparisons

Interventions

Included trials assessed various multi-component interventions primarily aimed at facilitating clinicians shared management of decisions to reduce antibiotics for ARIs and their related symptoms in primary care.

The delivery of interventions occurred in usual clinical settings or central locations, and varied in intervention elements and scope

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and the frequency and duration (i.e. intensity) of sessions. All studies provided education and communication skills training that aimed to improve GPs' understanding of topics such as: the probability of bacterial or viral ARI; evidence for the benefit/risk of antibiotics and/or other treatment options; risk communication techniques; information exchange about symptoms and natural disease course; methods of eliciting patients' concerns and expectations; and agreement with the patient about a management plan and summing up. Communicative techniques used were derived from various theoretical models or frameworks.

Training in specific education and communication skills was delivered through peer- or facilitator-led interactive workshops and seminars or via web-based platforms, and supported with the use of videos, interactive exercises and decision aids or interactive booklets to facilitate patient participation in treatment decisions. Other programme components in some studies included consensus procedures, simulated patient consultations, personal reflection on clinical practice, reminders of expected behaviours and provision of antibiotic resistance trend data. Several interventions contained materials developed for patients, including education materials in waiting rooms (poster and leaflet), an interactive booklet for use within the consultation and as a take home resource, or decision support tool).

A summary of the main intervention components is described using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Hoffmann 2014) (see Table 1).

Comparators

In all trials the comparator was usual care, with the exception of Briel 2006 where GPs received training in a two-hour seminar on evidence-based US guidelines for ARIs.

Excluded studies

We excluded six studies as shared decision making was not explicit or inferred in the interventions evaluated (Characteristics of excluded studies).

Risk of bias in included studies

The methodological characteristics of the studies are reported in the Characteristics of included studies table. The 'Risk of bias' summary and 'Risk of bias' graph are presented in Figure 2 and Figure 3, respectively.

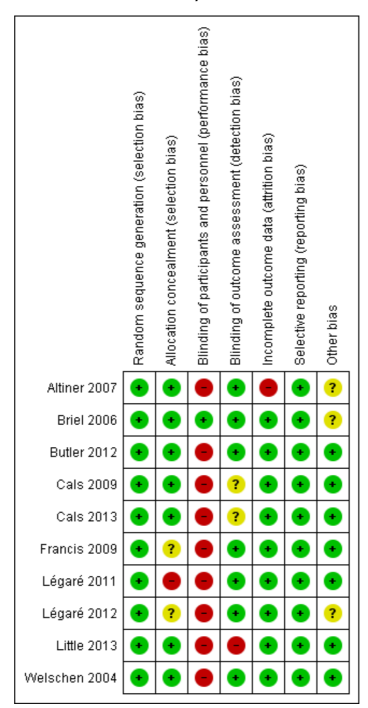
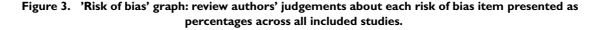
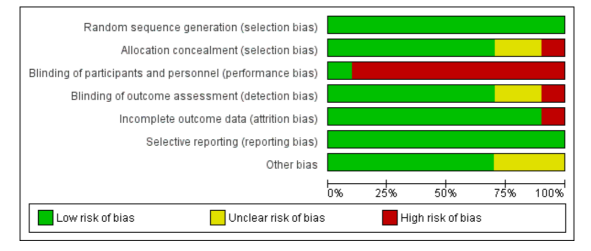


Figure 2. 'Risk of bias' summary: review authors' judgements about each risk of bias item for each included study.





Allocation

Methods of sequence generation comprised computer and/or program-generated methods (Altiner 2007; Briel 2006; Francis 2009; Légaré 2011; Légaré 2012; Little 2013). Studies used stratification and minimisation techniques (Little 2013), or dynamic block allocation (Butler 2012; Francis 2009), to achieve balanced groups on selected variables.

Concealed allocation occurred in most trials, with GPs blinded to group allocation until after randomisation, although methods of doing so were not clearly described in several trials (Altiner 2007; Briel 2006; Cals 2009; Francis 2009; Little 2013; Welschen 2004). In Légaré 2012, the family practice units were recruited before randomisation, but it is not clear when physicians in the units were recruited/consented. In Légaré 2011, individual family physicians were recruited after randomisation of the family medicine groups.

Blinding

The nature of the interventions meant blinding of the clinicians delivering the intervention was not possible. Briel 2006 reported blinding of general practitioners although this is not credible. Blinding of outcome assessment was not reported in Little 2013, although it was adequately described in all other included studies.

Incomplete outcome data

One study had high risk of attrition bias. Altiner 2007 reported that 17% of GPs were lost to follow-up at six weeks post-intervention and 41% at 12 months. The study authors explored the effect of high attrition by conducting a cluster level sensitivity analysis by imputing new values for missing average antibiotic rates: firstly, by performing a regression analysis according to GPs with complete data sets to receive a prediction rule of six weeks and 12 months prescribing rates from baseline prescribing rates and, secondly, by using these rules to estimate follow-up prescription rates for those physicians that dropped out of the study. Alternative estimates using last observations (baseline or six weeks) were similar, and the results of both sensitivity analyses were in line with reported results. Légaré 2012 reported that three of 12 randomised family practice teaching units were lost to follow-up. The loss of clusters was noted as a study limitation, but no further analysis was performed. Neither of these studies reported conducting statistical analysis on an intention-to-treat (ITT) basis.

The risk of attrition bias was low in the remaining studies.

Selective reporting

Several studies reported prospective trial registration (Butler 2012; Cals 2009; Cals 2013; Francis 2009; Légaré 2012; Little 2013), and/or had published trial protocols (Butler 2012; Cals 2009; Francis 2009; Légaré 2011; Légaré 2012). We detected no reporting bias by comparing these to the final reports. Only Briel 2006

neither reported trial registration nor published a protocol.

Other potential sources of bias

We considered recruitment bias to be minimal in the included trials as the unit of allocation was recruited into the trial before clusters were randomised. Similarly, we considered baseline imbalances between study group characteristics minimal as all studies disclosed baseline comparability and adjusted for important baseline differences in the analysis. In two studies there was sufficient loss of clusters following randomisation that may have introduced bias (Altiner 2007; Légaré 2012). All studies sufficiently reported the use of robust statistical methods to account for clustering in the analysis.

All studies reported a sample size calculation with the exception of Légaré 2011, which was designed as a pilot trial. An ITT analysis was pre-specified in all but two trials (Altiner 2007; Légaré 2012). Altiner 2007 included only practices with complete follow-up in the analysis and the method of analysis was not described in Légaré 2012.

The methods, timing and duration of patient recruitment varied across studies. Recruitment in some trials was planned to capture winter and/or autumn months (Cals 2009; Francis 2009; Légaré 2012; Welschen 2004). In the long-term follow-up study, Cals 2013, of the original cluster-RCT (Cals 2009), the end date of the follow-up period was chosen to ensure a similar number of winter days in each period. Recruitment in the Little 2013 study occurred at the end of the season for respiratory tract infections in participating European countries (February and May). One trial included registered practice populations over an entire year (Butler 2012). The timing and duration of participant recruitment (e.g. during limited/winter months versus annual periods) may influence study outcomes and seasonal variation in the frequency and severity of ARIs may affect results.

The possibility of selection bias remains a possibility, although trial authors report that the risk of bias was minimal as baseline GP and patient characteristics were disclosed in all studies and no systematic differences between known group characteristics or case-mix were observed. Altiner 2007 could not rule out that GPs, who were not monitored during the trial, may not have reported patients with acute cough who received an antibiotic. Participating GPs in Briel 2006 were considered highly motivated and several authors considered the possibility that GPs may have behaved differently while being monitored (Hawthorn effect) (Briel 2006; Francis 2009).

Intervention adherence was measured in only a few trials. Treatment fidelity was not measured in any of the included studies and sub-optimal exposure or delivery of the intervention as planned may dilute the observed effect.

Effects of interventions

See: Summary of findings for the main comparison Shared decision making compared to usual care for acute respiratory infections in primary care

Primary outcome

I. Prescription of antibiotics

There were data from all 10 included studies on antibiotic prescribing decisions for acute respiratory infection. However, they could not all be combined into one meta-analysis because of differences in adjusted effect estimates reported and outcome measurement time.

We extracted event and denominator data, and reported (or imputed) intra-class correlation coefficients, to calculate the risk ratio (RR) adjusted for the effects of clustering (Analysis 1.1; Analysis 1.2) to allow presentation of outcome data within a common scale. This also allowed us to combine trials reporting short (index consultation to \leq six weeks) and longer-term (\geq 12 months) intervention effects on antibiotic prescribing. Eight studies reporting short-term prescribing outcomes could be pooled in meta-analysis: the RR compared to usual care was 0.61, 95% confidence interval (CI) 0.55 to 0.68; P value = < 0.001 (Figure 4). There was a trend towards a reduction in antibiotic prescribing being maintained in the longer term: RR compared with usual care 0.74, 95% CI 0.49 to 1.11; P value = 0.14 (Figure 5). However, the non-significant results may be an artefact of the more conservative effect estimates using RR adjusted only for clustering.

Figure 4. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: I.I Antibiotics prescribed, dispensed or decision to use (short-term, index consultation to \leq 6 weeks).

			Shared decision making	Usual care		Risk Ratio	Risk Ratio	
Study or Subgroup	log[Risk Ratio]	SE	Tota	Total	Weight	IV, Random, 95% Cl	IV, Random, 95% Cl	
Francis 2009 (1)	-0.7867	0.4212	256	i 272	1.8%	0.46 [0.20, 1.04]		
Briel 2006 (2)	-0.1518	0.2619	259	ı 293	4.8%	0.86 [0.51, 1.44]		
Légaré 2012 (3)	-0.6931	0.2606	181	178	4.8%	0.50 [0.30, 0.83]		
Légaré 2011 (4)	-0.3738	0.204	81	70	7.9%	0.69 [0.46, 1.03]		
Cals 2009 (5)	-0.6846	0.19	201	230	9.1%	0.50 [0.35, 0.73]	-	
Altiner 2007 (6)	-0.6983	0.1666	1021	1143	11.8%	0.50 [0.36, 0.69]	+	
Little 2013 (7)	-0.3711	0.1251	2332	1932	20.9%	0.69 [0.54, 0.88]	-	
Welschen 2004 (8)	-0.4806	0.0917	905	i 818	38.9%	0.62 [0.52, 0.74]	•	
Total (95% CI)			5236	4936	100.0 %	0.61 [0.55, 0.68]	•	
Heterogeneity: Tau ² =	0.00; Chi² = 6.62	, df = 7 (F	° = 0.47); I² = 0%				0.005 0.1 1 10 20	<u>,</u>
Test for overall effect:	Z = 8.63 (P < 0.00	0001)					Shared decision making Usual care	.0

Footnotes

(1) Reported intra-class correlation co-efficient (ICC) = 0.24. Design effect and effective sample size calculated.

(2) Reported intra-class correlation co-efficient (ICC) = 0.04. Design effect and effective sample size calculated. Actual sample denomiator used to calculate risk ratio....

(3) Adjusted for cluster design, baseline values and patient age group (for analyses at teaching unit and physician levels).

(4) Reported intra-class correlation co-efficient (ICC) = 0.02. Design effect and effective sample size calculated.

(5) Reported intra-class correlation co-efficient (ICC) = 0.12. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in...

(6) Reported intra-class correlation co-efficient (ICC) = 0.20. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in... (7) Adjusted for baseline prescribing and clustering by physician and practice, age, smoking, sex, major cardiovascular or respiratory comorbidity, baseline symptoms,...

(r) regulated intra-class correlation co-efficient (ICC) = 0.09. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in...

Figure 5. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: 1.2 Antibiotics prescribed or dispensed (longer-term, \geq 12 months).

			Shared decision making	Usual care		Risk Ratio	Risk Ratio	
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% Cl	IV, Random, 95% Cl	
Cals 2013 (1)	-0.3817	0.2188	178	201	27.2%	0.68 [0.44, 1.05]		
Altiner 2007 (2)	-0.5625	0.1384	787	920	33.4%	0.57 [0.43, 0.75]	+	
Butler 2012 (3)	-0.0254	0.0053	239802	239700	39.4%	0.97 [0.96, 0.99]	•	
Total (95% Cl)			240767	240821	100.0%	0.74 [0.49, 1.11]	•	
Heterogeneity: Tau² = Test for overall effect:			(P = 0.0001); I ^z = 89%				0.005 0.1 1 10 2 Shared decision making Usual care	200

<u>Footnotes</u>

(1) Design effect (2.08) imputed from Cals (2009). Effective sample size calculated by imputing intra-class correlation co-efficient reported by Cals 2009 (0.12). Actual...
 (2) Reported intra-class correlation co-efficient (ICC) = 0.20. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in...
 (3) Numerators and denominators estimated from dispensing rates per 1000 registered patients and reported mean list sizes at baseline, respectively. Effective sample...

We also conducted a sensitivity analysis by pooling the results of trials reporting similar adjusted effect estimates (see Analysis 1.3; Analysis 1.4; Analysis 1.5). Three studies reported antibiotic prescription as an odds ratio (OR) adjusted for clustering and other covariates, and we were able to meta-analyse them: the pooled OR compared with usual care was 0.44, 95% CI 0.26 to 0.75; P value = 0.003 (Figure 6). Similarly a meta-analysis of two studies reporting a RR adjusted for clustering yielded a pooled RR compared with usual care of 0.64, 95% CI 0.49 to 0.84; P value = 0.001

(Figure 7). A meta-analysis of four studies reporting an adjusted risk difference (RD) yielded a pooled RD of -18.44%, 95% CI - 27.24 to -9.65% compared with usual care (Figure 8). The results of the primary meta-analysis (RR adjusted for clustering) are generally concordant with trials reporting comparable adjusted effect estimates, although not adjusting for covariates that may have differed slightly between randomised groups (which were adjusted for in the reports) results in some loss of precision and wider 95% CIs.

Figure 6. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: 1.3 Antibiotic prescriptions (index consultation) (adjusted odds ratio).

			Shared decision making	Usual care		Odds Ratio	Odds Ratio	
Study or Subgroup	log[Odds Ratio]	SE	Total	Total	Weight	IV, Random, 95% Cl	IV, Random, 95% Cl	
Briel 2006 (1)	-0.1508	0.3906	259	293	26.7%	0.86 [0.40, 1.85]	e	
Francis 2009 (2)	-1.2379	0.3716	256	272	28.1%	0.29 [0.14, 0.60]		
Altiner 2007 (3)	-0.9676	0.1936	1021	1143	45.2%	0.38 [0.26, 0.56]	-	
Total (95% CI)			1536	1708	100.0%	0.44 [0.26, 0.75]	◆	
Heterogeneity: Tau ² = Test for overall effect:			= 0.10); I ² = 57%				0.002 0.1 1 10	500
	2 0.02 () 0.00	-,					Shared decision making Usual care	

<u>Footnotes</u>

(1) Logistic regression with random effect for each cluster and patient covariates (age, sex, education, days with restriction at baseline).

(2) Odds ratio from multilevel modelling.

(3) Adjusted for patient's disease severity, average practice severity (severity of the disease rated by the GP), patients having fever (compared with no fever), and frequency of ...

Figure 7. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: 1.4 Antibiotic prescriptions (index consultation) (adjusted risk ratio).

			Shared decision making			Risk Ratio	Risk		
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% Cl	IV, Rando	m, 95% Cl	
Légaré 2012 (1)	-0.6931	0.2606	181	178	24.8%	0.50 [0.30, 0.83]			
Little 2013 (2)	-0.3711	0.1251	2332	1932	75.2%	0.69 [0.54, 0.88]			
Total (95% CI)			2513	2110	100.0%	0.64 [0.49, 0.84]	•		
Heterogeneity: Tau² =	0.01; Chi ² = 1.24	, df = 1 (F	° = 0.27); I² = 19%				0.01 0.1 1	10	100
Test for overall effect:	Z = 3.24 (P = 0.00	01)					Shared decision making		100

<u>Footnotes</u>

(1) Adjusted for cluster design, baseline values and patient age group (for analyses at teaching-unit and physician levels).

(2) Adjusted for baseline prescribing and clustering by physician and practice, age, smoking, sex, major cardiovascular or respiratory comorbidity, baseline symptoms,...

Figure 8. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: 1.5 Antibiotic prescriptions (index consultation or population rate per unit of time) (adjusted risk difference).

Study or Subgroup	Mean Difference	SE	Shared decision making Tota		Weight	Mean Difference IV, Random, 95% Cl	Mean Diff IV, Randon		
Butler 2012 (1)	-26.1	12.3363	23980:	2 239700	11.1%	-26.10 [-50.28, -1.92]			
Cals 2009 (2)	-26.1	5.6518	201	1 230	33.0%	-26.10 [-37.18, -15.02]			
Légaré 2011 (3)	-16	9.0754	8.	1 70	18.1%	-16.00 [-33.79, 1.79]			
Welschen 2004 (4)	-10.7	4.898	90:	5 818	37.9%	-10.70 [-20.30, -1.10]			
Total (95% CI) Heterogeneity: Tau² = Test for overall effect:			24098 = 0.19); I ^z = 37%	9 240818	100.0%	-18.44 [-27.24, -9.65]	-100 -50 0 Shared decision making	50 Usual care	100

Footnotes

(1) Analysis of covariance with the previous year's prescribing as a covariate.

(2) Crude 95%CI calculated and inflated for clustering by using standard deviation inflated by variance inflation factor. P value calculated from second order penalised..

(3) All P values adjusted for baseline values and the study's cluster design

(4) Intervention effect in multilevel analysis.

The absolute effect of the intervention for the outcome of antibiotics prescribed, dispensed, or decision to use, immediately after, or within six weeks, of the consultation was reduced from 47% to 29%.

Francis 2009 showed important reductions in antibiotics prescribed for children consulting for an ARI at the index consultation (intervention versus control: 19.5% versus 40.8%; adjusted

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OR 0.29; 95% CI 0.14 to 0.60. Francis 2009 was the only trial that also reported data on antibiotics taken (this was collected by telephone questionnaire). They reported the percentage of participants in each group that took antibiotics within the first two weeks (the data also include the antibiotics that were prescribed after the index consultation: 50 (19.5%) in the intervention group and 111 (40.8%) in the control group, with an adjusted OR of 0.35, 95% CI 0.18 to 0.66). A significant decrease in antibiotic prescriptions for acute cough was observed in Altiner 2007 at six weeks (adjusted OR 0.38, 95% CI 0.26 to 0.5; P value < 0.001) and 12 months (adjusted OR 0.55, 95% CI 0.38 to 0.80; P value = 0.002) post-intervention. Conversely, Briel 2006 was the only trial that found no significant reduction in antibiotics dispensed within two weeks of the index consultation (full intervention versus limited intervention: 13.5% and 15.7%; adjusted OR 0.86, 95% CI 0.40 to 1.93). DECISION+2 led to fewer patients deciding to use antibiotics immediately after the consultation (immediate versus no or delayed antibiotic use) for ARIs compared with usual care (27.2% versus 52.2%; adjusted RR 0.5, 95% CI 0.3 to 0.7) (Légaré 2012). Little 2013 demonstrated that antibiotic prescribing for predominately acute lower respiratory tract infections (LRTIs) and upper respiratory tract infections (URTIs) was lower in the intervention group compared with controls (36.1% versus 45.3%; adjusted RR 0.69, 95% CI 0.54 to 0.87). Cals 2009 demonstrated a reduction in antibiotic prescribing for patients with suspected LRTI recruited during the winters of successive years (2005 to 2006 and 2006 to 2007) (intervention versus control: 27.4%, 95% CI 25.6% to 36.6% versus 53.5%; 95% CI 43.8 to 63.2; P value < 0.01). Butler 2012 measured a mean 4.2% (95% CI 0.6% to 7.7%; P value = 0.02) reduction (as a

percentage of the mean in controls) in the total number of dispensed oral antibiotic items per 1000 registered patients for the year after the intervention practices were exposed to the STAR programme. A non-significant reduction in the decision to immediately use antibiotics was also observed in the pilot trial by Légaré 2011 (-16.0%; P value = 0.08). Welschen 2004 reported significantly reduced antibiotic prescribing rates for symptoms of ARIs (-10.7%, 95% CI -20.3% to -1.0%). In a long-term followup of Cals 2009, enhanced communication skills training showed sustained reduction in antibiotic prescribing at 3.67 years mean follow-up (intervention versus control: 26.3%, 95% CI 20.6% to 32.0 versus 39.1%, 95% CI 33.1% to 45.1%; corrected difference: -10.4%; P value = 0.02). See Table 2.

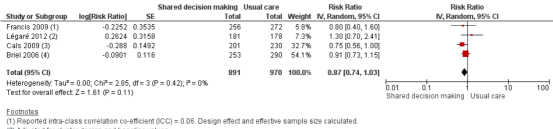
We graded the quality of evidence as moderate and low for antibiotic prescribing in the short term (less than six weeks) and long term (12 months or longer), respectively. See Summary of findings for the main comparison.

Secondary outcomes

I. Number or rate of patient-initiated re-consultations for unresolved ARI

Six studies reported adjusted effect estimates that we could not combine in a meta-analysis. We extracted data from four studies to calculate a RR adjusted for clustering, and pooled in meta-analysis. The RR compared to usual care was 0.87, 95% CI 0.74 to 1.03; P value = 0.11 (Analysis 1.6; Figure 9).

Figure 9. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: 1.6 Number or rate of re-consultations (risk ratio).



⁽²⁾ Adjusted for cluster design and baseline values.

(3) Reported intra-class correlation co-efficient (ICC) = 0.01. Design effect and effective sample size calculated.
 (4) Reported intra-class correlation co-efficient (ICC) = 0.04. Design effect and effective sample size calculated.

The proportion of re-consultations for the same illness episode reported in Briel 2006 was 44.7% versus 49.3% (adjusted RR compared to controls 0.97, 95% CI 0.78 to 1.21). The between-group consultation rates in Cals 2009 were 27.9% (95% CI 21.4 to 34.4)

and 37.0% (95% CI 30.4 to 43.6); P value = 0.14. Légaré 2012 reported no differences between groups (22.7% versus 15.2%; absolute difference 7.5%; adjusted RR compared to controls 1.3,

95% CI 0.7 to 2.3). Francis 2009 also reported no difference in the odds of re-consulting in primary care during the two weeks after the index consultation (12.9% versus 16.2%; adjusted OR 0.75 (0.41 to 1.38). Butler 2012 found no difference in median re-consultation rates after an index consultation for respiratory tract infections per 1000 registered patients at seven days (-0.65, 95% CI -1.69 to 0.55, P value = 0.446); 14 days (-1.33, -2.12 to 0.74; P value = 0.411); or 31 days (-2.32, 95% CI -4.76 to 1.95; P value = 0.503). Similarly, Little 2013 found the rates of new or worsening symptoms (including re-consultation in less than four weeks or hospital admission) did not differ significantly between groups (adjusted RR compared to controls 1.33, 95% CI 0.99 to 1.74; P value = 0.055). See Table 3.

We graded the quality of evidence as moderate. See Summary of findings for the main comparison.

2. Incidence of colonisation with, or infection due to, antibiotic-resistant organisms

No studies reported this outcome.

3. Incidence of hospital admission

Six trials reported serious adverse events (SAEs) requiring hospitalisation, although no significant differences between groups were observed. Butler 2012 reported a non-significant difference in the proportion of hospital admissions for possible respiratory tract infections and complications relative to the control group (-1.9%, 95% CI -13.2% to 8.2%; P value = 0.72). Briel 2006 reported that three patients were hospitalised (two patients in the full intervention group versus one in the limited intervention group). Six intervention and two usual care participants were hospitalised in Little 2013 (factorial analysis not reported). Francis 2009 reported seven hospitalisations (intervention = three, control = four). There were no occurrences of SAEs (death or admission to hospital) in Cals

2009. Cals 2013 reported five hospital admissions of 379 study participants: two patients receiving usual care (four exacerbations of chronic obstructive pulmonary disease (COPD) and one case of pneumonia), one randomised to C-reactive protein testing, and two episodes (pneumonia) in the combined intervention group (factorial analysis data not reported). See Table 4.

4. Incidence of pneumonia

Two studies reported on the incidence of pneumonia. Briel 2006 reported one case of pneumonia in the control group, and Cals 2013 reported two cases of pneumonia in patients receiving a combined intervention (factorial analysis data not reported) and two cases of pneumonia in those receiving usual care. See Table 5.

5. Incidence of acute otitis media complications

No studies reported on this outcome.

6. Mortality due to respiratory illness or similar

One study, Briel 2006, reported a fatal myocardial infarction following pneumonia in an elderly patient receiving a limited (control) intervention.

7. All-cause mortality

No studies reported on this outcome.

8. Measures of patient and caregiver satisfaction

The results from two studies could be pooled, giving an OR compared to controls of 0.86, 95% CI 0.57 to 1.30; P value = 0.47 (Analysis 1.7; Figure 10).

Figure 10. Forest plot of comparison: I Shared decision making versus usual care (control), outcome: 1.7 Patient satisfaction with the consultation.

			Shared decision making	Usual care		Odds Ratio	Odds Ratio
Study or Subgroup	log[Odds Ratio]	SE	Tota	Total	Weight	IV, Random, 95% Cl	IV, Random, 95% Cl
Francis 2009 (1)	-0.4463	0.338	246	263	34.3%	0.64 [0.33, 1.24]	
Briel 2006 (2)	0	0.2277	253	290	65.7%	1.00 [0.64, 1.56]	+
Total (95% CI)			499	553	100.0%	0.86 [0.57, 1.30]	•
Heterogeneity: Tau ² =	0.02; Chi ² = 1.20,	df=1 (P	= 0.27); I² = 17%				
Test for overall effect:	Z = 0.72 (P = 0.47))					Shared decision making Usual care

Footnotes (1) Odds Ration from multilevel modelling

(2) Proportion of patients with a maximum score of 70 (out of 70) used due to highly skewed scores. Logistic regression with random effect for each cluster and patient.

There were no differences observed between intervention and control groups in studies that reported this outcome. Briel 2006 found no difference in scores for patient satisfaction (Patient Satisfaction Questionnaire; score 0 to 70) between intervention and control groups (median 68 out of 70; % patients with 70 out of 70: 47.8% versus 49.0%; adjusted OR 1.00; 95% CI 0.64 to 1.31). Cals 2009 reported no differences in patient satisfaction with the index consultation (% at least very satisfied: 78.7%, 95% CI 72.5 to 84.9 versus 74.4%, 95% CI 68.2 to 80.6; P value = 0.88). In Francis 2009, the proportion of parents that were reported to be satisfied or very satisfied with the consultation were similar between groups (90.2% versus 93.5%; adjusted OR 0.64, 95% CI 0.33 to 1.22). Patient satisfaction (one = very dissatisfied, five = very satisfied) was also high and no between-group differences were observed in Welschen 2004 (adjusted mean difference (MD) -0.03, 95% CI -0.2 to 0.1). See Table 6.

We graded the quality of evidence as low. See Summary of findings for the main comparison.

9. Measures of patient and caregiver satisfaction with the decision reached, decisional conflict and decisional regret

Decisional conflict

One study measured GPs' decisional conflict using the Decisional Conflict Scale (DCS; 1 = low decisional conflict, 5 = very high decisional conflict) and found no difference between the intervention group and controls (MD 3.4, adjusted RR 3.5, 95% CI 0.3 to 38.0) (Légaré 2012). For patients' decisional conflict scores, the MD was 1.7 and the adjusted RR 0.8, 95% CI 0.2 to 2.4. See Table 7.

Decision regret (patients)

Légaré 2012 observed a clinically insignificant effect between the intervention and control groups on a decision regret measure (0 = very low regret, 100 = very high regret) with a mean of 12.4 in the intervention group and 7.6 in the control group; adjusted MD 4.8, 95% CI 0.9 to 8.7. Légaré 2011 also reported no difference in the proportion of patients with decisional regret between the study groups (7% in the intervention group versus 9% in the control; adjusted MD -2, 95% CI -12 to 5). See Table 8.

10. Measures of extent of patient involvement in the decision making process

Patient enablement

Three studies reported on patient enablement. Cals 2009 found no difference between intervention and control group scores on the Patient Enablement Instrument (PEI; score 0 to 12) (mean (SD): 3.29 (2.52) versus 3.06 (2.54); P value = 0.70). Francis 2009 found no between-group difference in a modified PEI measuring parent enablement (score 0 to 10; score greater or equal five: 40.2% versus 35.9%; adjusted OR 1.20, 95% CI 0.84 to 1.73). Briel 2006 found weak evidence for higher patient enablement on the PEI (median 8 out of 12; mean (SD) 8.49 (1.98) versus 8.15 (2.03); adjusted MD 0.35, 95% CI -0.05 to 0.75). See Table 9.

II. Measures of treatment compliance or adherence to decision reached

Decision quality

Légaré 2012 found no difference between GPs on a measure of GPs' decision quality (1 = very low quality, 10 = very high quality) (MD -0.2, 95% CI -0.6 to 0.2). The results were similar to the earlier pilot cluster-RCT, Légaré 2011 (MD -0.2, 95% CI -0.34 to 0.89; P value = 0.29). Similarly, there were no differences observed in patients' decision quality in Légaré 2012 (MD 0.0, 95% CI - 0.4 to 0.4) and Légaré 2011 (MD 0.1, 95% CI -0.88 to 0.94; P value = 0.57). See Table 10 and Table 11.

Adherence to decision

The only trial to measure adherence to the decision reached found no difference between intervention and control groups (87.7% of patients versus 91.5%; absolute difference of 3.8, adjusted RR 1.0, 95% CI 0.9 to 1.0) (Légaré 2012).

DISCUSSION

Summary of main results

Interventions aiming to promote shared decision making in primary care, as the focus or a core component of multi-faceted interventions, significantly reduced antibiotic prescribing for acute respiratory infections by almost 40% compared with usual care in the short term. There was insufficient evidence for sustained reductions in antibiotic prescribing over the longer term. There were no significant differences between groups receiving the intervention or usual care in clinical complications such as re-consultation for the same illness, or patient satisfaction with the consultation. There was also insufficient evidence to assess intervention effects on other clinically adverse or patient and/or caregiver shared decision process outcomes.

Overall completeness and applicability of evidence

A growing number of trials have examined the effect of interventions that aim to facilitate shared decision making, with all studies being conducted in the last 10 years (seven of 10 studies in the last five years), highlighting that shared decision making is a relatively new intervention.

All studies included acute upper or lower respiratory tract infection in children and/or adults consulting primary care or academic general practice. Trials were conducted in several high-income European countries and Canada. Applicability of findings to lowand middle-income countries and different cultural and healthcare settings is unknown.

We identified considerable heterogeneity in longer-term prescribing outcomes meta-analysed as risk ratio (see Analysis 1.2), and moderate to substantial heterogeneity in pooled results grouped under each reported effect estimate for the primary outcome (see Analysis 1.3; Analysis 1.4; Analysis 1.5). There was considerable diversity across included studies within each comparison in terms of the population (adults, children, or both), scale and composition of multi-component interventions evaluated, timing of the intervention and follow-up, outcome measures used and statistical techniques. The considerable heterogeneity observed in antibiotic prescribing rates over the longer term may be due to measurement differences in one study (Butler 2012) (all oral dispensed antibiotic items per 1000 registered patients for the year following exposure of practices to the intervention), or the low number of studies reporting longer-term sustainability of intervention effects. Substantial (although non-significant) heterogeneity apparent in studies reporting an adjusted odds ratio (OR) (see Analysis 1.3; I² statistic = 57%; P value = 0.10) may have resulted from the inclusion of one study reporting a statistically non-significant intervention effect (Briel 2006), where an unusually low antibiotic prescribing rate was noted (13.5% and 15.7% in the study groups) compared with other studies. Detecting an intervention effect may be difficult in a low prescribing setting. Some heterogeneity in pooled studies reporting an adjusted relative risk (Analysis 1.4; I² statistic = 19%; P value = 0.27) may result from true clinical and/or methodological diversity with the non-significant result being simply an artefact of only two studies being available for the comparison. Significant heterogeneity in four studies reporting adjusted risk differences (Analysis 1.5; I^2 statistic = 37; P value = 0.19) is likely due to inherent multiplicity of clinical and methodological factors.

The effect size of the included studies varied considerably, although there was general consistency in the direction of effects. The risk of bias overall in the included studies was low. Interventions varied markedly in the theoretical basis, and the components, scope, mode of delivery and duration. It is not possible, therefore, to identify which intervention components, combinations or modes of delivery most effectively promote shared decisions. Interventions and training were principally targeted at GPs. However, competence in the use of shared decision making was only reported in some trials, with no studies assessing intervention fidelity. Objective patient or clinician measures of adoption of shared decision making (e.g. OPTION (Elwyn 2003)) were not included in any studies. The usefulness of interventions aimed primarily at patients to help facilitate their role in initiating and making shared decisions remains unknown.

Quality of the evidence

We graded the quality of the evidence as moderate or low for all outcomes. All cluster-level randomised controlled trials (RCTs) used a method of sequence generation aimed at minimising chance between-group imbalance. All study participants (clusters) were randomised after they were enrolled and prior to group allocation to minimise selection bias. Blinding was not possible because of the nature of the interventions. We considered only two studies to have substantial loss to follow-up (Altiner 2007; Légaré 2012). Altiner 2007 did not conduct an intention-to-treat (ITT) analysis although they explored the effects of differential missing values in cluster-level sensitivity analysis. An ITT analysis was not reported by Légaré 2012.

Pooled studies for the primary outcome, antibiotic prescribing, were limited by the diversity in adjusted effect estimates reported and resulted in a low number of studies in each presented comparison (see Analysis 1.3; Analysis 1.4; Analysis 1.5). This was surmounted by calculating a risk ratio (RR) (using the design effect to adjust for clustering) for meta-analysis (see Analysis 1.1; Analysis 1.2; Analysis 1.6), which results in some loss of precision, although it is still robust (and more conservative at least). Similarly, metaanalysis could not be performed for several clinically important secondary outcomes due to variance in effect estimates reported or measurement differences, which resulted in only a small number of trials being included for patient satisfaction (see Analysis 1.7). The low number of trials in addition to the presence of considerable heterogeneity in the longer-term reduction in antibiotic prescribing suggests that the overall pooled results and meaningful exploration of heterogeneity was limited and should be interpreted with caution. See Summary of findings for the main comparison.

Potential biases in the review process

Combining trials under a common effect estimate (RR) for antibiotic prescribing in the longer term (≥ 12 months) required us to impute intra-class correlation coefficients for two studies (Butler 2012; Cals 2013), from similar studies, so that the design effect for adjustment of clustering effects could be calculated. The results for these outcomes should be interpreted with caution.

Agreements and disagreements with other studies or reviews

Other systematic reviews have assessed clinician- and/or patientoriented interventions to influence antibiotic prescribing for acute respiratory infections (ARIs) in primary care (Andrews 2012; Arnold 2005; Boonacker 2010; Ranji 2008; Thoolen 2013; van der Velden 2012; Vodicka 2013). Meaningful comparisons about the relative effectiveness of studies is limited by the diversity in study designs, interventions and outcome measures. Two reviews concluded that multiple component interventions that provided education to healthcare professionals and patients were most often

effective in reducing antibiotic use for respiratory tract infections (Arnold 2005; van der Velden 2012). Multi-faceted interventions and computer strategies aimed at healthcare professionals most effectively reduced antibiotic prescribing in children with upper respiratory tract infections (Boonacker 2010). Provision of patient information alone (Thoolen 2013), or in addition to physician education (van der Velden 2012), appears to offer only moderate or little additional benefit, respectively. However, reviews exclude many recent high quality intervention trials incorporating patient information materials and training explicitly aiming to facilitate shared decision making. Two reviews found that educational interventions directed at parents and/or caregivers were effective in modifying consulting behaviour and antibiotic use for children with ARIs, and may be more successful when they engage children (Andrews 2012; Vodicka 2013). Interventions were also more successful when they were delivered prior to the consultation and focused on specific symptoms (Andrews 2012). Several reviews concluded that a reduction in antibiotics was not at the expense of adverse clinical outcomes (Ranji 2008), or patient satisfaction (Andrews 2012; Ranji 2008; Thoolen 2013). Previous reviews have raised the importance of a patient-centred approach to help patients adopt a more active role in decision making about antibiotics for ARIs (Thoolen 2013), and communication skills training for physicians has been highlighted as a promising intervention element (van der Velden 2012).

AUTHORS' CONCLUSIONS

Implications for practice

Interventions that aim to facilitate shared decision making reduce antibiotic prescribing for acute respiratory infections (ARIs) in primary care in the short term by a relative risk reduction of almost 40% compared with usual care, without an increase in patient-initiated re-consultations for the same illness or a decrease in patient satisfaction. There is insufficient evidence that the effect may be sustained in the medium to longer term (~ one to three years). Whether the reduction in antibiotic prescribing achieved is sufficient, or sustained long enough, to reverse community-level resistance trends is not known as this was not measured in the included studies. We graded the quality of the evidence as moderate or low for all outcomes. The variety in the interventions and training components studied has important implications for knowing which intervention components should be used in clinical practice, or how best to adapt successful programmes to other primary care environments with different practice characteristics or access to financial and core support resources.

Implications for research

The addition of future trials into this systematic review may allow greater precision of the effects of shared decision making and an opportunity to explore reasons for the heterogeneity of the results. Evaluation of intervention adherence and fidelity (the degree to which the intervention was delivered as intended) should be incorporated into new studies. Further long-term follow-up of included studies would also provide greater certainty regarding the maintenance of intervention effects. Further research should also aim to determine which aspects of these interventions provide the greatest benefit to adapt programme implementation and uptake in diverse clinical settings. Research will also need to establish the link between a reduction in antibiotic prescribing for ARIs in primary care and the reversal in community-level antibiotic-resistance trends, to validate the usefulness and sustainability of programmes. Furthermore, while the interventions in studies are principally aimed at developing general practitioners' (GPs') communication skills to facilitate shared decision making, there appears to be scope to pursue ways of involving healthcare consumers in the design, planning and delivery of interventions to promote shared decision making for ARIs in primary care. Finally, although not relevant to the present review, the cost-effectiveness of establishing shared decision making training programmes in primary care to reduce antibiotic use for ARIs requires further research interest.

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* Indicates the major publication for the study

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Altiner 2007

Methods	Study design: cluster-randomised controlled trial Unit of randomisation: general practitioner (GP) Trial duration: November 2003 to March 2005 Recruitment: 2036 GPs from 9 regions in North-Rhine and Westphalia-Lippe, Germany, invited to participate (blinded to the primary outcome); of 239 GPs willing to participate and receiving baseline materials, 104 completed reliable baseline study documentation and were randomised (10 practice partners randomised as pairs) into intervention (GPs = 52, patients = 1389) and control groups (GPs = 52, patients = 1398) Methods of data collection: GPs recorded all consecutive and eligible patients during each documentation period on study specific paper documentation Data collection time points: 3 documentation intervals of 6 weeks each: baseline (before randomisation), and 6 weeks and 12 months post-intervention Length of follow-up: 12 months
Participants	GPs documented all consecutive and eligible patients: ≥ 16 years of age with an initial episode of acute cough (without prior episode < 8 weeks) and could comprehend German Exclusion: patients with underlying chronic lung diseases (e.g. asthma, chronic obstructive pulmonary disease), immune deficiency or malignant diseases
Interventions	Brief intervention name: complex, peer-led, educational intervention Recipients: GPs and patients (passive) Providers: GP peers were trained to provide (in 3 sessions) the outreach visits in clinics during normal working hours (methods of training these GP peers were not specified) Health professional components: focused on antibiotic 'misunderstanding' during a con- sultation, and aimed to motivate GPs to change attitudes to communication and em- power patients. Peers addressed GP beliefs and attitudes by exploring and evaluating GPs 'opposite' motivational background using a standardised dialogue script and com- munication techniques derived from the elaboration likelihood model. Aspects of the intervention were also informed by previous qualitative work Patients: waiting room poster and leaflet focusing on the patients' role within the antibi- otic misunderstanding (e.g. GP perceptions that patients expect an antibiotic) and also brief evidence-based information about acute cough and antibiotics to enable patients to raise and clarify issues and make a joint decision about antibiotic use with their doctor Materials: waiting room poster and leaflet (patient only); script used by GP peers Mode of delivery: face-to-face (GPs) and waiting room posters and leaflets (patients) Duration and intensity: 1 peer outreach visit per GP (duration not specified) Comparator: nil active comparator; GPs provided usual care
Outcomes	Primary: rate of antibiotic prescriptions per acute cough and by GP (study specific paper documentation) Secondary: nil
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: no

Altiner 2007 (Continued)

	Trial registration: yes Ethics approval: yes							
Risk of bias								
Bias	Authors' judgement	Support for judgement						
Random sequence generation (selection bias)	Low risk	Program-generated complete randomisa- tion list						
Allocation concealment (selection bias)	Low risk	Not described. However, GPs recruited prior to randomisation						
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (complex peer-led educational intervention)						
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Participating GPs sent data to researchers. Each patient was assigned a unique iden- tification number that could be connected with the patient only by the participating GP						
Incomplete outcome data (attrition bias) All outcomes	High risk	Randomised: 104 GPs (intervention = 52, 1389 patients; control = 52, 1398 patients) 6 weeks post-intervention: 86 GPs (inter- vention = 42 (80%), patients = 1021; con- trol = 44 (84%), patients = 1143) 12 months post-intervention: 61 GPs (in- tervention = 28 (54%); 787 patients; con- trol = 33 (63%); 920 patients) 17% (18/104) dropped out at 6 weeks and 41% (43/104) by 12 months (reasons for GPs' exclusion from analysis: poor data quality or did not return data) Cluster-level sensitivity analysis performed to explore effect of differential missing val- ues						
Selective reporting (reporting bias)	Low risk	All indicated results reported. Prospec- tive trial registration: Projektdatenbank Versorgungsforschung NRW, ID: 90/34/ CHANGE						
Other bias	Unclear risk	Sample size (power) calculation: yes. Sample size calculated on number of patients to detect a 10% difference in 6-month prescription rates (50% control, 40% intervention). Allowing for 20% drop-out rate,						

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Altiner 2007 (Continued)

Briel 2006

Methods	Study design: cluster-randomised controlled trial Unit of randomisation: general practitioner (GP) Trial duration: January to May 2004 Recruitment: 345 eligible GPs (criteria undefined) from 2 Swiss cantons (Basel-Stadt and Aargau), where self dispensation of drugs is not allowed. 30 GPs (providing written consent by 1 December 2003) were randomised to limited or full intervention groups (15 GPs each); the remaining 15 GPs (providing written consent by 1 January 2004) formed the non-randomised control group Methods of data collection: baseline data for eligible GPs obtained from the registry of the Swiss Medical Association; GPs recorded patient baseline data; medical students conducted standardised patient follow-up interviews at 7 and 14 days by telephone; pharmacists faxed all prescriptions with study labels to the study centre Length of follow-up: 14 days
Participants	GPs recruited all consecutive and eligible adult patients: \geq 18 years with symptoms of acute infections of the respiratory system (first experienced within the previous 28 days; including common cold, rhinosinusitis, pharyngitis, exudative tonsillitis, laryngitis, otitis media, bronchitis, exacerbated COPD or influenza) <i>Exclusion:</i> patients with pneumonia, not fluent in German, with intravenous drug use or psychiatric disorders, and not available for phone interviews or unable to give written informed consent

Interventions	Brief intervention name: patient-centred co Recipients: GPs Providers: unclear Health professional components: evidence-b based on existing US guidelines, adapted experts) presented as a booklet and in a 2-hot centred communication seminar in small g personal feedback by phone prior to the sta how to understand and modify patients' com for ARIs. Physicians were taught to practi to emotional clues and tailor information attitudes and readiness for behaviour chang DiClemente 1992) Patient components: nil Materials: evidence-based guidelines for the (http://www.bice.ch/publications/reports) Mode of delivery: booklet and face-to-face s munication seminar Duration and intensity: GPs attended 1 x 2 seminar and 1 x 6-hour small group interact Comparator 1 (Limited intervention): evide and in a 2-hour interactive seminar alone Comparator 2 (Non-randomised control): u	assed guidelines (developed by 3 trial authors to local conditions and reviewed by local ur interactive seminar, plus a 6-hour patient- roups (number not defined) and 2 hours of rt of the trial. Training aimed to teach GPs cepts and beliefs about the use of antibiotics ce elements of active listening, to respond given to patients. GPs identified patients' re using a theoretical model (Prochaska and e treatment of ARIs distributed as a booklet mall-group interactive patient-centred com- chour interactive evidence-based guidelines ive patient-centred communication seminar ence-based guidelines presented as a booklet	
Outcomes	Primary: antibiotic prescriptions dispensed by pharmacists < 2 weeks following initial consultation (prescriptions with study labels faxed by pharmacists to the study centre) Secondary: rates of different diagnoses of respiratory infections (GP records) Adherence to guidelines for antibiotic prescription (GP records) Days with restrictions from respiratory infection (patient follow-up interview at 7 and 14 days) Days off work (patient follow-up interview at 7 and 14 days) Re-consultation rates (patient follow-up interview at 7 and 14 days) Patient satisfaction (Patient Satisfaction Questionnaire; patient follow-up interview at 7 and 14 days) Patient enablement (Patient Enablement Instrument; patient follow-up interview at 7 and 14 days) Other: serious adverse events (independent monitoring board review of serious adverse events that occurred < 28 days of study enrolment)		
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: no Trial registration: not stated Ethics approval: yes		
Risk of bias			
Bias	Authors' judgement	Support for judgement	

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Briel 2006 (Continued)

Random sequence generation (selection bias)	Low risk	Computer-generated list created by an in- dependent institution
Allocation concealment (selection bias)	Low risk	Allocation to either intervention was con- cealed. However, method not stated. How- ever, GPs recruited prior to randomisation
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of general practitioners and trial staff reported. As this trial had 3 arms (2 intervention arms where the intervention in each involved a seminar and distribution of evidence guidelines; 1 usual care arm), it is possible that the GPs in the intervention arms would not have known which inter- vention group they were in
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Medical students, blinded to the goal of the trial, were trained to conduct standardised follow-up interviews at 7 and 14 days by phone Prescriptions with study labels faxed by pharmacists to the study centre were checked and entered into the database by a person blinded to the intervention group Trial authors assessed adherence of all pre- scriptions to guidelines independently and blinded to the intervention group
Incomplete outcome data (attrition bias) All outcomes	Low risk	GPs randomised into limited intervention (GPs = 15; patients = 293) and full in- tervention groups (GPs = 15; patients = 259); 15 GPs (285 patients) participated as non-randomised controls (data not ex- tracted). All GPs completed the trial. There were 290, 253 and a convenience sample of 93 patients (stratified by physician), re- spectively, interviewed at 7 days; and 287, 245 and 92 patients interviewed at 14 days. Reasons for loss to follow-up reported
Selective reporting (reporting bias)	Low risk	All indicated results reported. Trial registra- tion or published trial protocol not stated
Other bias	Unclear risk	Sample size (power) calculation: yes ITT or per protocol analysis: ITT Intraclass correlation (co-efficient) re- ported: 4.0% and a design effect of 1.6% Low study baseline prescribing rates - full

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Briel 2006 (Continued)

		intervention (13.5%), limited intervention (15.7%) and non-randomised control (21. 4%) Highly motivated GPs: recruitment coin- cided with introduction of a new nation- wide computer-based reimbursement sys- tem and due to increased workload partic- ipating GPs considered to be highly moti- vated
Butler 2012		
Methods	Study design: cluster-randomised controlled trial Unit of randomisation: general practices Trial duration: conducted during 2007 and 2008 Recruitment: 212 general practices approached at random from 454 eligible practices in Wales, UK. 102 practices expressed interest to participate; 70 recruited; 68 practices (-480,000 patients) randomised to intervention or control groups (34 each) Methods of data collection: routine administrative systems (see 'Outcomes') Data collection time points: total numbers of dispensed oral antibiotic items (primary) and hospital admissions for possible RTIs and their complications (secondary): rate per 1000 patients for the year after the intervention practices were exposed to the interven- tion; re-consultation for RTIs: (secondary; 7, 14 and 31 days after initial consultation). Cost data not extracted Length of follow-up: 12 months	
Participants	Clinicians (general practitioners (GPs) and nurse practitioners) and all patients registered with and consulting a participating general practice in Wales (practice list)	
Interventions	Brief intervention name: Stemming the Tide of Antibiotic Resistance (STAR) educational programme: multifaceted flexible blended learning approach to continuing education for clinicians Recipients: clinicians (GPs and nurse practitioners) Providers: web-based modules and practice-based seminar led by a facilitator Health professional components: the programme is a blended learning experience, and based on Social Learning Theory to develop GPs sense of importance about change (the 'why' of change) and confidence in their ability to achieve change (the 'how' of change). The intervention consist of 7 parts (5 online, 1 face-to-face and 1 facilitator-led practice-based seminar): case-scenarios and updated summaries of research evidence and guidelines; reflections on clinical judgement on antibiotic prescribing; a facilitator-led practice-based seminar presenting regional, local and practice-level antibiotic prescribing and resistance data; novel communicative consulting skills and information exchange based on motivational interviewing; personal reflections on clinical practice; web-based forum to share experiences and views; and a booster module completed 6 to 8 months after completion of the initial training to reinforce previously outlined communication skills. GPs had to complete each online learning component before the software would allow them access to the next. The intervention was flexible to allow GPs to access online components and try out new skills with patients at their convenience	

Butler 2012 (Continued)

	Patient components: nil Materials: web-based materials Mode of delivery: interactive web-based modules (including online videos in addition to a facilitator-led practice-based seminar Duration and intensity: not specified Comparator: usual care
Outcomes	Primary: total number of dispensed oral antibiotic items per 1000 registered patients for the year after practices were exposed to the STAR programme (Prescribing Audit Reports and Prescribing Catalogues; www.nhsbsa.nhs.uk/prescriptions) Secondary: hospital admission rates for possible RTIs and their complications per 1000 registered patients for the year after practices were exposed to the STAR programme. (Patient Episode Database for Wales); and practice re-consultation rates (for patients with RTIs, practice re-consultation rates were identified using diagnostic READ codes recorded by the general practitioner over 7, 14 and 31 days after an initial consultation) Costs data not extracted
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: yes Trial registration: yes Ethics approval: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was conducted once all practices were recruited and all participat- ing physicians had provided written con- sent. Dynamic block allocation was used to achieve balance between groups of prac- tices for the potential confounders of previ- ous rate of antibiotic dispensing (averaged over the past year), practice size (number of whole time equivalent staff at recruitment), and proportion of clinicians in the practice registered for the study. The practices were divided into 3 sets of 24, 22 and 22 prac- tices; within each set we generated all pos- sible allocations into 2 groups and selected the 1000 allocations within each set with the best balance with respect to the speci- fied confounders. The independent statisti- cian on the trial steering committee selected 1 allocation at random for each set and ran- domly assigned intervention or control to the 2 groups in each set to construct the final allocation

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Butler 2012 (Continued)

Allocation concealment (selection bias)	Low risk	Clinicians and researchers were blinded to group allocation until after randomisation
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (multifaceted intervention programme)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Data on antibiotic dispensing, hospital ad- missions and re-consultations were col- lected through routine administrative sys- tems that were not influenced by the study research process
Incomplete outcome data (attrition bias) All outcomes	Low risk	68 practices (~480,000 patients) ran- domised to intervention (34 practices; 137 GPs, 2 nurse practitioners) or control (34 practices; 122 GPS, 2 nurse practitioners) groups. 2 practices (one in each group; in- cluding 12 intervention GPs and 7 con- trol GPs) withdrew after randomisation but were included in the ITT analyses
Selective reporting (reporting bias)	Low risk	All indicated results reported. Published trial protocol available
Other bias	Low risk	Sample size (power) calculation: yes ITT or per protocol analysis: ITT analysis for primary outcome

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Cals 2009

Methods	 Study design: cluster-randomised controlled trial (factorial design) Unit of randomisation: general practices (cluster of 2 general practitioners (GPs) per practice) Trial duration: conducted during the winters of 2005 to 2006 and 2006 to 2007 Recruitment: 54 general practices within a large suburban region of the Netherlands were assessed for eligibility; 20 eligible general practices (with 2 participating GPs per practice) were randomised into groups of 10 practices per intervention (resulting in 4 trial arms of 5 general practices and 10 GPs): use of C-reactive protein (CRP) testing; training in enhanced communication skills; use of CRP and training in enhanced communication skills; control (usual care) Methods of data collection: antibiotic prescribing and re-consultation data obtained from patient medical records. Patients rated symptoms (cough, phlegm, shortness of breath, disturbance of daily activities, sleeping problems and generally feeling unwell), satisfaction and enablement, on a 28-day daily diary validated for use in a RCT on management of LRTI in primary care
Participants	General practitioners recruited sequential eligible adults within regular consultation hours during the winters of 2005 to 2006 and 2006 to 2007 Eligibility: suspected lower respiratory tract infection (LRTI) with a cough lasting < 4 weeks together with1 focal and 1 systemic symptom
Interventions	Brief intervention name: enhanced communication skills training Recipients: GPs Providers: seminars led by a moderator Health professional components: enhanced communication skills training involved 1 x 2-hour training seminar at a central location, preceded and followed by consulting with simulated patients in routine surgeries and peer-review of transcripts. The moderator- led seminar on shared decision making (within 1 week of simulated patient consultation) comprised GPs' reflection on simulated patient transcript, current views and insights on LRTI (highlighting contrast between research and practice), outline of elicit-provide- elicit framework (elicit patient's main worries and expectations and conveying the balance of possible antibiotic benefits and harms, provide information relevant to the patients' individual understanding and interest, and elicit patients' interpretation about what has been said and done and discusses implications for help seeking behaviour), videos presenting practice-based examples and GPs identifying specific aspects during their consultations that need most attention Patient components: nil Materials: desk reminder for GPs Mode of delivery: face-to-face seminar and simulated patient consultations with peer- review of transcripts Duration and intensity: 1 x 2-hour moderator-led training seminar; pre- and post- seminar simulated patient consultations with peer- review of transcripts Comparator 1: C-reactive protein point of care testing (date not extracted) Comparator 2: enhanced communication skills training plus C-reactive protein point of care testing (date not extracted) Comparator 4: usual care (Dutch guidelines for managing acute cough, including diag-

Cals 2009 (Continued)

	nostic and therapeutic advice for lower respiratory tract infection are distributed to all GPs in the Netherlands)
Outcomes	Primary: antibiotic prescribing in the index consultation (medical records) Secondary: antibiotic prescribing during 28 days' follow-up (medical records) Re-consultation (medical records) Clinical recovery data not extracted Patients' satisfaction (Likert scale; 28-day daily diary) Patients' enablement (Patient Enablement Index; 28-day daily diary)
Notes	Funding: yes Conflict of interest: none declared Published trial protocol: yes Trial registration: yes Ethics approval: yes Main comparator reported in this review: communication skills training (n = 201) versus no communication skills training (n = 230)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	General practices randomised into 2 groups of 10 practices per intervention, balanced for recruitment potential, resulting in the 4 trial arms. The balancing factor used for randomisation was the amount of GP's consultation time (expressed as full time equivalent (FTE)) that the practice was contributing to the study (which equated to between1 and 2 FTEs for clinical con- tact time. The randomisation was balanced for those with 1.5 or less FTEs and those with more than 1.5 FTEs
Allocation concealment (selection bias)	Low risk	All practices and general practitioners were recruited before randomisation
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (due to the nature of the in- tervention)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not described
Incomplete outcome data (attrition bias) All outcomes	Low risk	20 practices (40 GPs) randomised to each of the 4 trial arms (5 practices, 10 GPs each) and recruited 431 patients. 37 GPs com-

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Cals 2009 (Continued)

		pleted the trial (3 left on maternity leave in the enhanced communication skills group) . All patients (100%) had data for the pri- mary outcome, 90% (mean) had 28-day diary data For the communication skills training group (10 GPs, 84 patients), there was 100% prescribing data and 88% returned diaries
Selective reporting (reporting bias)	Low risk	All indicative results reported. Published study protocol. Prospective trial registra- tion
Other bias	Low risk	Sample size (power) calculation: yes ITT or per protocol analysis: the primary analysis was ITT

Cals 2013

Methods	Study design: 3.5 year follow-up of a cluster-randomised controlled trial (factorial design) (Cals 2009) Trial duration: 3.5 years (mean 3.67 years) Recruitment: patients recruited in the winter periods from September 2005 until March 2007 (Cals 2009), were observed until July 2010 Methods of data collection: medical records Data collection time points: recorded consultations for RTI from original 28-day follow- up period until July 2010 (follow-up period); recorded consultation for RTI for the exact same period preceding the consultation in which the patient was recruited in the original trial (baseline period). Deceased patients and patients that moved practices and whose medical records could not be retrieved were excluded Length of follow-up: mean 3.67 years
Participants	General practices: see Cals 2009 Patients: of the original 431 patients enrolled in the trial, 379 patients (87.9%) had ac- cessible medical records for the follow-up period. Only data for the enhanced communi- cation training (178) versus no enhanced communication skills training (201) extracted
Interventions	See Cals 2009
Outcomes	Primary outcome: average number of episodes of RTIs during the follow-up period for which patients consulted their physician per patient per year (PPPY) and the proportion of these episodes that resulted in an antibiotic prescription Secondary outcome: nil
Notes	Funding: yes Conflict of interest: RH received travel/lecture funds from Axis-shield (Norway) and Orion Diagnostica (Finland), both manufacturers of C-reactive protein devices Trial registration: yes

Cals 2013 (Continued)

	Ethics approval: yes		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	See Cals 2009	
Allocation concealment (selection bias)	Low risk	See Cals 2009	
Blinding of participants and personnel (performance bias) All outcomes	High risk	See Cals 2009	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Data were extracted, by 2 researchers, from the patients' medical records system. No mention if these researchers were blind to the practices' original allocation	
Incomplete outcome data (attrition bias) All outcomes	Low risk	379 of 431 patients enrolled in the orig- inal trial (87.9%) had accessible medical records for the follow-up period	
Selective reporting (reporting bias)	Low risk	See Cals 2009	
Other bias	Low risk	Sample size (power) calculation: see Cals 2009 ITT or per protocol analysis: see Cals 2009	

Francis 2009

Methods	Study design: cluster-randomised controlled trial Unit of randomisation: general practices Trial duration: October 2006 to April 2008 Recruitment: half of all general practices from 9 local health boards in Wales (n = 147) were randomly selected to be sent study information (the other half were provided infor- mation about a related RCT conducted in parallel); 49 returned a practice agreement and were randomised. 4 primary care research networks in England also recruited practices; 34 returned practice agreement and were randomised. All randomised practise (83) were allocated to intervention (41 practices; 30 recruited patients; patients = 274) or control (42 practices; 31 recruited patients; patients = 284) Methods of data collection: baseline data (age, duration of illness, symptoms) collected by GPs. Follow-up via a telephone administered questionnaire (or self completion ques- tionnaire contact unsuccessful by telephone) with child's parent or guardian Data collection time points: index consultation and 14 days after recruitment Length of follow-up: 14 days
Participants	Participating clinicians recruited sequential eligible children (6 months to 14 years) consulting with a respiratory tract infection (cough, cold, sore throat, earache for 7 days or less) and their parents <i>Exclusion:</i> children with asthma and those with serious ongoing medical conditions such as malignancy or cystic fibrosis
Interventions	Brief intervention name: interactive booklet on respiratory tract infections in children for use within the consultation and provided as a take home resource Recipients: parents and clinicians Providers: not stated Health professional components: the online training described the content and aims of the booklet, and encouraged its use within the consultation to facilitate the use of certain communication skills, mainly exploring the parent's main concerns, asking about their expectations, and discussing prognosis, treatment options and any reasons that should prompt re-consultation Patient components: use of the booklet in the consultation and as a take home resource Materials: 8-page interactive booklet (see www.whenshouldiworry.com) Mode of delivery: 8-page interactive booklet and online training for clinicians in use of the booklet Duration and intensity: not stated Comparator: usual care (clinicians were asked to conduct consultations in usual manner)
Outcomes	Primary: re-consultation (primary or secondary care) during the 2 weeks after the index consultation (telephone administered questionnaire) Secondary: antibiotic prescriptions (telephone administered questionnaire) Antibiotic consumption (telephone administered questionnaire) Future consulting intention (telephone administered questionnaire) Parental satisfaction with the index consultation (5-point Likert; telephone administered questionnaire) Parental enablement (modified Patient Enablement Instrument; telephone administered questionnaire) Pareception of the usefulness (value) of the information received during the index con- sultation (5-point Likert; telephone administered questionnaire)

Francis 2009 (Continued)

	Parental reassurance (3-point Likert; telephone administered questionnaire)	
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: yes Trial registration: yes Ethics approval: yes	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Practices were randomised by a statistician using block randomisation with random block sizes and stratification by practice list size, antibiotic prescribing rate for 2005, and country
Allocation concealment (selection bias)	Unclear risk	It is reported that practices were ran- domised after agreeing to take part, but no other details are provided
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (training and use of an inter- active booklet for use within consultations and as a take home resource)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Telephone interviewers were blinded to treatment group and asked to record any subsequent unblinding of allocation (e.g. parent talking about receiving a booklet) . Interviewers reported becoming aware of participants treatment group in 34/509 (6. 7%) of interviews
Incomplete outcome data (attrition bias) All outcomes	Low risk	83 practices were randomised to inter- vention (41) or control (42) groups; 61 practices, 30 intervention and 31 control practices, recruited 274 and 284 patients, respectively. Primary outcome data were available for 256 patients (93%) in the intervention group (246 completed tele- phone interviews, 10 postal questionnaire returned) and 272 (96%) control group patients (262 completing telephone inter- views, 9 postal questionnaires returned)
Selective reporting (reporting bias)	Low risk	All indicted outcomes reported. Published trial protocol

Francis 2009 (Continued)

Other bias	Low risk	Sample size (power) calculation: yes ITT or per protocol analysis: primary anal- ysis was ITT
Little 2013		
Methods	Unit of randomisation: general Trial duration: October 2010 Recruitment: all general pract proached, and all clinicians (a prescribed antibiotics for respi ity: practices that had not pre- ing and could include > 10 p were selected separately in Ant z (Poland), Southampton (UK ish Society of Family Medicin gions of Europe (north, south enrolled; 246 were randomised protein (CRP) test at point of (n = 61), or in both CRP and Methods of data collection: ca	•
Participants	for RTIs consecutively recruit first 5 with URTI presenting a attending a first consultation clinician believed to be an acut most prominent symptom; an respiratory tract infection (e.g illness) <i>Exclusion:</i> patients with a work embolus, heart failure, oesoph month; inability to provide in	d nurse prescribers in the UK) who prescribed antibiotics ted up to the first 30 patients with LRTI and up to the at each practice. Eligible patients were \geq 18 years of age, for acute cough of up to 28 days' duration or what the te LRTI as the main diagnosis, despite cough not being the ad diagnosis judged by the physician to be an acute upper c, sore throat, otitis media, sinusitis, influenza and coryzal king diagnosis of a non-infective disorder (e.g. pulmonary mageal reflux, or allergy); use of antibiotics in the previous formed consent (e.g. due to dementia, psychosis or severe mmunological deficiencies. Pneumonia was not an exclu-
Interventions	Recipients: GPs Providers: n/a Health professional componen tients' concerns and expectatio course and treatments; agreen guidance about when to re-cor to use during consultations th	unced communication skills training ats: training focused on the gathering of information on pa- ons; exchange of information on symptoms, natural disease ment of a management plan, summing up and providing asult. Physicians were provided with an interactive booklet nat included information on symptoms, use of antibiotics help measures, and when to re-consult. The training was

Little 2013 (Continued)

	supported by video demonstrations of consultation techniques. The Internet modules and materials were translated into the relevant national language and mainly addressed lower respiratory tract infections, although many of the issues were relevant to all respi- ratory tract infections Patient components: interactive booklet used within consultations Materials: interactive booklet for use within consultations Mode of delivery: Internet training supported by video demonstrations of consultation techniques Duration and intensity: not described Comparator: 1. Usual care 2. Training in use of C-reactive protein (CRP) test at point of care (data not extracted for this review)
	3. Both CRP and enhanced communication skills training (data not extracted for this review)
Outcomes	Primary: antibiotic use (index consultation; case-report form) Secondary: new or worsening symptoms defined as re-consultation for new or worsening symptoms < 4 weeks, new signs or hospital admission (review of medical notes) Symptom severity and duration defined as the severity of symptoms in the 2 to 4 days after seeing the physician (case report form; 0 = no problem to 4 = severe problem)
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: no Trial registration: yes Ethics approval: yes ITT or per protocol analysis: ITT analysis

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation of practices was done by 2 study authors, and was achieved by com- puter generation of random numbers, strat- ified by network. Minimisation was ap- plied, on the basis of the proportion of pa- tients prescribed antibiotics from the base- line audit, the number of participating physicians per practice, and the number of patients recruited
Allocation concealment (selection bias)	Low risk	Physicians and patients were unaware of initial group allocation
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (due to the nature of the in- tervention)

Little 2013 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	High risk	GPs recorded data on a case-report from, during the index consultation
Incomplete outcome data (attrition bias) All outcomes	Low risk	259 practices enrolled and provided base- line data (6771 patients); 13 practices re- cruited < 10 patients each) were excluded Remaining were 246 practices randomised to CRP (62), enhanced communication training (61), both interventions combined (62), or usual care (61) Antibiotic prescription documentation was available for 58 CRP practices (1062 pa- tients), 55 (90%) enhanced communica- tion skills practices (1170 patients), 62 combined intervention practices (1162 pa- tients) and 53 (87%) usual care practices (870 patients). Reasons for exclusion were reported as recruiting no patients
Selective reporting (reporting bias)	Low risk	All indicated outcomes are reported
Other bias	Low risk	Sample size (power) calculation: not stated ITT analysis: yes

Légaré 2011

Methods	Study design: cluster-randomised controlled trial (pilot) Unit of randomisation: family medicine groups (FMGs) Trial duration: during November 2007 and March 2008 Recruitment: 24 FMGs (group of family physicians who work closely with nurses to offer family medicine services to registered individuals) from the greater urban area of Quebec City, Canada, were invited to participate; 4 participating FMGs were randomised either to a group immediately exposed to the DECISION+ program (n = 2) or to a control group which exposure to DECISION+ program was delayed for 6 months (n = 2) Methods of data collection: self administered questionnaire completed following the consultation at each time point Data collection time points: baseline, following exposure of the intervention group to DECISION+ (~ 6 months), and following delayed exposure of DECISION+ to controls (~ 12 months) Length of follow-up: 12 months
Participants	Eligible general practitioners (no previous participation in an implementation trial of SDM and planned to remain in clinical practice for the trial duration) recruited eligibility patients consulting their GP for an ARI: no age restriction, patients or their guardians had to be able to read, understand and write French and had to give informed consent to participate in the trial Exclusion: patients with a condition requiring emergency care. A research professional

Légaré 2011 (Continued)

	waited in the FMG's waiting room and recruited patients of enrolled FPs during walk- in clinic hours; 15 patients were recruited per GP: 5 at baseline, 5 after the GPs in the experimental group were exposed to DECISION +, and 5 after the FPs in the control group were exposed to DECISION+
Interventions	 Brief intervention name: DECISION+ Recipients: GPs Providers: principal investigators (or co-trainers) Health professional components: DECISION+ is made up of 3 main components 1. Interactive workshops addressed the probability of bacterial versus viral ARIs in primary care, evidence of the benefit/risk of the various treatment options, risk communication techniques and strategies for fostering patient participation in the decision making process. Workshops included videos of simulated patient-GP consultations for each ARI and distinguished 2 approaches (usual care or SDM), and exercises to facilitate group discussion about facilitators and barriers to SDM. GPs were trained to use decision support tools (though video examples and group exercises) developed for each of the 4 targeted ARIs (rhinosinusitis, pharyngitis, bronchitis and acute otitis media) and 1 integrating all 4 ARIs 2. Reminders of expected behaviours: a reminder printed on a letter-sized piece of paper emphasised the use of the decision support tools, reiterated the expected SDM-related behaviours, and highlighted new studies relevant to the pilot trial topics (e.g. new evidence on the risks and benefits of antibiotics). These reminders were mailed to GPs between each workshop. A second reminder was postcards that participants had written to themselves in the last workshop to remind themselves of what they needed to implement in their practice. The research team collected the postcards and mailed them 6 to 8 weeks later 3. Feedback to GPs on the agreement between their decisional conflict scores and that of their first 5 patients Patient components: decision support tools Materials: a booklet summarising the content of the workshop and decision support tools was developed for physician participants and training manuals for the co-trainers Mode of delivery: interactive workshops led by 2 study principal investigators (or co-trainers) and conducted face-to-face in a group forma
Outcomes	Primary: decision about using antibiotics (immediate use, delayed use or no use) (GP/ patient; self administered questionnaire) Secondary: Perception of the quality of the decision (GP/patient; single item on a 10-point Likert scale; self administered questionnaire) Decisional conflict (GP/patient; Decisional Conflict Scale) Patients' intention to engage in SDM in future consultations concerning antibiotics for ARIs (3-item, 7-point Likert scale; self administered questionnaire) GPs' intentions to engage in SDM and comply with clinical practice guidelines regarding prescribing antibiotics for ARIs (3-item, 7-point Likert scale) Decision Regret Scale (patients; telephone interview; 2 weeks following consultation)

Légaré 2011 (Continued)

	Perception of health changes since the consultation (patients; telephone interview; 2 weeks following consultation) Number of prescriptions filled by patients covered by Quebec's public drug insurance plan (Regie de l'Assurance-Maladie du Quebec medication claims database) (during the 3 months preceding baseline and during the 3 months after FPs in the experimental group were exposed to DECISION+) Script concordance test (probes whether respondents' knowledge is efficiently organised to take appropriate clinical action by placing respondents in written, but authentic, clin- ical situations in which they must interpret data to make decisions. It measures the con- cordance between respondents' scripts and the scripts of a panel of experts (administered to GPs at each data collection point)
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: yes Trial registration: not reported Ethics approval: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A biostatistician simultaneously randomised all FMGs using Internet-based software
Allocation concealment (selection bias)	High risk	A biostatistician allocated FMGs to groups using Internet-based software. There was concealed allocation of the Fam- ily Medicine Groups, but not the family physicians
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (multiple-component, con- tinuing professional development pro- gramme in shared decision making)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Codes were attributed to the trial groups and the bio-statistician analysed the data blindly. Team members accessed the codes only after having completed the analyses and interpreting the results
Incomplete outcome data (attrition bias) All outcomes	Low risk	4 FMGs randomised to intervention (2; GPs = 18; patients = 245) or control groups (2; GPs = 15; patients = 214). 3/33 (9%) GPs dropped out of the trial 20/245 pa- tients in the intervention group and 14/ 214 controls could not be contacted over the 2-week follow-up

Légaré 2011 (Continued)

Selective reporting (reporting bias)	Low risk	All indicated outcomes reported. Published trial protocol
Other bias	Low risk	Sample size (power) calculation: no Primary analysis was ITT
Légaré 2012		
Methods	Study design: cluster-randomised controlled trial Unit of randomisation: family practice teaching units Trial duration: July 2010 to April 2011 Recruitment: the network of 12 family practice teaching units in 6 regions of Quebec, Canada, were randomised to intervention (6) or control (6) groups Methods of data collection: following the consultation, patients and GPs independently completed self administered questionnaires (primary and secondary outcomes). 2 weeks later, a telephone follow-up interview was conducted by a research assistant (secondary outcomes) Data collection time points: immediately following consultation and 14 days	
Participants	GPs, including physician teachers and residents, who provide care in the walk-in clinics of the 12 family practice teaching units. GPs participating in the pilot trial (Légaré 2011) or those not expecting to practice in the teaching unit during the trial period were excluded. Patients with symptoms suggestive of an ARI were recruited by a research assistant in the waiting room prior to consultation with a physician. Eligible patients were adults (and children who were accompanied by a parent/legal guardian) with a diagnosis of ARI (e.g. bronchitis, otitis media, pharyngitis or rhinosinusitis) and for which the use of antibiotics was subsequently considered either by the patient or physician during the visit. The patient, parent or legal guardian had to be able to read, understand and write French	
Interventions	 Brief intervention name: DECISION+2 shared decision making program Recipients: GPs Providers: trained facilitators Health professional components: an online tutorial comprised of 5 modules addressing key components of the clinical decision making process about antibiotic treatment for ARI in primary care: introduction to shared decision making and ARIs, estimating diagnostic probabilities for ARIs, therapeutic options, effective strategies to communicate risk and benefits, identify patients' values and preferences; and use of decision support tools that promote shared decision making. Participants had 1 month to complete the online tutorial. The on-site facilitator-led interactive workshop aimed to help physicians review and integrate the concepts they acquired during the online training Patient components: decision support tools Materials: both the online tutorial and workshop included videos, exercises and decision aids to help physicians communicate to their patients the probability of a bacterial acute respiratory infection and the benefits and harms associated with the use of antibiotics Mode of delivery: online tutorial and facilitator-led interactive workshop Duration and intensity: 2-hour online tutorial followed by a 2-hour on-site interactive 	

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Légaré 2012 (Continued)

	workshop Comparator: usual care
Outcomes	 Primary: proportion of patients who decided to use antibiotics immediately after consultation (GP and patient self administered questionnaire) Secondary: decisional conflict (GP/patient; Decisional Conflict Scale) Perception that shared decision making occurred (GP/patient; modified Control Preference Scale) Quality of decision made (GP/patient; single question Likert scale) Adherence to the decision (patient; single-item asking if decision made was maintained) Repeat consultation (for the same reason) (patient) Decisional Regret (patient; Decisional regret Scale) Quality of life (patient; SF-12) Intention to engage in SDM in future consultations regarding the use of antibiotics for ARIs (patients; questions based on Theory of Planned Behaviour) Intentions to engage in shared decision making (GP) Intention to adhere to clinical practice guidelines (GP) Preferred role in decision making (Control Preference Scale)
Notes	Funding: yes Conflict of interest: none disclosed Published trial protocol: yes Trial registration: yes Ethics approval: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A biostatistician used Internet-based soft- ware to simultaneously randomise all 12 family practice teaching units to either the intervention group (DECISION+2) or control group. The teaching units were stratified according to rural or urban loca- tion
Allocation concealment (selection bias)	Unclear risk	The family practice teaching unites were recruited prior to randomisation, but it is not clear when the physicians in the units were recruited
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (due to the nature of the in- tervention and the self administered out- comes)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Statistical analysis was performed by a statistician who was unaware of the teach- ing unit allocations

Légaré 2012 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	12 family practice teaching units ran- domised; 9 participated in the study and all clusters completed the trial
Selective reporting (reporting bias)	Low risk	All indicated outcome reported. Prospec- tive trial registration. Published trial proto- col
Other bias	Unclear risk	Sample size calculation: yes ITT or per protocol analysis: not stated

Welschen 2004

Methods	Study design: cluster-randomised controlled trial Unit of randomisation: GP peer review group Trial duration: 2000 to 2002 Recruitment: general practitioners' (GP) peer review groups, with collaborating phar- macists (which aim to promote rational prescribing through audit and feedback), in the region of Utrecht, Netherlands, if the group consisted of ≥ 4 GPs and all agreed to participate Methods of data collection: during a 3-week period during 2000 and 2001 Data collection time points: index consultation Length of follow-up: nil
Participants	Primary care setting type: recruited from general practitioner (GP) peer review groups General practitioners: 100 GPs Patients: all registered patients presenting with acute symptoms of the respiratory tract *Relatively low prescription rates in the Netherlands
Interventions	 Brief intervention name: multiple intervention Recipients: GPs and patients Providers: GP peer facilitators Health professional components: a) Group education meeting (jointly led a GP and pharmacist in each peer review group) included a review of previous years claims data, discussion of evidence-based medicine and communication of evidence for treatment benefit and risk to inform group consensus about the indication and first choice of antibiotics per indication (AOM, sinusitis, tonsillitis and acute cough); communication skills training (how to explore patients' worries and expectations and to inform patients about the natural course of the symptoms, self medication and alarm symptoms). GPs received a summary of their group's guidelines by mail 1 week after the meeting, and received the results of the baseline measurement (to reinforce the consensus reached) after 2 months b) Monitoring and feedback on prescribing behaviour (6 months post-intervention) based on insurance claims data comparing the period after the intervention (March to May 2001) with the same period before the intervention (March to May 2001) with the same period before the intervention (March to May 2000). Volumes of different kinds of antibiotics and the extent to which prescribed antibiotics were in line with the consensus about first choice antibiotics were presented at practice level c) Group education for assistants of GPs and pharmacists attended a 2-hour group

Welschen 2004 (Continued)

	education session informing them about Dutch guidelines for GPs, followed by skills training in educating patients Patient components: education material for patients consisted of a brochure and accom- panying posters (also translated into Turkish and Arabic) available in waiting rooms of intervention group general practices, pharmacies and municipal health services, aiming to inform patients about the self limiting character of most respiratory tract symptoms, self medication and serious symptoms ("alarm signals") necessitating a consultation with the GP Materials: consensus guidelines for GPs and education material for patients Mode of delivery: GP and pharmacist-led group education meeting for GPs and assis- tants, and patient education brochure and posters Duration and intensity: 1 x group education meetings for GPs (duration not stated) and 1 x 2-hour group education meetings for assistants Comparator: usual care
Outcomes	Primary: proportion of practice encounters for acute symptoms of the respiratory tract for which antibiotics were prescribed (patient records) Patient satisfaction (self reported questionnaire; 1 = very dissatisfied to 5 = very satisfied) Secondary: administrative claims data (from regional health insurance company, Agis, over the period 2000 to 2002) (March to May, 2000 and March to May, 2001)
Notes	Funding: yes Conflict of interest: none declared Published trial protocol: not reported Trial registration: not reported Ethics approval: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The 12 peer review groups were allocated to groups A or B. All possible composi- tions of groups A and B were considered and the option chosen of those groups re- sulting in comparability between group A and B in groups with a high or low vol- ume of antibiotic prescribing (above or be- low the median), rural or urban working groups, and number of general practition- ers per group (above or below the median) . MMK, who was blinded to the composi- tion of the groups, flipped a coin to deter- mine whether group A became the inter- vention or control group
Allocation concealment (selection bias)	Low risk	Not stated. However, practices recruited prior to randomisation

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Welschen 2004 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	High risk	Not possible (multiple intervention)		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Research assistants blinded to the interven- tion status of the practices extracted infor- mation from patient records (age, sex, diag- noses, antibiotic prescriptions and referrals to hospital doctors) Patient satisfaction questionnaires returned directly to the investigators without being shown to the GP		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Of the 42 of 48 peer-review groups in the Utrecht region that were invited to participate, 30 groups refused or were unable to participate. The 12 remaining peer-review groups were randomised to intervention (6 groups, 46 GPs) or control (6 groups, 54 GPs). All clusters and 89/100 GPs completed the study (intervention = 42, control = 49), with loss to follow-up due to retirement (n = 1), removal outside the region (n = 3), illness (n = 3), motivational problems (n = 2)		
Selective reporting (reporting bias)	Low risk	All indicative results reported		
Other bias	Low risk	Sample size (power) calculation: yes ITT of per protocol analysis: yes		

AOM: acute otitis media ARI: acute respiratory infection COPD: chronic obstructive pulmonary disease CRP: C-reactive protein FP: family physician GP: general practitioner ITT: intention-to-treat LRTI: lower respiratory tract infection n/a: not applicable OTC: over-the-counter RCT: randomised controlled trial RTI: respiratory tract infection SDM: shared decision making URTI: upper respiratory tract infection

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Bourgeois 2010	Shared decision making not explicit or inferred
Gonzales 2013	Shared decision making not explicit or inferred
Pshetizky 2003	Shared decision making not explicit or inferred
Regev-Yochay 2011	Shared decision making not explicit or inferred
Samore 2005	Shared decision making not explicit or inferred
Taylor 2005	Shared decision making not explicit or inferred

Characteristics of ongoing studies [ordered by study ID]

Altiner 2012

Trial name or title	Converting habits of antibiotic prescribing for respiratory tract infections in German primary care - the cluster-randomised controlled (CHANGE-2) trial
Methods	3-arm cluster-randomised controlled trial
Participants	GPs (n = 94) or practice-based paediatricians (n = 94) and their patients (~ 30,000 children and adults) who consult in general practices located in 2 German regions (Baden-Württemberg and Mecklenburg-Western Pomerania) for an ARI
Interventions	Communication training versus communication training and point of care testing (C-reactive protein and rapid antigen detection testing) versus control
Outcomes	Primary: physician antibiotic prescription rate for ARI at 2-year follow-up (post-intervention) derived from data of the statutory health insurance company Secondary: 1. Re-consultation rate 2. Use of medical services 3. Hospital admissions
Starting date	GP and paediatrician recruitment commenced October 2012; patient recruitment over 3 successive winter periods
Contact information	Prof Attila Altiner; Institute for General Practice, Rostock University Medical Center; POB 100888; Rostock 18055 Germany Phone: +49 (0)381 494 2481 Fax: +49 (0)381 494 2482 Email: ifa.sekretariat@med.uni-rostock.de

Altiner 2012 (Continued)

Notes -

ARI: acute respiratory infection

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DATA AND ANALYSES

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Antibiotics prescribed, dispensed or decision to use (short-term, index consultation to ≤ 6 weeks)	8	10172	Risk Ratio (Random, 95% CI)	0.61 [0.55, 0.68]
2 Antibiotics prescribed or dispensed (longer-term, ≥ 12 months)	3	481588	Risk Ratio (Random, 95% CI)	0.74 [0.49, 1.11]
3 Antibiotic prescriptions (index consultation) (adjusted odds ratio)	3	3244	Odds Ratio (Random, 95% CI)	0.44 [0.26, 0.75]
4 Antibiotic prescriptions (index consultation) (adjusted risk ratio)	2	4623	Risk Ratio (Random, 95% CI)	0.64 [0.49, 0.84]
5 Antibiotic prescriptions (index consultation or population rate per unit of time) (adjusted risk difference)	4	481807	Mean Difference (Random, 95% CI)	-18.44 [-27.24, -9. 65]
6 Number or rate of re-consultations (risk ratio)	4	1861	Risk Ratio (Random, 95% CI)	0.87 [0.74, 1.03]
7 Patient satisfaction with the consultation	2	1052	Odds Ratio (Random, 95% CI)	0.86 [0.57, 1.30]

Comparison 1. Shared decision making versus usual care (control)

Analysis I.1. Comparison I Shared decision making versus usual care (control), Outcome I Antibiotics prescribed, dispensed or decision to use (short-term, index consultation to \leq 6 weeks).

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: I Antibiotics prescribed, dispensed or decision to use (short-term, index consultation to \leq 6 weeks)

Study or subgroup	Shared decision making N	Usual care N	log [Risk Ratio] (SE)	Risk Ratio IV,Random,95% Cl	Weight	Risk Ratio IV,Random,95% CI
Francis 2009 (1)	256	272	-0.7867 (0.4212)		1.8 %	0.46 [0.20, 1.04]
Briel 2006 (2)	259	293	-0.1518 (0.2619)	-	4.8 %	0.86 [0.51, 1.44]
L gar 2012(3)	181	178	-0.6931 (0.2606)		4.8 %	0.50 [0.30, 0.83]
L gar 2011(4)	81	70	-0.3738 (0.204)	-=-	7.9 %	0.69 [0.46, 1.03]
Cals 2009 (5)	201	230	-0.6846 (0.19)	+	9.1 %	0.50 [0.35, 0.73]
Altiner 2007 (6)	1021	1143	-0.6983 (0.1666)	-	11.8 %	0.50 [0.36, 0.69]
Little 2013 (7)	2332	1932	-0.3711 (0.1251)	-	20.9 %	0.69 [0.54, 0.88]
Welschen 2004 (8)	905	818	-0.4806 (0.0917)	-	38.9 %	0.62 [0.52, 0.74]
Total (95% CI)	5236	4936		•	100.0 %	0.61 [0.55, 0.68]
Heterogeneity: $Tau^2 = 0.0$;	Chi ² = 6.62, df =	7 (P = 0.47); I ²	=0.0%			
Test for overall effect: $Z =$	8.63 (P < 0.0000)				
Test for subgroup difference	es: Not applicable					
			(0.005 0.1 1 10 200)	
			Shared	decision making Usual care		

(1) Reported intra-class correlation co-efficient (ICC) = 0.24. Design effect and effective sample size calculated.

(2) Reported intra-class correlation co-efficient (ICC) = 0.04. Design effect and effective sample size calculated. Actual sample denominator used to calculate risk ratio. Actual sample denominators reported in forest plot.

(3) Adjusted for cluster design, baseline values and patient age group (for analyses at teaching unit and physician levels).

(4) Reported intra-class correlation co-efficient (ICC) = 0.02. Design effect and effective sample size calculated.

(5) Reported intra-class correlation co-efficient (ICC) = 0.12. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in forest plot.

(6) Reported intra-class correlation co-efficient (ICC) = 0.20. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in forest plot.

(7) Adjusted for baseline prescribing and clustering by physician and practice, age, smoking, sex, major cardiovascular or respiratory comorbidity, baseline symptoms, crepitations, wheeze, pulse higher than 100 beats per minute, temperature higher than 37.8 degrees Celcius, respiratory rate, blood pressure, physician's rating of severity, and duration of cough.

(8) Reported intra-class correlation co-efficient (ICC) = 0.09. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in forest plot.

Analysis 1.2. Comparison I Shared decision making versus usual care (control), Outcome 2 Antibiotics prescribed or dispensed (longer-term, ≥ 12 months).

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: 2 Antibiotics prescribed or dispensed (longer-term, \geq 12 months)

Study or subgroup	Shared decision making N	Usual care N	log [Risk Ratio] (SE)	Risk Ratio IV,Random,95% Cl	Weight	Risk Ratio IV,Random,95% Cl
Cals 2013 (1)	178	201	-0.3817 (0.2188)	-	27.2 %	0.68 [0.44, 1.05]
Altiner 2007 (2)	787	920	-0.5625 (0.1384)	-	33.4 %	0.57 [0.43, 0.75]
Butler 2012 (3)	239802	239700	-0.0254 (0.0053)	+	39.4 %	0.97 [0.96, 0.99]
Total (95% CI) Heterogeneity: Tau ² = 0 Test for overall effect: Z Test for subgroup differe	= 1.47 (P = 0.14)	,	015); I ² =89%	•	100.0 %	0.74 [0.49, 1.11]
				.005 0.1 I I0 200 lecision making Usual care)	

(1) Design effect (2.08) imputed from Cals (2009). Effective sample size calculated by imputing intra-class correlation co-efficient reported by Cals 2009 (0.12). Actual sample denominators reported in forest plot.

(2) Reported intra-class correlation co-efficient (ICC) = 0.20. Design effect and effective sample size used to calculate risk ratio. Actual sample denominators reported in forest plot.

(3) Numerators and denominators estimated from dispensing rates per 1000 registered patients and reported mean list sizes at baseline, respectively. Effective sample size calculated by imputing design effect at 12 months (6.9) reported by Altiner (2007).

Analysis I.3. Comparison I Shared decision making versus usual care (control), Outcome 3 Antibiotic prescriptions (index consultation) (adjusted odds ratio).

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: 3 Antibiotic prescriptions (index consultation) (adjusted odds ratio)

Study or subgroup	Shared decision making N	Usual care N	log [Odds Ratio] (SE)	Odds Ratio IV,Random,95% Cl	Weight	Odds Ratio IV,Random,95% Cl
Briel 2006 (I)	259	293	-0.1508 (0.3906)		26.7 %	0.86 [0.40, 1.85]
Francis 2009 (2)	256	272	-1.2379 (0.3716)	-	28.1 %	0.29 [0.14, 0.60]
Altiner 2007 (3)	1021	1143	-0.9676 (0.1936)	•	45.2 %	0.38 [0.26, 0.56]
Total (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z = Test for subgroup differen	= 3.02 (P = 0.002	5)	l ² =57%	•	100.0 %	0.44 [0.26, 0.75]
				0.002 0.1 I IO 500 decision making Usual care)	

(1) Logistic regression with random effect for each cluster and patient covariates (age, sex, education, days with restriction at baseline).

(2) Odds ratio from multilevel modelling.

(3) Adjusted for patient's disease severity, average practice severity (severity of the disease rated by the GP), patients having fever (compared with no fever), and frequency of fever in practice.

Analysis I.4. Comparison I Shared decision making versus usual care (control), Outcome 4 Antibiotic prescriptions (index consultation) (adjusted risk ratio).

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: 4 Antibiotic prescriptions (index consultation) (adjusted risk ratio)

Study or subgroup	Shared decision making N	Usual care N	log [Risk Ratio] (SE)	IV,F	Risk Ratio Random,95% Cl	Weight	Risk Ratio IV,Random,95% Cl
L gar 2012(1)	181	178	-0.6931 (0.2606)			24.8 %	0.50 [0.30, 0.83]
Little 2013 (2)	2332	1932	-0.3711 (0.1251)		+	75.2 %	0.69 [0.54, 0.88]
Total (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = Test for subgroup differen	= 3.24 (P = 0.001	2)	l ² = 19%		•	100.0 %	0.64 [0.49, 0.84]
			Shared	0.01 0.1 d decision makir	I IO Ig Usual care	100 e	

(1) Adjusted for cluster design, baseline values and patient age group (for analyses at teaching-unit and physician levels).

(2) Adjusted for baseline prescribing and clustering by physician and practice, age, smoking, sex, major cardiovascular or respiratory comorbidity, baseline symptoms, crepitations, wheeze, pulse higher than 100 beats per minute, temperature higher than 37.8 degrees celcius, respiratory rate, blood pressure, physician's rating of severity, and duration of cough.

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Analysis 1.5. Comparison I Shared decision making versus usual care (control), Outcome 5 Antibiotic prescriptions (index consultation or population rate per unit of time) (adjusted risk difference).

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: 5 Antibiotic prescriptions (index consultation or population rate per unit of time) (adjusted risk difference)

Study or subgroup	Shared decision making	Usual care	Mean Difference (SE)	Diffe	Mean erence	Weight	Mean Difference
	N	Ν		IV,Rando	om,95% Cl		IV,Random,95% CI
Butler 2012 (1)	239802	239700	-26.1 (12.3363)			11.1 %	-26.10 [-50.28, -1.92]
Cals 2009 (2)	201	230	-26.1 (5.6518)	-		33.0 %	-26.10 [-37.18, -15.02]
L gar 2011(3)	81	70	-16 (9.0754)		-	18.1 %	-16.00 [-33.79, 1.79]
Welschen 2004 (4)	905	818	-10.7 (4.898)	-		37.9 %	-10.70 [-20.30, -1.10]
Total (95% CI)	240989	240818		•		100.0 %	-18.44 [-27.24, -9.65]
Heterogeneity: Tau ² = 29	$17; Chi^2 = 4.7;$	5, df = 3 (P = 0). 9); ² =37%				
Test for overall effect: Z =	= 4.11 (P = 0.00	0040)					
Test for subgroup differen	ices: Not applic	able					
				-100 -50 0) 50 100		
			Shared	decision making	Usual care		

(1) Analysis of covariance with the previous year's prescribing as a covariate.

(2) Crude 95%Cl calculated and inflated for clustering by using standard deviation inflated by variance inflation factor. P value calculated from second order penalised quasi-likelihood multilevel logistic regression model for variance at general practitioner and practice level (random intercept at practice and general practitioner level). Models included both interventions and interaction term of intervention.

(3) All P values adjusted for baseline values and the study's cluster design.

(4) Intervention effect in multilevel analysis.

Analysis 1.6. Comparison I Shared decision making versus usual care (control), Outcome 6 Number or rate of re-consultations (risk ratio).

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: 6 Number or rate of re-consultations (risk ratio)

Study or subgroup	Shared decision making	Usual care	log [Risk Ratio]	Risk Ratio	Weight	Risk Ratio
	Ν	Ν	(SE)	IV,Random,95% CI		IV,Random,95% CI
Francis 2009 (1)	256	272	-0.2252 (0.3535)		5.8 %	0.80 [0.40, 1.60]
L gar 2012(2)	181	178	0.2624 (0.3158)		7.3 %	1.30 [0.70, 2.41]
Cals 2009 (3)	201	230	-0.288 (0.1492)	-	32.7 %	0.75 [0.56, 1.00]
Briel 2006 (4)	253	290	-0.0901 (0.116)	•	54.1 %	0.91 [0.73, 1.15]
Total (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z =	= 1.61 (P = 0.11)		2 =0.0%	•	100.0 %	0.87 [0.74, 1.03]
Test for subgroup differer	nces: Not applicab	ble				
			Share	0.01 0.1 I I0 I0 d decision making Usual care	0	

(1) Reported intra-class correlation co-efficient (ICC) = 0.06. Design effect and effective sample size calculated.

(2) Adjusted for cluster design and baseline values.

(3) Reported intra-class correlation co-efficient (ICC) = 0.01. Design effect and effective sample size calculated.

(4) Reported intra-class correlation co-efficient (ICC) = 0.04. Design effect and effective sample size calculated.

Analysis I.7. Comparison I Shared decision making versus usual care (control), Outcome 7 Patient satisfaction with the consultation.

Review: Interventions to facilitate shared decision making to address antibiotic use for acute respiratory infections in primary care

Comparison: I Shared decision making versus usual care (control)

Outcome: 7 Patient satisfaction with the consultation

-

Study or subgroup	Shared decision making N	Usual care N	log [Odds Ratio] (SE)	IV	Odds Ratio Random,95% Cl	Weight	Odds Ratio IV,Random,95% Cl
Francis 2009 (1)	246	263	-0.4463 (0.338)		-	34.3 %	0.64 [0.33, 1.24]
Briel 2006 (2)	253	290	0 (0.2277)		-	65.7 %	1.00 [0.64, 1.56]
Total (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z : Test for subgroup differen	= 0.72 (P = 0.47)		l ² = 17%		•	100.0 %	0.86 [0.57, 1.30]
				0.005 0.	I I I0	200	
			Shan	ed decision mal	ting Usual care		

(1) Odds Ration from multilevel modelling

(2) Proportion of patients with a maximum score of 70 (out of 70) used due to highly skewed scores. Logistic regression with random effect for each cluster and patient covariates (age, sex, education, days with restrictions at baseline.

ADDITIONAL TABLES

Table 1. TIDieR intervention summary (Hoffmann 2014)

Au- thor year	Brief name	Recip- ient	Why	What (mate- rials)	What (pro- ce- dures)	Who pro- vided	How	Where	When and how much	Tailor- ing	Mod- ifica- tion of inter- ven- tion through out trial	Strate- gies to im- prove or main- tain inter- ven- tion fi- delity	Ex- tent of inter- ven- tion fi- delity
Al- tiner 2007	Com- plex GP peer- led ed-	GPs and pa- tients	Fo- cused on com- muni-	Peers used a semi- struc- tured	GP peer- led out- reach	5 prac- tising GPs and teach-	Face- to-face out- reach visits	GP clinics during normal work-	1 out- reach visit per- formed	Not de- scribed	Not de- scribed	Not de- scribed	51/52 GPs re- ceived inter-

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			hoped that doc- tors and pa- tients would openly talk about the issue and thus reduce unnec- essary antibi- otic pre- scrip- tions	and antibi- otics	con- sulta- tion Patient mate- rials were aimed at em- power- ing pa- tients to raise and clarify issues within the con- sulta- tion								
Briel 2006	Brief train- ing pro- gramme in pa- tient- cen- tred com- muni- cation	GPs	Fo- cused on teach- ing GPs how to under- stand and modify pa- tients' con- cepts and beliefs about the use of antibi- otics for ARIS.	by	and to tailor infor- mation given to pa- tients. Physi-	Not speci- fied	Semi- nar in small groups (num- ber not spec- ified) and per- sonal feed- back by tele- phone prior to the start of the trial. Evi- dence- based	Not speci- fied	Atten- dance at 1 x 6-hour sem- inar and 1 x 2-hour tele- phone call to give per- sonal feed- back prior to the trial start	Not de- scribed	Not de- scribed	Not de- scribed	Not de- scribed

Table 1.	TIDieR intervention summary (Hoffmann 2014)	(Continued)
Table 1.	Tibler mervenelon summary (Tommann 2011)	(Communa)

			GPs were intro- duced to a model (Prochas 1992) for iden- tifying pa- tients' atti- tude and readi- ness for be- haviour change	as a book- let [URL pro- vided is no longer active]	used a model were intro- duced to a model (Prochas 1992) to iden- tify pa- tients' atti- tude and readi- ness for be- haviour change		guide- lines were dis- tributed as a book- let						
Butler 2012	Multi- faceted flexible blended learn- ing ap- proach for clini- cians	GPs and nurse practi- tioners	Blended learn- ing expe- rience to de- velop clini- cians' sense of the impor- tance about change and their confi- dence in their ability to achieve change based	of re- search evi- dence and guide- lines, web- based mod- ules using video- rich ma- terial pre- senting novel com- muni- cation	Inter- ven- tion consist of 7 com- po- nents: experi- ential learn- ing, up- dated sum- maries of re- search evi- dence and guide- lines; web- based	A facil- itator con- ducted face- to-face semi- nar	Inter- ven- tion con- sisted of 7 parts (5 online mod- ules, 1 face- to-face sem- inar and 1 facili- tator- led prac- tice- based semi- nar)	The face- to-face and facili- tator- led sem- inars were pre- sented at the general prac- tice	mod- ule (6	pa-	Not de- scribed	Not de- scribed	138/ 139 com- pleted all online train- ing and up- loaded de- scrip- tions of con- sulta- tions for the port- folio tasks; 129/ 139 at-

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			prin- ciples of mo- tiva- tional inter- view- ing		to share experi- ences and views								
Cals 2009	En- hanced com- muni- cation skills train- ing	GPs	Fo- cused on infor- mation ex- change based on the elicit- pro- vide- elicit frame- work from coun- selling in be- haviour change - explor- ing pa- tients' fears and ex- pecta- tions, pa- tients' opin- ion on antibi- otics and outlin- ing the	Pre and post- work- shop tran- scripts of sim- ulated pa- tients	Brief con- text- learn- ing based work- shop in small groups (5 to 8 GPs) , pre- ceded and fol- lowed by prac- tice- based con- sulta- tions with simu- lated pa- tients. GPs re- flected on own tran- scripts of con-	Expe- ri- enced mod- erator to lead semi- nars	Brief work- shop (5 to 8 GPs) , pre- ceded and fol- lowed by prac- tice- based con- sulta- tion with simu- lated pa- tients	Gen- eral prac- tice	1 x 2- hour mod- erator- led small groups work- shop, pre- ceded and fol- lowed by prac- tice- based con- sulta- tion with simu- lated pa- tients	Not de- scribed	Not de- scribed	Not de- scribed	66% of pa- tients re- cruited by GPs allo- cated to train- ing in en- hanced com- muni- cation skills re- called their GP's use at least 3 of 4 spe- cific com- muni- cation skills com- muni- cation skills com- muni- cation skills

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			natural dura- tion of cough in lower respi- ratory tract infec- tions		sulta- tions with simu- lated pa- tients, which were also peer- re- viewed by col- leagues								19% in the no train- ing group
Francis 2009	Inter- active book- let for parents and clini- cian train- ing in its use	GPs and pa- tients	Fo- cused on spe- cific com- muni- cation skills, such as explor- ing par- ent's main con- cerns, asking about their ex- pecta- tions, and dis- cussing prog- nosis, treat- ment op- tions and	(now at www.wh); online train- ing in use of the book- let in- cluded videos to demon- strate use	con- sulta- tion and as a take- home re- source (no further details pro- vided) Online train- ing on the use of the book- let was	N/A (online train- ing)	Parents used the book- let face- to-face in the con- sulta- tion with GPs and took it home; GP train- ing in use of book- let was online	Gen- eral prac- tice; par- ents' homes	1 x 40- minute online train- ing mod- ule	Not de- scribed	Not de- scribed	Online clin- ician train- ing moni- tored through study web- site: whether a GP has logged on to the site, how much time spent on it and which pages were viewed	sured so that asses-

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Table 1. TIDi	eR intervention summ	ary (Hoffmann 2014)	(Continued)
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			reasons that should prompt re-con- sulta- tion	tures and	de- scrib- ing the con- tent and aims of the book- let, and en- cour- aging use within the con- sulta- tion to facil- itate use of spe- cific com- muni- cation skills								
Légaré 2012	Shared deci- sion mak- ing train- ing pro- gram (DE- CI- SION+2	physi- cians (in- clud- ing teach- ers and resi- dents)	A shared deci- sion mak- ing train- ing pro- gram that aimed to help physi- cians com- mu- nicate to pa-	Online tuto- rial and work- shop in- cluded videos, exer- cises and deci- sion aids to help physi- cians	Online self tu- torial com- prising 5 mod- ules 2- hour online tuto- rial fol- lowed by a facili- tator- led on- site inter-	Trained facili- tators	On- line tu- torial and face- to-face work- shop	Family prac- tice teach- ing units	1 x 2- hour online tuto- rial, fol- lowed by 1 x 2-hour on-site inter- active work- shop. Partic- ipants had 1	Not de- scribed	Not de- scribed	Not de- scribed	Of the 162 physi- cians, 103 com- pleted both the online tuto- rial and work- shop; 16

Table 1. TID	ieR intervention su	ummary (Hoffmann 2014)	(Continued)
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			tients the proba- bility of a bac- terial ARI and the bene- fits and harms asso- ciated with the use of antibi- otics	com- muni- cate to their pa- tients the proba- bility of bac- terial ARIs and ben- efits/ harms of an- tibiotic use. Deci- sion aids were avail- able in the con- sulta- tion rooms in all family prac- tice teach- ing units	con- cepts ac- quired during online train-				month to com- plete the pro- gramme				com- pleted only the work- shop; 15 only the tu- torial; and 28 com- pleted none of the train- ing com- po- nents
Légaré 2011	Mul- tiple- com- po- nent, con- tin- uing profes- sional devel-	medicine		Work- shops in- cluded videos (sim- ulated con- sulta- tions of	Inter- active work- shops and related mate- rial, re- minders of ex- pected	Trained facili- tators	Face- to-face work- shop	Family medicino groups		Not de- scribed	4 pilot work- shops held rather than 3 as the second work- shop	Not de- scribed	Not de- scribed

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op-	proba-	usual	be-	addi-	was re-
ment	bility	care	haviours	tion	designec
pro-	of bac-	and	and	to re-	and re-
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sion	fits and		agree-	haviours	on its
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 Table 1. TIDieR intervention summary (Hoffmann 2014)
 (Continued)

Little 2013	Inter- net- based train- ing in en- hanced com- muni- cation skills	GPs	Ratio- nale was that Inter- net- based train- ing can be more widely dis- semi- nated than face- to-face train- ing fo- cused on elicit- ing pa- tients' expec- tations and con- cerns, natural disease course, treat- ments, agree- ment agre- ment plan, sum- ming	Inter- active book- let for use by GPs within con- sulta- tions Train- ing sup- ported by video demon- stra- tions of con- sulta- tion tech- niques	ap-	N/A (online mod- ules) other than lead GP at each prac- tice to organ- ise a meet- ing (not spe- cific to just this arm of the in- terven- tion though)	Online mod- ules (and GP-led struc- tured prac- tice- based meet- ing)	Gen- eral prac- tice	Inter- net mod- ules com- pleted alone or in a group	Not de- scribed	Not de- scribed	Not de- scribed	94/ 108 prac- tices (87%) com- pleted the com- muni- cation train- ing. Mean (SD) time spent on the web- site was 37 (29) min- utes
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			up and guid- ance on when to re- con- sult										
Welsche 2004	Group e edu- cation meet- ing with con- sensus proce- dure and com- muni- cation skills train- ing	GPs/ phar- ma- cists and their assis- tants, and pa- tients	GPs dis- cussed evi- dence for an- tibiotic ben- efit/ risk, and learned com- muni- cation tech- niques to explore pa- tients' expec- tations and con- cerns, inform about natural course of symp- toms, self- medi- cation and alarm	Group con- sensus guide- lines and patient wait- ing room mate- rials (poster/ leaflets)	Group edu- cation meet- ing with con- sensus proce- dure, with a sum- mary, and guide- lines mailed 1 month later to rein- force con- sensus reached; feed- back on pre- scrib- ing be- haviour (post- and pre-in- terven- tion insur- ance	Jointly led by GP and phar- macist	Group edu- cation meet- ing for GPs with con- sensus proce- dure and com- muni- cation skills train- ing, Group edu- cation for GPs' and phar- ma- cists' assis- tants, moni- toring and feed- back on pre- scrib- ing be-	Not de- scribed	1 x group edu- cation meet- ing with con- sensus proce- dure; 1 x 2- hour group edu- cation session for GP and phar- ma- cists' assis- tants; moni- toring and feed- back of pre- scrib- ing be- haviour at 6 months post-	Not de- scribed	Not de- scribed	Not de- scribed	Not de- scribed

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Table 1. TIDieR intervention summary (Hoffmann 2014) (Continued)

toms.	claims	haviour,	inter-	
Patient	data)	and	ven-	
edu-	and	patient	tion	
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vided	level	mate-		
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self-	pre-			
lim-	scrib-			
iting	ing be-			
nature	haviours			
or ARIs,	aligned with			
self-				
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	reached;			
and	group			
alarm	edu-			
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	terial			
	for pa-			
	tients			

ARI: acute respiratory infection

Author	Outcome	Measurement time point	Intervention (n)	Control	Effect estimate	P value	Notes
					Adjusted odds ratio (95% CI)		
Francis (2009)	Antibiotics prescribed at the index con- sultation	14 days	(30 practices) Patients = 50/ 256 (19.5%)	(31 practices) Pa- tients = 111/ 272 (40.8%)	0.29 (0.14 to $0.60)^a$	NR	ICC = 0.24
Altiner (2007)	antibiotic pre- scriptions (per	6 weeks	GPs = 42 Patients = 1021	GPs = 44 Patients = 1143	0.38 (0.26 to 0.56) ^b	< 0.001	ICC=0.20
	acute cough and per GP)	12 months	GPs = 28 Patients = 787	GPs = 33 Patients = 920	$0.55 (0.38 to 0.80)^b$	0.002	
Briel (2006)	Uptake of antibiotic pre- scriptions as reported by pharmacists < 2 weeks after the consulta- tion	14 days	GPs = 15 Patients = 259	GPs = 15 Patients = 293	0.86 (0.40 to 1.93) ^c	NR	ICC = 0.04 Design effect = 1.6
					Ad- justed risk ra- tio (95% CI)		
Little (2013)	Antibiotic prescription	index consul- tation	Practices = 61 Patients = 2332	Practices = 61 Patients = 1932	0.69 (0.54 to $0.87)^d$	< 0.0001	-
Légaré (2012)	% pa- tients who de- cided to use antibiotics im- mediately af- ter the consul- tation	Index consul- tation	Practice units = 6 GPs = 77 Patients = 181	= 6 GPs = 72	0.50 (0.30 to 0.70) ^e	-	-

Table 2. Antibiotic prescriptions per index consultation or population rate over time

Adjusted risk difference (95% CI) Légaré % pa- Index consul- Medicine Medicine -16 (-31 to 1) 0.08 groups = 2(2011)tients who detation groups cided to use GPs = 18GPs = 14antibiotics im-Patients = 81 Patients = 70 mediately after the consultation Butler (2012) Total no. dis- 12-month pe- Practices Practices -4.2 (-0.6 to - 0.02 = 34 Patients = = 34 Patients = 7.7) pensed riod oral antibiotic 7053 7050 items per 1000 registered patients for the year after the intervention Cals (2009) Antibiotic Index consul- n/N = 55/201 n/N = 123/-26.1 $(\% < 0.01^{h})$ ICC = 0.12prescribing at tation % crude (95% 230 crude) the index con- $CI)^G$ % crude (95% sultation 27.4 (25.6 to CI)g 36.6) 53.5 (43.8 to 63.2) Cals (2013) Proportion -10.4^{i} 0.02^{i} Mean n = 178n = 201% (95% CI) of episodes of 3.67 years fol-% (95% CI) 26.3 (20.6 to 39.1 (33.1 to respilow-up ratory tract in-32.0) 45.1) fections during follow-up for which a GP was seen and that antibiotics were prescribed for Welschen % practice en-Index consul-Review groups Review groups -10.7 (-20.3 to -Practice = (2006)countation = 6 = 6 $-1.0)^{j}$ 0.17 ters for acute Review group = 0.09 symptoms of the respiratory tract for which antibiotics were pre-

Table 2. Antibiotic prescriptions per index consultation or population rate over time (Continued)

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Table 2. Antibiotic prescriptions per index consultation or population rate over time (Continued)

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^aTwo level (practice and patient) random intercept logistic regression models.

^{*b*}After backward elimination, four explanatory variables remained in the model: patients' disease severity, measured on a four-point scale (odds ratio 4.8, 95% CI 3.9 to 5.9 per step on scale, P value < 0.001), and average practice severity (severity of the disease rated by the GP) (odds ratio 0.14, 95% CI 0.06 to 0.33, P value < 0.001 per category step on the scale), patients having fever (odds ratio 1.80, 95% CI 1.35 to 2.39, P value < 0.001 compared with no fever) and frequency of fever in practice, as determined by the log odds (odds ratio 1.31, 95% CI 1.08 to 1.59, P value = 0.007 per category step on the scale).

^cLogistic regression with random effects for each cluster and patient covariates (age, sex, education, days with restrictions at baseline). ^dThe adjusted model adjusted for baseline prescribing and clustering by physician and practice, and additionally controlled for age, smoking, sex, major cardiovascular or respiratory comorbidity, baseline symptoms, crepitations, wheeze, pulse higher than 100 beats per minute, temperature higher than 37.8°C, respiratory rate, blood pressure, physician's rating of severity and duration of cough. ^eAdjusted for cluster design, baseline values and patient age group (for analyses at teaching unit and physician levels).

^f P value adjusted for baseline values and the study's cluster design.

^gCalculated and inflated for clustering by using standard deviation inflated by variance inflation factor.

^hCalculated from second order penalised quasi-likelihood multilevel logistic regression model adjusted for variance at general practitioner and practice level (random intercept at practice and general practitioner level). Models included both interventions and interaction term of interventions.

^{*i*}P values from multilevel linear regression model to account and correct for variation at the level of family physician, and to adjust for both interventions, RTI-episodes treated with antibiotics during baseline period, chronic obstructive pulmonary disease comorbidity. ^{*j*}Intervention effect in multi-level analysis

CI: confidence interval

GP: general practitioner

NR: not reported

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Briel (2006)	Re- consultations	Within 14 days	. ,	n/N (%) 143/290 (49. 3)		NR	-
Butler (2013)	Re-consulta- tions after in- dex consulta- tion) ^b	days	2.66 (1.88 to 4.25) 5.10 (4.70 to 7.92)	3.35 (2.16 to	ence (95% CI) c -0.65 (-1.69 to 0.55)	446^{d} P value = 0. 411^{d} P value = 0.	-

Table 3. Number or rate of re-consultations

Table 3. Number or rate of re-consultations (Continued)

Cals (2009)	Re- consultations	Within 28 days		% crude (95% CI) ^e	Absolute dif- ference 9.1 (% crude)	0.14 ^{<i>f</i>}	ICC = 0.01
Francis (2009)	Re- consultation ^g	Within 14 days	n/N (%) 33/256 (12.9)	n/N (%) 44/272 (16.2)	Adjusted odds ratio (95% CI) 0.75 (0.41 to 1.38)	NR	ICC = 0.06
Légaré (2012)	Re- consultation	Baseline (pre)	21.6 (12.1 to 29.7)	22.7 (10.3 to 27.3)	justed risk ra- tio (95% CI) ^h 1.3 (0.7 to 2.	NR	-
		Within 14 days (post)	13.4 (9.9 to 15.9)	15.2 (11.9 to 19.4)	3) Absolute dif- ference = 7.5		
Little (2013)	New or wors- ening symptoms ⁱ	-	n/N (%) 451/2242 (20%)	n/N (%) 309/1879 (16%)	Ad- justed risk ra- tio (95% CI) ^j 1.33 (0.99 to 1.74)	P value = 0. 055	-

^{*a*} Poisson regression with random effects for each cluster and patient covariates (age, sex, education, days with restrictions at baseline). ^{*b*} Collected from the electronic records of a subsample of 37 general practices (20 intervention/17 control). 47 patients (10.9%) reconsulted more than once within 28 days with pattern similar across groups.

^{*c*}Computed with bootstrapping methods.

^dFrom Mann-Whitney U test.

^eCalculated and inflated for clustering by using standard deviation inflated by variance inflation factor.

^f Calculated from second order penalised quasi-likelihood multilevel logistic regression model adjusted for variance at general practitioner and practice level (random intercept at practice and general practitioner level). Models included both interventions and interaction term of interventions.

^g Parental report that child attended a face-to-face consultation with a primary care clinician in their general practice, or with an out of hours provider, in the 2 weeks after registration.

^hAdjusted for cluster design and baseline values.

ⁱDefined as re-consultation for new or worsening symptoms within 4 weeks, new signs or hospital admission.

^{*j*}The adjusted model adjusted for baseline prescribing and clustering by physician and practice, and additionally controlled for age, smoking, sex, major cardiovascular or respiratory comorbidity, baseline symptoms, crepitations, wheeze, pulse higher than 100 beats per minute, temperature higher than 37.8°C, respiratory rate, blood pressure, physician's rating of severity and duration of cough. CI: confidence interval

ICC: intra-class correlation co-efficient

IQR: interquartile range

NR: not reported

Table 4. Incidence of hospital admissions

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Briel (2006)	Hospital admissions	< 28 days of study enrol- ment	n/N = 2/253	n/N = 1/290	NR	NR	-
Butler (2012)	Hospital admissions ^a	Baseline Follow-up	Mean 7.7 7.5	Mean 8.7 8.0	% reduc- tion (interven- tion relative to controls ^b (95% CI) -1.9 (-13.2 to 8.2)	P value = 0.72	-
Cals (2013)	Hospital admissions	Mean 3.67 year follow-up		n/N 5/201	NR	NR	-
Francis (2009)	Hospital admissions (or observed in a paediatric as- sessment unit)	< 14 days	n/N 3/256	n/N 4/272	NR	NR	-
Little (2013)	Hospital admissions ^c	< 4 weeks	n/N 6/1170	n/N 2/870	NR	-	-

^{*a*} Annual number of hospital episodes for possible respiratory tract infections and complications of common infections per 1000 registered patients. A single admission occurred if patient admitted to hospital for a possible RTI or complication. If patient admitted more than once, and gap between admissions was 30 days or more, this was considered a separate complication episode.

^bDifference between means in intervention group and control group as percentage of mean control group.

^cFactorial analysis data not reported

NR: not reported

RTI: respiratory tract infection

SAEs: serious adverse events

Table 5. Incidence of pneumonia

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Briel (2006)	Pneumonia	< 28 days	n/N = 0/253	1/290	NR	NR	-
Cals (2013)	Pneumonia	Mean 3.67 year follow-up	n/N = 0/178	n/N = 1/201	NR	NR	-

NR: not reported

Table 6. Patient satisfaction

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Briel (2006)	Patient satis- faction (Patient Satis- faction Ques- tionnaire) ^a	7 and 14 days	121/253 (47. 8)	142/290 (49. 0)	Adjusted OR (95% CI) ^b 1.00 (0.64 to 1.31)	NR	-
Cals (2009)	Patient satis- faction (% at least 'very sat- isfied' on Lik- ert scale) ^c	28 days	n/N = 144/ 201 % (crude 95% CI) ^d 78.7 (72.5 to 84.9)	n/N = 151/230 % (crude 95% CI) ^d 74.4 (68.2 to 80.6)	4.3	P value = 0.88 e	-
Francis (2009)	Parent satis- faction (Likert scale) ^f	14 days	n/ N (%) = 222/ 246 (90.2)	n/ N (%) = 246/ 263 (93.5)	Adjusted OR (95% CI) ^g 0.6 (0.3 to 1. 2)	NR	-
Welschen (2006)	Patient satis- faction (Likert scale) ^h	Index consul- tation	Patient satis- faction (%) Baseline (pre) = 4.3 (0.3) Follow- up (post) = 4. 3 (0.3) % change (SD) = 0 (0.4)	Patient satis- faction (%) Baseline (pre) = 4.2 (0.4) Follow- up (post) = 4. 2 (0.3) % change (SD): 0 (0.4)	difference of	NR	-

^{*a*}% patients with a maximum score of 70 reported, as satisfaction scores (scale 14 to 70; median 68/70) were highly skewed.

^bLogistic regression with random effects for each cluster and patient covariates (age, sex, education, days with restrictions at baseline). ^c% at least 'very satisfied'.

^dCalculated and inflated for clustering by using standard deviation inflated by variance inflation factor.

^eCalculated from models adjusted for variance at general practitioner and practice level.

^fTransformed into binary outcomes: 'very satisfied' and 'satisfied' versus 'neutral', 'dissatisfied' and 'very dissatisfied'.

^gOdds ratio (95% CI) from multilevel modelling.

 $^{h}1$ = very dissatisfied to 5 = very satisfied.

^{*i*}Intervention effect in multilevel analysis.

CI: confidence interval

OR: odds ratio

SD: standard deviation

Table 7. Decisional conflict

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Légaré (2012)	Deci- sional conflict (GPs) ^a	Immediately after consulta- tion	(0 to 9.0)	Baseline: 3.0 (0 to 5.9) Follow-up:1.1 (0 to 2.4)	3.4 (0.3 to 38.	NR	-
Légaré (2012)		Immediately after consulta- tion	(0 to 13.5)	Follow-up: 6.	0.8 (0.2 to 2.	NR	-
Légaré (2011)	Correlation of deci- sional conflict between GPs and patients ^a				ence at follow-	0.06	-

^{*a*} Proportion of participants who had a value of 2.5 or more on the Decision Conflict Scale (where 1 = low decisional conflict and 5 = very high decisional conflict).

^bPresented as correlation of family physicians' and patient's DCS scores (Pearson's r).

CI: confidence interval

GP: general practitioner

NR: not reported

RR: risk ratio

Table 8. Decisional regret

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Légaré (2012)	Decisional re- gret ^a	2 weeks after consultation	Baseline: 10.5 ± 15.4 Follow-up: 12.4 ± 19.1	Baseline: 10.8 ± 20.8 Follow-up: 7. 6 ± 13.7	mean	-	-
Légaré (2011)	Patients (%) with deci- sional regret	2 weeks after consultation	Baseline: 1 Follow-up: 7	Baseline: 1 Follow-up: 9	Differ- ence at follow- up (95% CI) -2 (-12 to 5)	0.91	-

a = Decisional Regret Scale used, where 0 = very low regret and 100 = very high regret CI: confidence interval

Table 9. Patient enablement

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Briel (2006)	Patient en- ablement (Pa- tient Enable- ment Instru- ment; scale 0 to 12)	7 and 14 days	Mean (SD): 8. 49 (1.98)	Mean (SD): 8. 15 (2.03)	Adjusted co- efficient (95% CI) ^{<i>a</i>} 0.35 (-0.05 to 0.75)	NR	-
Cals (2009)	Patient en- ablement (Pa- tient Enable- ment In- strument; max score is 12)	28 days		Median (IQR) score: $3 (4)^d$ Mean (SD) score: $3.06 (2.$ 54)	-	NR 0.70 ^b	-
Francis (2009)	Parent enable- ment (Modified Pa- tient Enable- ment Instru- ment, scale 1 to 10) ^c	14 days	n/N (%): 99/ 246 (40.2)	n/N (%): 94/ 262 (35.9)	Adjusted OR (95% CI) 1.20 (0.84 to 1.73)	NR	-

^{*a*}Linear regression with random effects for each cluster and patient covariates (age, sex, education, days with restrictions at baseline). ^{*b*}Calculated from models adjusted for variance at general practitioner and practice level.

^cPresented results are % with parent enablement score of 5 or more (binary outcome).

^dComparator is 'no skills training'.

CI: confidence interval

IQR: interquartile range

NR: not reported

OR: odds ratio

SD: standard deviation

Table 10. Quality of the decision made (GPs)

Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Légaré (2012)	Quality of decision made (GPs) (0 to 10 Likert scale)	After consul- tation	1.5	Baseline: 8.7 ± 1.5 Follow-up: 8. 5 ± 1.5	mean	NR	-

Table 10.	Quality of the decision made (GPs)	(Continued)
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Légaré	Ouality of	After consul-	Decoline, Q.Q.	Decoline 92	Differ	0.29	
0	C /		Dasenne: 0.0 ±			0.29	-
(2011)	decision made	tation	1.1	1.4	ence at follow-		
	(GPs) (0 to 10		Follow-up: 8.	Follow-up: 8.	up (95% CI)		
	Likert scale)		7 ± 1.2	5 ± 1.3	0.2 (-0.34 to		
					0.89)		

CI: confidence interval GP: general practitioner

NR: not reported

Table 11.	Quality of t	he decision	made (patients)
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Author	Outcome	Measurement time point	Intervention	Control	Effect estimate	P value	Notes
Légaré (2012)	Quality of de- ci- sion made (pa- tients) (0 to 10 Likert scale) ^a	After consul- tation	1.1	1.4 Follow-up: 8.	mean difference	NR	-
Légaré (2011)	Quality of the decision made (patients) (0 to 10 Likert scale) ^{<i>a</i>}	After consul- tation	2.1 Follow-up: 8.	Baseline: 8.4 ± 1.9 Follow-up: 8. 6 ± 1.9	ence at follow- up (95% CI)	0.57	-

^{*a*}Likert scale where 0 = very low quality and 10 = very high quality.

CI: confidence interval

NR: not reported

APPENDICES

Appendix I. MEDLINE (Ovid) search strategy

```
1 exp Respiratory Tract Infections/ (297579)
2 (respiratory adj2 (infection* or inflam*)).tw. (31350)
3 pharyngitis.tw. (4164)
4 sinusit*.tw. (11403)
5 (acute adj2 rhinit*).tw. (174)
6 (rhinosinusit* or nasosinusit*).tw. (4197)
7 common cold*.tw. (2806)
8 coryza.tw. (379)
9 (throat* adj2 (sore* or inflam* or infect*)).tw. (3897)
10 laryngit*.tw. (1305)
11 tonsillit*.tw. (4080)
12 bronchit*.tw. (18478)
13 bronchiolit*.tw. (8053)
14 pneumon*.tw. (133425)
15 (bronchopneumon* or pleuropneumon*).tw. (5382)
16 Cough/ (12409)
17 cough*.tw. (34227)
18 exp Otitis Media/ (21649)
19 otitis media.tw. (16032)
20 (aom or ome).tw. (6083)
21 Croup/ (970)
22 (croup or pseudocroup or laryngotracheobronchit* or laryngotracheit*).tw. (1971)
23 or/1-22 (451019)
24 exp Anti-Bacterial Agents/ (537825)
25 antibiotic*.tw,nm. (242634)
26 or/24-25 (640170)
27 23 and 26 (79549)
28 exp Decision Making/ (122846)
29 exp decision support techniques/ (62827)
30 exp Decision Theory/ (9884)
31 (decision* or decid* or option* or choice* or choose* or deliberat*).tw. (618268)
32 exp Informed Consent/ (35917)
33 (informed adj3 (consent* or agree* or assent*)).tw. (23002)
34 Health Knowledge, Attitudes, Practice/ (74387)
35 "Attitude of Health Personnel"/ (92103)
36 professional-patient relations/ or physician-patient relations/ (82522)
37 exp Consumer Participation/ (32440)
38 ((patient* or consumer* or carer* or parent* or child* or individual* or person* or interpersonal*) adj5 (participat* or involv* or
collabor* or cooperat* or co-operat* or engag* or consult* or feedback* or interaction*)).tw. (184609)
39 (values* or prefer*).tw. (981018)
40 exp Communication/ (369188)
41 (communicat* or negotiat* or facilitat* or discuss*).tw. (1366627)
42 health education/ or exp consumer health information/ or patient education as topic/ (125443)
43 ((patient* or consumer* or parent*) adj3 (educat* or informat*)).tw. (58615)
44 (shar* adj2 information*).tw. (3292)
45 sdm.tw. (869)
46 ((patient* or client* or subject or person or consumer* or family or families or carer* or care giver*) and (professional* or physician*
or clinician* or practitioner*)).tw. (327702)
47 Risk Assessment/ (180413)
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48 ((check or clarify) adj3 understanding).tw. (222)
49 (patient adj2 (understanding or expect*)).tw. (3479)
50 problem defin*.tw. (230)
51 (ask adj2 question*).tw. (1819)
52 (assess* adj2 risk*).tw. (50234)
53 self-manag*.tw. (8193)
54 equipoise.tw. (596)
55 checklist*.tw. (18085)
56 (goal adj2 set*).tw. (2180)
57 consensus.tw. (98026)
58 concordance.tw. (26142)
59 agreement*.tw. (155845)
60 (action* adj2 plan*).tw. (5452)
61 or/28-60 (3975067)
62 27 and 61 (14717)

Appendix 2. EMBASE (Elsevier) search strategy

#53 #23 AND #26 AND #52 28861 #52 #27 OR #28 OR #29 OR #30 OR #31 OR #32 OR #33 OR #34 OR #35 OR #36 OR #37 OR #38 OR #39 OR #40 OR #41 OR #42 OR #43 OR #44 OR #45 OR #46 OR #47 OR #48 OR #49 OR #50 OR #51 3678076 #51 'self-manage':ab,ti OR equipoise:ab,ti OR checklist:ab,ti OR consensus:ab,ti OR concordance:ab,ti OR agreement*:ab,ti OR (action* NEAR/2 plan*):ab,ti OR (goal NEAR/2 set*):ab,ti AND [embase]/lim298331 #50 (assess* NEAR/2 risk*):ab,ti AND [embase]/lim53641 #49 (ask NEAR/2 question*):ab,ti AND [embase]/lim1736 #48 (problem NEAR/1 defin*):ab,ti AND [embase]/lim327 #47 (patient NEAR/2 (understanding OR expect*)):ab,ti AND [embase]/lim4348 #46 ((check OR clarify) NEAR/3 understanding):ab,ti AND [embase]/lim203 #45 'risk assessment'/de AND [embase]/lim276013 #44 patient*:ab,ti OR client*:ab,ti OR subject:ab,ti OR person:ab,ti OR consumer*:ab,ti OR family:ab,ti OR families:ab,ti OR carer*: ab,ti OR 'care giver':ab,ti OR 'care givers':ab,ti AND (professional*:ab,ti OR physician*:ab,ti OR clinician*:ab,ti OR practitioner*: ab,ti) AND [embase]/lim349162 #43 sdm:ab,ti AND [embase]/lim776 #42 (shar* NEAR/2 information*):ab,ti AND [embase]/lim2631 #41 ((patient* OR consumer* OR parent*) NEAR/3 (educat* OR informat*)):ab,ti AND [embase]/lim59024 #40 'patient education'/de OR 'consumer health information'/de AND [embase]/lim41224 #39 communicat*:ab,ti OR negotiat*:ab,ti OR facilitat*:ab,ti OR discuss*:ab,ti AND [embase]/lim1335786 #38 'interpersonal communication'/de OR 'communication skill'/de OR 'nonverbal communication'/exp OR 'persuasive communication'/de OR 'verbal communication'/de OR 'conversation'/de AND [embase]/lim117435 #37 values*:ab,ti OR prefer*:ab,ti AND [embase]/lim995711 #36 ((patient* OR consumer* OR carer* OR parent* OR child* OR individual* OR person* OR interpersonal*) NEAR/5 (participat* OR involv* OR deliberat* OR collabor* OR cooperat* OR 'co-operate' OR 'co-operates' OR 'co-operation' OR engag* OR consult* OR feedback* OR interaction*)):ab,ti AND [embase]/lim197067 #35 'patient participation'/de AND [embase]/lim6904 #34 'doctor patient relation'/de AND [embase]/lim39102 #33 'attitude to health'/de AND [embase]/lim7634 #32 (treatment* NEAR/2 option*):ab,ti AND [embase]/lim65145 #31 (informed NEAR/3 (consent* OR agree*)):ab,ti AND [embase]/lim32077 #30 'informed consent'/de AND [embase]/lim39300 #29 decision*:ab,ti OR decid*:ab,ti OR option*:ab,ti OR choice*:ab,ti OR choice*:ab,ti OR deliberat*:ab,ti AND [embase]/lim639112 #28 'decision support system'/de AND [embase]/lim4763 #27 'decision making'/de OR 'patient decision making'/de OR 'medical decision making'/de OR 'clinical decision making'/de AND [embase]/lim158809

#26 #24 OR #25 892667 #25 antibiotic*:ab,ti AND [embase]/lim219681 #24 'antibiotic agent'/exp AND [embase]/lim842466 #23 #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21 OR #22 518323 #22 croup:ab,ti OR pseudocroup:ab,ti OR laryngotracheobronchit*:ab,ti OR laryngotracheit*:ab,ti AND [embase]/lim1492 #21 'otitis media':ab,ti OR aom:ab,ti OR ome:ab,ti AND [embase]/lim19731 #20 'otitis media'/exp AND [embase]/lim21150 #19 cough*:ab,ti AND [embase]/lim37668 #18 'coughing'/de AND [embase]/lim52337 #17 bronchopneumon*:ab,ti OR pleuropneumon*:ab,ti AND [embase]/lim3817 #16 pneumon*:ab,ti AND [embase]/lim131768 #15 bronchiolit*:ab,ti AND [embase]/lim8788 #14 bronchit*:ab,ti AND [embase]/lim15885 #13 tonsillit*:ab,ti AND [embase]/lim3497 #12 laryngit*:ab,ti AND [embase]/lim1237 #11 (throat* NEAR/2 (sore* OR inflam* OR infect*)):ab,ti AND [embase]/lim4582 #10 'sore throat'/de AND [embase]/lim8854 #9 'common cold':ab,ti OR 'common colds':ab,ti OR coryza:ab,ti AND [embase]/lim2828 #8 'common cold symptom'/de AND [embase]/lim269 #7 rhinosinusit*:ab,ti OR nasosinusit*:ab,ti AND [embase]/lim4585 #6 (acute NEAR/2 rhinit*):ab,ti AND [embase]/lim179 #5 sinusit*:ab,ti AND [embase]/lim11343 #4 pharyngit*:ab,ti AND [embase]/lim4248 #3 (respiratory NEAR/2 (infection* OR inflam*)):ab,ti AND [embase]/lim33268 #2 'respiratory tract inflammation'/exp AND [embase]/lim275986 #1 'respiratory tract infection'/exp AND [embase]/lim198937

Appendix 3. Web of Science (Thomson Reuters) search strategy

#7	#6 AND #5 DocType = All document types; Language = All languages;
#6	TOPIC: (random* or placebo* or ((singl* or doubl*) NEAR/1 blind*) or allocat* or crossover* or "cross over") OR TITLE: (trial) DocType=All document types; Language=All languages;
#5	#4 AND #3 DocType=All document types; Language=All languages;
#4	TOPIC: (sdm or decision* or decid* or choice* or prefer* or option*) OR TOPIC: ((informed NEAR/3 (consent* or agree*))) OR TOPIC: ((patient* or consumer* or parent* or personal* or individual* or interpersonal*) NEAR/2 (participat* or involv*)) DocType=All document types; Language=All languages;
#3	#2 AND #1 DocType=All document types; Language=All languages;
#2	TOPIC: (antibiotic* or antibacterial* or anti-bacterial*) DocType=All document types; Language=All languages;

(Continued)

#1 TOPIC: (pharyngit* or sinusit* or "acute rhinitis" or rhinosinusit* or nasosinusit* or "common cold*" or coryza or laryngit* or tonsillit* or bronchit* or bronchiolit* or pneumon* or bronchopneumon* or pleuropneumon* or cough* or "otitis media" or aom or ome or croup or pseudocroup or laryngotracheit* or laryngotracheobronchit*) OR TOPIC: ((respiratory NEAR/2 (infect* or inflam*))) or (throat* NEAR/2 (sore* or inflam* or infect*))) DocType=All document types; Language=All languages;

WHAT'S NEW

Date	Event	Description
8 February 2017	Amended	Minor formatting change to Table 1 with rows and columns swapped, to improve ease of reading

CONTRIBUTIONS OF AUTHORS

Tammy Hoffmann (TH) conceived the original idea for the review.

Peter Coxeter (PC) was responsible for drafting the protocol.

TH and Chris Del Mar (CDM) contributed content and methodological expertise and provided advice and guidance on the development of the draft protocol and final editing.

Elaine Beller (EB) provided statistical advice and guidance.

Leanne McGregor conducted independent screening (titles/abstract and full text) and data extraction.

DECLARATIONS OF INTEREST

Peter Coxeter: none declared

Chris B Del Mar: none declared

Leanne McGregor: none declared

Elaine M Beller: none declared

Tammy Hoffmann: none declared

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• No sources of support supplied

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INDEX TERMS

Medical Subject Headings (MeSH)

*Decision Making; Acute Disease; Anti-Bacterial Agents [*therapeutic use]; Patient Participation; Primary Health Care; Randomized Controlled Trials as Topic; Respiratory Tract Infections [*drug therapy]

MeSH check words

Humans