Accepted Manuscript

Treatment of Interstitial Lung Disease associated cough: CHEST guideline and expert panel report

Surinder S. Birring, MD, Joanne E. Kavanagh, MBChB, Richard S. Irwin, MD, Master FCCP, Karina Keogh, MD, Kaiser G. Lim, MD, Jay H. Ryu, MD

PII: S0012-3692(18)31075-4

DOI: 10.1016/j.chest.2018.06.038

Reference: CHEST 1826

To appear in: CHEST

Received Date: 23 May 2018

Accepted Date: 22 June 2018

Please cite this article as: Birring SS, Kavanagh JE, Irwin RS, Keogh K, Lim KG, Ryu JH, Treatment of Interstitial Lung Disease associated cough: CHEST guideline and expert panel report, *CHEST* (2018), doi: 10.1016/j.chest.2018.06.038.

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.







	A CORDER MANUCODIT
	ACCEPTED MANUSCRIPT
1	Treatment of Interstitial Lung Disease associated cough:
2	CHEST guideline and expert panel report
3	
4	Authors
5	Surinder S Birring, MD ¹
6	Joanne E Kavanagh, MBChB ²
7	Richard S. Irwin, MD, Master FCCP ³
8	Karina Keogh, MD ⁴
9	Kaiser G Lim, MD ⁴
10	Jay H Ryu, MD ⁴
11	
12 13 14 15 16	¹ Division of Asthma, Allergy & Lung Biology, School of Transplantation, Immunology, Infection & Inflammation Sciences, Faculty of Life Sciences & Medicine, King's College London, King's Health Partners, London, UK ² Chest Department, St Thomas' Hospital, London, UK
17 18 19	³ Division of Pulmonary, Allergy, and Critical Care Medicine, UMass Memorial Medical Center, Worcester, MA, USA
1) 20 21 22	(KK, KGL, JHR) Division of Pulmonary and Critical Care Medicine, Mayo Clinic, Rochester MN, USA
23 24	Correspondence to: Professor Surinder S Birring
25 26 27 28 29	Division of Asthma, Allergy & Lung Biology, School of Transplantation, Immunology, Infection & Inflammation Sciences, Faculty of Life Sciences & Medicine, King's College London, King's Health Partners, Denmark Hill, London, SE5 9RS, United Kingdom. Telephone: (+44) 203 299 4630. Email: surinder.birring@nhs.net
30 31 32 33	DISCLAIMER: American College of Chest Physician guidelines are intended for general information only, are not medical advice, and do not replace professional medical care and physician advice, which always should be sought for any medical condition. The complete disclaimer for this guideline can be accessed at http://www.chestnet.org/Guidelines-and-Resources/Guidelines-and-Consensus-Statements/CHEST-Guidelines.

- not medical advice, and do not replace professional medical care and physician advice, which always should be
- sought for any medical condition. The complete disclaimer for this guideline can be accessed at
- http://www.chestnet.org/Guidelines-and-Resources/Guidelines-and-Consensus-Statements/CHEST-Guidelines.



34 Abbreviations

ATS	American Thoracic Society
CDC	Centers for Disease Control and Prevention
CIC	Chronic Idiopathic Cough
CINAHL	Cumulative Index of Nursing and Allied Health Literature
CQLQ	Cough Quality of Life Questionnaire
CTD	Connective Tissue Disease
CXR	Chest Radiograph
EB	Eosinophilic Bronchitis
EMBASE	Excerpta Medica dataBASE
ERS	European Respiratory Society
GERD	Gastro-Esophageal Reflux Disease
GI	Gastro-Intestinal
ILD	Interstitial Lung Disease
IPF	Idiopathic Pulmonary Fibrosis
MCID	Minimal Clinically Important Difference
LCQ	Leicester Cough Questionnaire
PICO	Population, Intervention, Comparison, Outcome
PPI	Proton Pump Inhibitor
PSALTI	Physiotherapy and Speech And Language Therapy Intervention
RCT	Randomized Control Trial
SGRQ	St George's Respiratory Questionnaire
SPT	Speech Pathology Therapy
SSc	Systemic Sclerosis
TLCO	Transfer Factor of the Lung for Carbon Monoxide
UACS	Upper Airway Cough Syndrome
VAS	Visual Analogue Score
VC	Vital Capacity



37 ABSTRACT

BACKGROUND: Chronic cough in interstitial lung disease (ILD) causes significant
impairment in quality of life. Effective treatment approaches are needed for cough associated
with ILD.

41 **METHODS:** This systematic review asked: Is there evidence of clinically relevant treatment 42 effects for therapies for cough in ILD? Studies of adults aged > 18 years with a chronic cough 43 > 8 weeks duration were included and assessed for relevance and quality. Based on the 44 systematic review, guideline suggestions were developed and voted on by using CHEST 45 guideline methodology.

46 **RESULTS:** Eight Randomized Controlled Trials (RCT) and 2 case series (>10 patients) were 47 included that reported data on patients with Idiopathic Pulmonary fibrosis (IPF), Sarcoidosis 48 and Scleroderma-related ILD who received a variety of interventions. Study quality was high 49 in all 8 RCTs. Inhaled corticosteroids were not supported for cough associated with 50 Sarcoidosis. Cyclophosphamide and Mycophenolate were not supported for solely treating 51 cough associated with scleroderma associated ILD. A recommendation for thalidomide to 52 treat cough associated with IPF did not pass the panel vote. In view of the paucity of 53 antitussive treatment options for refractory cough in ILD, the guideline panel suggested that 54 the CHEST unexplained cough guideline be followed, by considering options such as the 55 neuromodulator gabapentin and speech pathology management. Opiates were also suggested 56 for patients with cough refractory to alternative therapies.

57 **CONCLUSIONS:** The evidence supporting the management of chronic cough in ILD is 58 limited. This guideline presents suggestions for managing and treating cough on the best 59 available evidence but future research is clearly needed.

KEY WORDS: chronic cough, interstitial lung disease, sarcoidosis, scleroderma, treatment,
 refractory, unexplained.



62 SUMMARY OF RECOMMENDATIONS AND SUGGESTIONS

63

64 1. For patients with ILD who present with a troublesome cough, we suggest that 65 patients be assessed for progression of their underlying ILD, or complications from immunosuppressive treatment (e.g. drug side effect, pulmonary infection) and also be 66 considered for further investigation / treatment trials for their cough according to 67 68 guidelines for acute, subacute and chronic cough. (Ungraded Consensus- Based Statement) 69 70 2. For patients with IPF, chronic cough and a negative workup for acid gastroesophageal reflux, we suggest that proton pump inhibitor therapy should not be 71 72 prescribed. (Ungraded Consensus- Based Statement) 73 74 3. For patients with pulmonary sarcoidosis, we suggest that inhaled corticosteroids 75 should not be routinely prescribed to treat the chronic cough. (Grade 2C). 76 77 4. For patients with ILD and refractory chronic cough, we suggest trials of therapies 78 recommended for patients with unexplained chronic cough according to the CHEST guidelines, with treatments such as gabapentin and multimodality speech pathology 79 80 therapy, or entering into clinical trials if available. (Ungraded Consensus- Based 81 Statement) 82 83 5. For patients with chronic cough due to ILD, when alternative treatments have failed 84 and the cough is adversely affecting their quality of life, we suggest that opiates be

recommended for symptom control in a palliative care setting with reassessment of the



- 86 benefits and risks at 1 week and then monthly before continuing. (Ungraded Consensus-
- 87 Based Statement)



89 INTRODUCTION

90 Interstitial lung disease (ILD) comprises a wide range of acute and chronic pulmonary 91 disorders that affect both the airways and lung parenchyma with variable amounts of 92 inflammation and fibrosis. Cough is a common symptom associated with ILD, and may be 93 the presenting symptom in some patients. Cough in idiopathic pulmonary fibrosis (IPF) has been the best studied of all ILDs and a chronic cough is present in up to 80% of patients with 94 IPF, although the proportion of patients with a troublesome cough is likely to be less than 95 this¹. Cough in IPF has been assessed with validated tools, such as 24-hour cough monitoring 96 and health-related Quality of Life questionnaires². The health-related quality of life 97 98 impairment in cough associated with IPF is significant and comparable with unexplained chronic cough 2,3 . 99

100

101 The presence of cough in IPF also has prognostic significance. A recent study of 242 patients with IPF found that cough predicted disease progression, independent of disease severity⁴. 102 103 Cough was more prevalent in patients with more advanced pulmonary fibrosis. An important 104 challenge when assessing cough in patients with ILD is to establish whether the cough is a 105 consequence of the underlying inflammation or fibrosis, or due to the presence of a co-106 morbidity. In IPF, while the mechanism of cough may be mechanical distortion associated with lung parenchymal fibrosis⁵, heightened cough reflex sensitivity, gastro-esophageal reflux 107 108 and airway inflammation have all been reported in patients with IPF and may therefore also be potentially important mechanisms⁶⁻⁸. 109

110

111 Cough is also a common symptom in patients with pulmonary sarcoidosis. A recent study by 112 Sinha et al reported that 50% of patients had a chronic cough and this was associated with 113 poor health-related quality of life⁹. Furthermore, there was a significant association between



the frequency of cough measured objectively with 24-hour cough monitoring and healthrelated quality of life. In the study by Sinha et al, cough reflex sensitivity assessed with capsaicin was the only independent predictor of the frequency of coughing. Lung function, disease activity or severity were not predictors of cough frequency. In scleroderma-related ILD, cough is also a prevalent symptom, and is associated with more severe interstitial lung disease^{10,11}.

120

121 The present systematic review addresses the problem of chronic cough associated with 122 interstitial lung diseases, including sarcoidosis and scleroderma, in the areas of management 123 and future directions.

124

125 **METHODS**

126 The methodology of the CHEST Guideline Oversight Committee was used to select the 127 Expert Cough Panel chair and the international panel of experts to perform the systematic 128 review, synthesis of the evidence, and development of the recommendations and 129 suggestions¹².

130

131 Systematic Review Question

The key clinical question for this systematic review, developed by the writing group, was: "Is there evidence of clinically relevant treatment effects for therapies for cough in ILD". The literature search was generated following the creation of a PICO (population, intervention, comparison, outcome) element table (Table 1).

- 137
- 138



139 Literature search

The search methods used for this systematic review conformed to those outlined in "Methodologies for the development of CHEST guidelines and expert panel reports"¹². PubMed and the Cochrane Database of Systematic Reviews were searched for systematic reviews, and Scopus (includes EMBASE), PubMed, CINAHL (Cumulative Index of Nursing and Allied Health Literature) and the Cochrane Central Register of Controlled Trials were searched for other papers. All databases were searched from the earliest available date until February 2016. The search terms used are included in e-Appendix 1.

147

148 Eligibility criteria were reached by consensus between all of the authors (table 1). Given the 149 relative paucity of data in this field, we included all study types including case series of more 150 than 10 patients. The titles and abstracts were reviewed independently by two of the authors 151 (SB and JK) to identify potentially useful publications, based on the inclusion and exclusion 152 criteria. Where differences occurred, consensus was achieved between the two reviewers. 153 While a third independent reviewer was available if consensus could not be reached, lack of 154 consensus did not occur. The full text of those abstracts selected as potentially relevant were then reviewed independently (again by SB and JK) against the eligibility criteria. The 155 156 inclusion of a standardized cough outcome measure was acceptable as a primary or secondary 157 endpoint. Systematic reviews fulfilling the PICO criteria were selected in the preliminary 158 search for further analysis.

159

160 **Quality assessment**

All studies included at full text stage were randomized control trials and a quality assessment for each study was carried out using the Cochrane risk of bias tool¹³. To assess the quality of any systematic reviews, the Documentation and Appraisal Review Tool (DART) was used¹⁴.



164 There is no standardized tool available for quality assessing case series, and these studies 165 were assessed on criteria suggested by the Agency for Healthcare Research and Quality 166 (AHRQ)¹⁵.

167

168 Intervention Fidelity Assessment

During full text review, the methodology was examined (including online supplementation where available) to determine whether the investigators had systematically excluded common alternative causes of cough in their patient population prior to the intervention.

172

173 Grading Recommendations

174 Each recommendation is graded for the strength of the recommendation and quality of evidence, as is the case for all CHEST guidelines¹². The strength of recommendation is graded 175 176 as either strong (grade 1; likely to apply to almost all patients) or weak (grade 2; conditional 177 and only applying to selected patients). This grading is based on the quality of the evidence, the balance of risk versus potential benefit, the acceptability to patients and resource 178 179 considerations (for example: cost, availability, practicality / burden). The overall quality of 180 evidence for each recommendation is graded as high (grade A), moderate (grade B) or low 181 (grade C). A structured consensus-based Delphi approach was used to provide expert advice on guidance statements ¹⁶. In this regard, for a recommendation or suggestion to be approved 182 by the Expert Cough Panel, 75% of the eligible Panel members had to vote and 80% of those 183 184 voting had to strongly agree or agree with the statement. All panelists had the opportunity to 185 vote during the Delphi exercise; none were recused because of any conflicts of interest (e-Table 1). A patient representative who is a member of the Cough Panel provided patient-186 187 centered input for this guideline and approved of the suggestions contained herein.



189 **RESULTS**

190 Figure 1 summarizes the results of the systematic review. The majority of publications were 191 excluded by title or abstract alone, because they did not meet the inclusion criteria (702 of the 192 723 identified by the literature search). Twenty-six full text publications were reviewed and a 193 further 16 excluded. The most common reasons for exclusion at this stage were studies that 194 did not use a standardized or validated cough end-point or reviews that were non-systematic. Finally, 3 systematic reviews that met our inclusion criteria were excluded because the only 195 relevant studies they contained were already included in our review¹⁷⁻¹⁹. Three papers²⁰⁻²² 196 197 met the inclusion criteria but were published after the initial search and were subsequently 198 included on agreement of all the authors of the expert panel report. This process resulted in 10 199 studies included in this review (table 2). Of these, 5 concerned idiopathic pulmonary fibrosis, 200 3 investigated the use of inhaled corticosteroids in pulmonary sarcoidosis and 2 investigated 201 immunosuppressive therapies in scleroderma-related ILD.

202

203 Quality Assessment.

204 In general, the study quality was high in the trials selected for inclusion in the systematic 205 review. In the 8 RCTs included, the risk of bias was unclear in four of the studies in the area 206 of selection (table 3). A high risk of bias was identified in one study in selective outcome reporting ²³. This paper presents further, new analyses of the patients reported in an earlier 207 study ²⁴ and a post-hoc subgroup analysis. The methodology used to determine the subgroups 208 209 was unclear and this paper was therefore categorized as high risk of selective outcome 210 reporting. The two observational studies included have an inherent risk of bias, having no 211 placebo group and therefore have to be interpreted with caution.

212

213 Diagnosis of cough.



11

Cough in a patient with interstitial lung disease may be due to their underlying lung disease. However, because chronic cough occurs so commonly in the general population ²⁵, it is likely that in some cases other diseases may be responsible for the cough in patients with ILD (asthma, eosinophilic bronchitis, upper airway causes, drug therapy) or that cough may be unexplained. Patients with some ILDs are commonly treated with immunosuppressive therapies, and therefore, infectious causes should also be considered.

220

Only one of the included studies ²⁶ reported using a standardized diagnostic protocol to assess 221 cough prior to their intervention, otherwise it is unclear whether alternative causes of cough 222 were fully excluded (e.g. asthma, upper airway cough syndrome). In 4 studies ^{23,27-29}, cough 223 224 was not a major focus of the study and was only examined as a secondary or tertiary end-point 225 and often as part of a composite symptom score, so it is unlikely a cough diagnostic protocol was used. Horton et al and Birring et al did complete an assessment for identifiable causes of 226 cough but details of this assessment was not recorded. In Theodore, et al's study 30 , it is 227 228 possible that enrolled patients had been assessed; however, this was not reported.

229

Suggestion 1. For patients with ILD who present with a troublesome cough, we suggest that patients be assessed for progression of their underlying ILD, or complications from immunosuppressive treatment (e.g. drug side effect, pulmonary infection) and also be considered for further investigation / treatment trials for their cough according to guidelines for acute, subacute and chronic cough. (Ungraded Consensus- Based Statement)

236

237 Cough in IPF.



While the major trials of pirfenidone ^{24,31,32} and nintedanib ³³ have not reported data on the 238 impact of treatment on cough (table 4), Azuma et al ²³ used data from an earlier Japanese 239 pirfenidone trial²⁴ to report on the impact of pirfenidone on cough in IPF. These data were 240 not published in the initial study, but cough was included as part of a composite symptoms-241 242 based tertiary endpoint, together with dyspnea. Azuma et al report no significant difference overall in cough severity between the treatment arms (low and high dose pirfenidone) and 243 placebo. A recent study by van Manen et al ²² investigated the effect of pirfenidone on IPF 244 245 associated cough using validated cough outcome tools (Leicester Cough Monitor and Leicester Cough Questionnaire). There was a 34% reduction in objective cough frequency 246 247 and a clinically significant improvement in cough-related quality of life. The limitation of this 248 study was the lack of a control group.

249

250 Horton et al assessed cough as a primary endpoint in patients with cough and IPF³⁴. This was 251 a crossover trial where all patients received 12 weeks of treatment with thalidomide 50mg 252 daily, increasing in all but 1 patient to 100 mg if there was no improvement in cough after 2 253 weeks. All patients receiving thalidomide routinely received sodium docusate and vitamin B supplementation. The trial recruited small numbers of patients (24 patients, of whom 20 254 completed the trial). There was a statistically significant improvement in CQLQ score (a 255 validated quality of life questionnaire for cough) (decreased 11.37 points, p<0.001), cough 256 257 VAS (decreased by 31.2mm, p<0.001) and the SGRQ (decreased 11.7 points, p=0.001).

258

The side effect profile of thalidomide is an obvious concern, including its potential teratogenicity in women of childbearing age. Significantly more patients receiving thalidomide than placebo reported adverse events (77% vs 22%, p=0.001). The commonest side effects were constipation, dizziness and malaise. There were no serious adverse events



during treatment. Three patients required a dose reduction and the four withdrawals from the study were due to lack of interest (n=1) or disease progression and inability to return for study visits (n=3). Despite most patients experiencing side effects, all participants accepted the offer of continuing thalidomide treatment at the end of the trial.

267

Based on this evidence, the authors made a recommendation to the CHEST Expert Cough Panel that thalidomide be considered in those with IPF and chronic cough where alternative causes of cough were ruled out (Grade 2c). Only 67% of the panel voted in favor of this recommendation, and so it failed to pass and was removed (an approval score of 80% is required). It is likely that the practical barriers to prescribing Thalidomide in many countries, side effects and the limited evidence from a single, small trial were concerns amongst the CHEST Expert cough panel. The potential of thalidomide should be further investigated.

275

276 A recent proof of concept study investigated the efficacy of a novel inhaled cromolyn sodium formulation (PA101) in IPF patients with chronic cough ²⁰. PA101 is a high concentration 277 278 Cromolyn formulation delivered via a high efficiency eFlow nebulizer that achieves significantly higher drug deposition in the lung compared to existing formulations. There was 279 a 31% decrease in daytime mean cough frequency (the primary outcome measure) at day 14 280 281 with treatment when adjusted for placebo, but no statistically significant change in cough specific quality of life (measured by LCQ) or cough severity (measured by VAS score). The 282 283 drug was well tolerated with no severe or serious adverse events reported. PA101 Cromolyn 284 is not currently available to prescribe and further studies investigating its efficacy are awaited. 285



287 Proton Pump Inhibitors (PPIs) have long been used in IPF, and have been recommended in the 2015 American Thoracic Society IPF guidelines³⁵. 288 This was a conditional 289 recommendation and was based on two observational studies suggesting a slower decline in Vital Capacity in IPF patients taking PPI therapy. Kilduff et al ²⁶ investigated the efficacy of 290 291 high dose PPI and H2 receptor antagonist for 8 weeks in a small group of 18 patients with IPF. They assessed cough and gastro-esophageal reflux (GERD) objectively with esophageal 292 pH and impedance testing and cough monitoring. This was an uncontrolled study, but there 293 294 was no change in objective cough counts with PPI despite a decrease in acid reflux events. 295 This finding is in keeping with current evidence in unexplained chronic cough, where no 296 benefit in cough severity or quality of life has been observed with high dose esomeprazole in RCTs ^{36,37}. The CHEST unexplained chronic cough guideline therefore recommends against 297 using PPIs in patients with a negative workup for acid gastro-esophageal reflux. 298

299

Suggestion 2. For patients with IPF, chronic cough and a negative workup for acid
 gastroesophageal reflux, we suggest that proton pump inhibitor therapy not be
 prescribed. (Ungraded Consensus- Based Statement)

303

304

305 Cough in Sarcoidosis

Three trials investigating inhaled steroids in sarcoidosis were included (table 4). The type of patients included in each study differed, as did the dose of inhaled steroid used. Baughman et al studied 22 patients with acute sarcoidosis ²⁷. All patients were given concomitant oral steroids according to a standardized protocol. Ten patients were randomized to also take 880µg of fluticasone twice a day for 48 weeks, while the other 12 took a placebo inhaler. Cough was evaluated in a systematic way, although not using a validated cough tool. Cough



was reported as 'better' in more patients in the treatment arm (8 out of 10 in the treatment arm versus 6 out of 11 in the placebo arm); however, there was no statistically significant difference (p=0.36). There was no reduction in oral steroid requirements in the treatment arm.

DuBois et al also investigated the efficacy of inhaled fluticasone²⁸; but, in contrast, in patients 316 317 with established sarcoidosis (> 1 year from time of diagnosis) and at a slightly higher dose 318 compared to Baughman et al (2000 mcg daily versus 1760 mcg daily). Seventy-five percent of 319 patients were concomitantly taking oral prednisolone; but this was balanced across the 320 placebo and treatment groups. Cough was evaluated by a 4-point severity scale. This study 321 also recruited a relatively small number of patients (n=43), and although all patients reported 322 a decrease in cough severity at the end of the trial there was no significant difference between 323 the fluticasone and placebo arms.

324

Milman et al conducted a study in 1994 29 to investigate inhaled budesonide in established sarcoidosis. The number of subjects recruited was small, with 29 patients enrolled of whom 14 were treated with budesonide. There was no difference in cough severity, assessed with a 4-point scale, between the treatment and placebo arms (p=0.87). In the small subgroup on oral prednisolone (eight patients) there was no significant difference in the change in oral prednisolone dose with treatment versus placebo.

331

332 Suggestion 3. For patients with pulmonary sarcoidosis, we suggest that inhaled
333 corticosteroids should not be routinely prescribed to treat the chronic cough. (Grade
334 2C).

335

336 Cough in Scleroderma-related ILD.



337 The only studies included relating to connective tissue disease both concern Sclerodermarelated ILD. Theodore et al ³⁰ reported cough data from the Scleroderma Lung Study ¹⁰ that 338 339 randomized 158 patients to receive either cyclophosphamide or placebo for one year (with a 340 further 12 months follow up). A cough index, (which included a score of severity, frequency 341 and presence of phlegm) was assessed at baseline and 3 monthly intervals. Patients with 342 cough (73%) were more likely to have severe lung disease (established with TLCO and CT scans) and more diffuse scleroderma. The change in cough index with cyclophosphamide 343 therapy compared to placebo was not significantly different. There was a trend for greater 344 improvement in reported cough frequency in the treatment group, but this did not reach 345 346 statistical significance (p=0.56).

347

Tashkin et al reported cough data from the Scleroderma Lung Study II¹¹ that randomized 142 348 349 patients to either cyclophosphamide for 1 year followed by placebo for one year, or to 350 mycophenalate for 2 years. Given the interesting findings concerning cough in the first 351 scleroderma lung study, the investigators included the LCQ as a secondary outcome measure. 352 The presence of 'frequent cough' was determined by the responses to 2 questions in the St George's questionnaire. The paper reported the number of frequent coughers and LCQ scores 353 354 from both treatment arms combined and compares these to pre-treatment scores, not against 355 placebo. There was a decrease in the number of patients who reported frequent cough (from 356 61.7% to 45.7%, p = .0051); but there was no significant difference in the LCQ scores.

357

The Tashkin paper also reported an association between the presence of GERD-related GI symptoms and frequent cough. Of those patients with frequent cough, 77% reported GERDrelated GI symptoms at baseline compared with 59% of non-coughers (p= 0.025). In the small number of patients in whom GERD had resolved by 24 months (n=8), there was a



362 clinically significant improvement in mean LCQ score (from 14.3 to 17.9). In contrast, LCQ
363 scores were unchanged in patients who had persistent GERD at the end of the study.

364

365 **Refractory or Unexplained Cough in ILD.**

Patients with ILD and refractory cough have severely impaired quality of life ². In view of the limited treatment options for cough, the authors felt it was reasonable to manage such patients according to the CHEST unexplained cough guideline, particularly in those with advanced disease and poor quality of life. ³⁸

370

371 One treatment option recommended in the CHEST unexplained cough guideline is multimodality speech pathology therapy (SPT)³⁹ that has been reported to decrease objective 372 cough frequency and improve quality of life in patients with unexplained chronic cough ⁴⁰. A 373 374 similar efficacy has been reported for physiotherapy, speech and language therapy intervention (PSALTI)⁴¹. These therapies include educating patients about cough, teaching 375 376 cough suppression techniques, vocal hygiene and psychoeducational counseling. Gabapentin, 377 a neuromodulator, is also recommended, and in one RCT it decreased cough severity, frequency and improved quality of life in patients with refractory chronic cough ⁴². Side 378 379 effects were experienced by 10 of the 32 patients in the treatment group and were most 380 commonly confusion, dizziness, dry mouth, fatigue and nausea. There was a clinically and 381 statistically significant improvement in the mean LCQ score of 1.8 adjusted for placebo 382 effect. A more recent study combined the use of pregabalin and speech therapy and found a 383 larger improvement in LCQ and cough VAS compared to speech therapy alone, suggesting a combined approach may offer additional benefit ⁴³. 384



Suggestion 4. For patients with ILD and refractory chronic cough, we suggest trials of therapies recommended for patients with unexplained chronic cough according to the CHEST guidelines, with treatments such as gabapentin and multimodality speech pathology therapy, or entering into clinical trials if available. (Ungraded Consensus-Based Statement)

391

This review found no studies that met the inclusion criteria on the use of opiates for 392 393 management of chronic cough in ILD patients. There is a single RCT investigating slow release morphine (5mg twice daily) in refractory chronic cough patients ⁴⁴. This was a 394 395 positive study with a significant improvement in quality of life (a mean increase of 3.2 points 396 in the LCQ score). The most common side effects were constipation and drowsiness; 397 however, the drug was well tolerated and no patients withdrew from the study due to adverse 398 events. Opiates were not recommended by the CHEST Cough Panelists for unexplained 399 cough, because such a recommendation narrowly failed the guideline acceptance voting 400 threshold of 80%. Nevertheless, the authors of this guideline agree that opiates should be 401 considered for ILD patients with intractable cough when the cough has a substantial impact on 402 quality of life and when all alternative treatments have failed. Low dose opiates may be 403 helpful for symptomatic relief, and this will be particularly appropriate in the palliative care 404 setting. There are guidelines for the use of opiates for symptomatic relief of dry cough in patients receiving palliative care ⁴⁵; and, in the US, the CDC has published primary care 405 406 physician guidelines for opiate use in the treatment of chronic pain that includes helpful advice regarding the potential harms of opiate use and managing risk ⁴⁶. 407

408

409 Suggestion 5. For patients with chronic cough due to ILD, when alternative treatments
410 have failed and the cough is adversely affecting their quality of life, we suggest that



- 411 opiates be recommended for symptom control in a palliative care setting, with
- 412 reassessment of the benefits and risks at 1 week and then monthly before continuing.
- 413 (Ungraded Consensus- Based Statement)
- 414
- 415



416 **DISCUSSION**

4	1	7
4	T	1

418 **Diagnosis**

419 Interstitial lung diseases are a broad range of conditions that can affect the airways, lung 420 parenchyma and pulmonary vasculature. The involvement of any of these compartments can 421 lead to development of cough. Cough in patients with ILD can also be caused by ILD drug 422 therapy, infections and the presence of co-morbid conditions, such as GERD, upper airways 423 disease and asthma. It can therefore be challenging to determine the cause of a cough in a 424 patient with ILD. The approach to investigating patients with ILD-associated cough needs to 425 be individualized to the patient, and therefore a general approach such as that suggested in 426 Figure 2 has limitations and should be used only as a guide. ILD is likely to be the cause of 427 cough in a patient with evidence of disease progression, temporal association between onset of cough and disease progression and a favorable response to ILD therapy. 428

429

430 When cough persists, other causes should be investigated as per the CHEST Chronic Cough 431 Guidelines, specifically evaluating patients for the presence of asthma, non-asthmatic eosinophilic bronchitis, upper airways cough syndrome due to a variety of rhinosinus 432 conditions, and/or GERD ³⁸. If a cause is not established and the patient is significantly 433 434 troubled by their cough, in the opinion of this guideline group, it is reasonable to follow the 435 CHEST Unexplained Cough Guideline approach for managing cough patients because it is 436 possible that some patients with ILD and an unexplained cough have dysfunctional cough sensory nerves ⁴⁷. The term 'cough hypersensitivity syndrome' has been proposed to address 437 such patients ⁴⁸. In IPF, there is evidence of increased cough reflex sensitivity to both 438 439 capsaicin and nerve derived mediator substance P and increased levels of nerve growth factor have been reported in the airways of patients with IPF⁴⁹. 440



441	
442	Therapy
443	The approach to treating cough depends on a number of factors that include the type of ILD,
444	the availability of effective treatments for ILD, the risk/benefit profile of ILD and anti-tussive
445	treatments and the presence of co-morbid conditions that can cause cough. When ILD is the
446	suspected cause of cough, treatment for the underlying ILD should be considered on an
447	individual basis, particularly in patients with clear evidence of disease progression.
448	
449	<u>IPF</u>
450	In IPF, anti-fibrotic therapy with pirfenidone and nintedanib should be prescribed according
451	to ATS/ERS Guidelines, and not specifically for cough ³⁵ . It is possible that anti-fibrotic
452	therapy may reduce the severity of cough as suggested by the findings of an uncontrolled
453	study ²² but this needs confirmation in larger clinical trials.
454	
455	There are no controlled studies that have evaluated the use of systemic corticosteroids for IPF-
456	associated cough. The efficacy of corticosteroids for suppressing cough was evaluated in an
457	open label study of 6 patients by Hope-Gill et al, which did report a reduction in cough;
458	however, this did not meet our review inclusion criteria 49. Corticosteroids in IPF (in
459	combination with azathioprine and N-Acetyl-Cysteine; "triple therapy") have been associated
460	with increased mortality compared to placebo ⁵⁰ and the use of corticosteroids in IPF should
461	therefore be limited to patients that may be experiencing an exacerbation of IPF or who have
462	co-existing asthma or eosinophilic bronchitis.

463

Thalidomide has been investigated in IPF-associated cough in a single-center trial that 464 included a small number of patients³⁴. Thalidomide led to a significant improvement in 465



quality of life. The mechanism of action is thought to be anti-inflammatory, and possibly reduced cough sensory nerve activity ^{51,52} Thalidomide was, however, associated with significant side-effects ³⁴. While Thalidomide is not suggested for use in IPF-associated cough, the Committee does encourage further study of Thalidomide for cough because it might be helpful.

471

A recent pilot study of a novel formulation of inhaled cromolyn sodium (PA101) showed promise ²⁰. There was a >30% reduction in objective cough frequency between the treatment and placebo periods that is likely to be clinically significant. Although there was no significant change in the subjective cough severity and quality of life measures with treatment, the study was not powered to detect these and the treatment duration was short. The drug was well tolerated in this pilot study. A larger trial with a longer treatment period is needed to further assess its efficacy.

479

480 <u>Sarcoidosis</u>

In sarcoidosis, three trials of inhaled corticosteroids did not demonstrate a significant reduction in cough. The findings of the trials by duBois et al and Baughman et al did report trends towards improvement in cough^{27,28}. Further trials are needed to clarify the efficacy of inhaled corticosteroids, ideally in carefully selected patients with active, airway-centered disease and a clinically significant cough.

486

487 <u>Scleroderma</u>

488 Two connective tissue disease ILD studies met the inclusion criteria of this Guideline. They 489 evaluated immunosuppressive therapy in Scleroderma-associated ILD ^{21,30}. Both studies 490 demonstrated an association between cough and the severity of ILD. Theodore et al found no



491 significant improvement in cough with cyclophosphamide. Tashkin et al, however, did report 492 a decrease in subjective cough frequency with treatment (cyclophosphamide or 493 mycophenolate), but this was not associated with a significant improvement in quality of life. 494 The clinical significance of these therapies is therefore not clear and we suggest that 495 immunosuppressive therapy be prescribed for the underlying lung disease, rather than 496 specifically for cough. Further trials in scleroderma-associated ILD are needed, with cough 497 as the primary focus.

498

499 <u>Gastro-esophageal reflux Disease</u>

500 GERD is highly prevalent in both patients with ILD, such as IPF and scleroderma-associated 501 ILD and is therefore a potential cause of cough. There were no randomized controlled trials of proton pump inhibitor (PPI) therapy in ILD-associated cough. There was a small open 502 503 label study of high dose acid suppression therapy in IPF cough that did utilize validated objective and subjective cough end-points ²⁶; however, there was no improvement in cough 504 with PPI therapy. In patients with unexplained chronic cough and no ILD, two randomized 505 controlled trials did not demonstrate anti-tussive efficacy ^{36,37}. The CHEST Unexplained 506 507 Chronic Cough Guidelines therefore recommended that, in patients with a negative work-up 508 of acid gastro-esophageal reflux that included esophageal pH monitoring, PPI therapy should 509 not be prescribed 38 . A similar approach in IPF cough seems reasonable. In patients with 510 scleroderma-associated ILD, esophageal dysfunction is a key feature of the disease and 511 improvement in cough has been associated with improvements in GERD in the Tashkin study²¹. Therefore, a thorough approach investigating both acid and non-acid reflux in 512 513 scleroderma-associated ILD is reasonable until studies that guide best management become 514 available.



517 The severity of cough in ILD, particularly IPF, and its impact on quality of life warrants 518 consideration of further anti-tussive therapy options. In unexplained chronic cough, there are 519 several randomized controlled trials that support the use of neuromodulator drugs, such as gabapentin and pregabalin ^{42,43}, morphine ⁴⁴ and non-pharmacological interventions such as 520 speech pathology therapy ^{39,40} and Physiotherapy and Speech and Language Intervention 521 522 (PSALTI). ⁴¹ There are no studies that evaluated the efficacy of these therapies in ILD, and 523 they are urgently needed. The Guideline Panel acknowledged the paucity of evidence for use of general anti-tussive therapy in ILD, but recommended that they should be considered 524 525 because of the lack of availability of treatment options, and for use in a palliative care setting. 526 Opiates are likely the most controversial of these treatment options, due to concerns about 527 safety, and the potential for abuse and addiction. Indeed, morphine narrowly missed the 80% voting endorsement requirement of the ACCP Unexplained Cough Guidelines due to such 528 concerns³⁸. However, given the severity of lung disease and the poor prognosis associated 529 530 with many ILDs, the panel felt that morphine should be considered as a potential treatment for 531 ILD cough when quality of life is severely impacted. It is already used to some extent by clinicians to relieve both cough and dyspnea in ILD and for IPF patients with debilitating 532 cough 53 . 533

534

535 SUMMARY OF SYSTEMIC REVIEW RESULTS AND ITS LIMITATIONS

The present systemic review evaluated ten trials that investigated therapeutic interventions in ILD-associated cough. Cough was often not the primary focus of these studies, and was reported in retrospective analyses. The sample sizes were relatively small, and a variety of outcome assessments were used, not all of which were adequately validated. None of the interventions have been replicated in other RCTs. It is likely that there was heterogeneity in



the patient population under study that restricts the general reliability of the results. Intervention fidelity is recognized as a key aspect in the diagnosis of unexplained chronic cough and this factor was incompletely reported in the studies included in this review that indicates a possibility for indication-bias in the studies evaluated. These aspects of study design limit the strength of the conclusions.

546

547 **FUTURE DIRECTIONS**

A better estimate of the prevalence, severity and predictors of cough in ILD is needed in 548 larger population studies. Cough may be a potential biomarker in ILD, and therefore its 549 550 potential to assess disease severity and guide prognosis should be evaluated. There is a clear 551 need for randomized controlled trials of anti-tussive therapy in ILD. Validated cough outcome measures such as objective cough monitoring, visual analogue severity scales and 552 quality of life questionnaires, such as the CQLQ and LCQ should be used ⁵⁴⁻⁵⁶. Registries for 553 554 patients with ILD, such as IPF, should record the presence and severity of cough, and this can 555 be done with very simple tools such as the VAS or the Borg scale. There are now numerous 556 targets identified for novel anti-tussive therapies for unexplained cough. The antagonist of the P2X3 sensory nerve ion channel is one of the most promising and advanced in development. 557 558 A randomized controlled trial of a P2X3 antagonist in IPF-associated cough has been completed, but not reported ⁵⁷. The novel Cromolyn formulation PA101 shows promise in a 559 proof of concept trial but needs further evaluation ²⁰. Other anti-tussive targets include the 560 Neurokinin-1, TRPV4 and alpha-7 nicotinic receptors ⁵⁸⁻⁶⁰. Further studies of Thalidomide 561 and similar drugs are also warranted. 562

563

564



566

567

568 CONCLUSIONS

569 Cough associated with ILD can be due to underlying lung disease, and co-morbid conditions 570 such as upper airway disease or GERD. In some patients it remains unexplained. The 571 approach to managing ILD-associated cough needs to be individualized for the patient. 572 Further clinical trials of neuromodulator therapies and speech pathology/physiotherapy-based 573 cough suppression are needed.

574

575



Author contributions: All authors participated in developing the key clinical questions. The titles and abstracts and full text reviews and quality assessments of potentially relevant articles were independently carried out by SB and JK. SB and JK wrote the first draft of the manuscript. All authors reviewed and approved subsequent drafts and the final version of the manuscript.

583

584 Financial/nonfinancial disclosures:

The authors have reported to *CHEST* the following: S.S.B has received scientific advisory board/consultancy fees from Merck, Bayer, Patara and Menlo; speaker fees from Roche and conference travel from Boehringer Ingleheim. None declared (JEK, RSI, KK, KGL, JHR).

588 Endorsements:

589 None

590 **Collaborators:**

591 Todd M. Adams, MD (Webhannet Internal Medicine Associates of York Hospital), Kenneth

592 W. Altman, MD, PhD (Baylor College of Medicine, Houston, TX), Elie Azoulay, MD, PhD

593 (University of Paris, Paris, France), Alan F. Barker, MD (Oregon Health & Science

594 University, Portland, OR), Surinder S. Birring, MBChB, MD (Division of Asthma, Allergy

and Lung Biology, King's College London, Denmark Hill, London, United Kingdom), Fiona

596 Blackhall, MD, PhD (University of Manchester, Department of Medical Oncology,

597 Manchester, England), Louis-Philippe Boulet, MD, FCCP (Institut universitaire de

598 cardiologie et de pneumonlogie de Québec, Quebec, [IUCPQ], QC, Canada), Sidney S.

599 Braman, MD, FCCP (Mount Sinai Hospital, New York, NY), Christopher Brightling, MBBS,

600 PhD, FCCP (University of Leicester, Glenfield Hospital, Leicester, United Kingdom),

601 Priscilla Callahan-Lyon, MD (Adamstown, MD), Anne B. Chang, MBBS, PhD, MPH (Royal

602 Children's Hospital, Queensland, Australia), Paul Davenport, PhD (Department of



603	Physiological Sciences, University of Florida, Gainesville, FL), Ali A. El Solh, MD, MPH
604	(University at Buffalo, State University of New York, Buffalo, NY), Patricio Escalante, MD,
605	MSc, FCCP (Mayo Clinic, Rochester, MN), Stephen K. Field, MD (University of Calgary,
606	Calgary, AB, Canada), Dina Fisher, MD, MSc (University of Calgary, Respiratory Medicine,
607	Calgary, AB, Canada), Cynthia T. French, PhD, FCCP (UMass Memorial Medical Center,
608	Worcester, MA), Cameron Grant, MB ChB, PhD (University of Aukland, New Zealand),
609	Susan M. Harding, MD, FCCP, (Division of Pulmonary, Allergy and Critical Care Medicine,
610	University of Alabama at Birmingham, Birmingham, AL), Philip Gold, MD, MACP, FCCP
611	(Loma Linda University, Loma Linda, CA), Anthony Harnden, MB ChB, MSc (University of
612	Oxford, Oxford, England), Adam T. Hill, MB ChB, MD (Royal Infirmary and University of
613	Edinburgh, Edinburgh, Scotland), Richard S. Irwin, MD, Master FCCP (UMass Memorial
614	Medical Center, Worcester, MA), Peter J. Kahrilas, MD (Feinberg School of Medicine,
615	Northwestern University, Chicago, IL), Joanne Kavanagh, MBChB, (Division of Asthma,
616	Allergy and Lung Biology, King's College London, Denmark Hill, London, United
617	Kingdom), Karina A. Keogh, MD (Mayo Clinic, Rochester, MN), Kefang Lai, MD, PhD
618	(First Affiliated Hospital of Guangzhou Medical College, Guangzhou, China), Andrew P.
619	Lane, MD (Johns Hopkins University School of Medicine, Baltimore, MD), Kaiser Lim, MD
620	(Mayo Clinic, Rochester, MN), J. Mark Madison, MD, FCCP, (UMass Memorial Medical
621	Center, Worcester, MA), Mark A. Malesker, PharmD, FCCP (Creighton University School of
622	Pharmacy and Health Professions, Omaha, NE), Stuart Mazzone, PhD, FCCP (University of
623	Melbourne, Victoria, Australia), Lorcan Mc Garvey, MD (The Queens University Belfast,
624	Belfast, United Kingdom), Alex Molasoitis, PhD, MSc, RN (Hong Kong Polytechnic
625	University, Hong Kong, China), Abigail Moore, BM BCh, (University of Oxford, Oxford,
626	England), M. Hassan Murad, MD, MPH (Mayo Clinic, Rochester, MN), Mangala
627	Narasimhan, DO, FCCP (Hofstra-Northwell Health, Manhasset, NY), Huong O, Nguyen,

627 Narasimhan, DO, FCCP (Hofstra-Northwell Health, Manhasset, NY), Huong Q. Nguyen,



628	PhD, RN (Kaiser Permanente, Pasadena, CA), Peter Newcombe, PhD (School of Psychology
629	University of Queensland, Queensland, Australia), John Oppenheimer, MD (UMDNJ-Rutgers
630	University), Marcos I. Restrepo, MD, MSc, FCCP (South Texas Veterans Health Care
631	System, San Antonio), Mark Rosen, MD, Master FCCP (Icahn School of Medicine at Mount
632	Sinai, New York, NY), Bruce Rubin, MEngr, MD, MBA (Virginia Commonwealth
633	University, Richmond, VA), Jay H. Ryu, MD, FCCP (Mayo Clinic, Rochester, MN), Sonal
634	Singh, MD, MPH (UMass Memorial Medical Center, Worcester, MA), Jaclyn Smith, MB
635	ChB, PhD (University of Manchester, Manchester, England), Susan M. Tarlo, MBBS, FCCP
636	(Toronto Western Hospital, Toronto, ON, Canada), Julie Turmel, PhD (Quebec Heart and
637	Lung Institute, Laval University, Quebec), Anne E. Vertigan, PhD, MBA, BAppSc (SpPath)
638	(John Hunter Hospital, New South Wales, Australia), Gang Wang MD, PhD (Sichuan
639	University, West China Hospital, Chengdu, China), Miles Weinberger, MD, FCCP
640	(University of Iowa Hospitals and Clinics, Iowa City, IA).
641	
642	Role of sponsors
643	None
644	Other contributions
645	Additional information
646	
647 648 649 650 651 652 653 654 655 656	
657 658	





71 Table 1. Eligibility criteria

Criteria	Study requirements
Inclusion	 Published any time prior to time of search (February 2016) English language Any study design, case series (>10 subjects)
Population	 Age > 18 years Diagnosis of ILD, IPF or Sarcoidosis
Intervention	 Any pharmacological intervention or non-pharmacological intervention
Control / comparison	 Usual care / standard therapy or placebo (if applicable to study design)
Outcome	 Assessment of cough (measured using validated/standardized tool)





http://guide.medlive.cn/

Table 2. Study characteristics

Study	Study design	Anatomical work up for chronic cough	Duration	Intervention	Number randomised [intervention arm ; placebo arm]
IPF			I		
Azuma 2011	Post-hoc analysis of a multicenter randomized double blind placebo controlled trial	No	12 months	Pirfenidone 1200mg or 1800mg/day	275 [166 ; 109]
Horton 2012	Randomized double blind placebo controlled crossover	No	12 weeks	Thalidomide 100mg	24*
Kilduff 2014	Cohort study	Yes	8 weeks	High dose PPI (omeprazole 40mg bd or lansoprazole 30mg bd) plus ranitidine 300mg nocte	18
Birring 2017	Multicenter, randomized, double blind, placebo controlled, 2-cohort, 2-period cross-over trial	Yes	14 days	Inhaled cromolyn sodium (PA101) via eFlow nebulizer 40mg tds	24*
Van Manen 2017	Multi-center, prospective, observational study	No	12 weeks	Pirfenidone 2403mg/day**	43



SARCOID					
Milman 1994	Randomized double blind placebo controlled	No	12 months	Inhaled budesonide 1.2 or 2.0mg/day	21 [9 ; 12]
Du Bois 1999	Two center randomized double blind placebo controlled	No	6 months intervention + 2 months follow up	Inhaled fluticasone 2mg /day	43 [21 ; 22]
Baughman 2002	Multicenter randomized double blind placebo controlled	No	48 weeks	Inhaled fluticasone 880mcg bd	22 [10;12]
SCLERODERMA				<u> </u>	
Theodore 2012	Randomized double blind placebo controlled	No	12 months	Oral cyclophosphamide 2mg/kg monthly	158 [79 ; 79]
Tashkin 2017	Multicenter randomized double blind trial, 2 treatment arms	No	24 months	Mycophenylate target dose 1500mg bd for 24 months OR Cyclophosphamide target dose 2mg/kg/day for 12 months then placebo 12 months	142***

*Cross over trial design ** Communication with first author. Target dose; patients were up titrated from 801mg/day starting dose over a 2 week period. *** This study pooled the cough data from both treatment arms

Table 3. Risk of bias - RCTs

Azuma et al, 2011	Baughman et al, 2002	Birring et al, 2017	Du Bois et al, 1999	Horton et al, 2012	Milman et al, 1994	Tashkin et al, 2017	Theodore et al, 2012	
L	U	L	U	L	U	L	L	Random sequence generation (selection bias)
U	U	L	U	L	U	L	L	Allocation concealment (selection bias)
L	L	L	L	L	L	L	L	Blinding of participants and personnel (performance bias)
L	L	L	L	L	L	L	L	Blinding of outcome assessment (detection bias)
L	L	L	L	L	U	L	L	Incomplete outcome data addressed (attrition bias)
Н	L	L	L	L	L	L	L	Selective outcome reporting (reporting bias)
L	L	L	L	L	L	L	L	Other bias

H = high risk of bias

L = low risk of bias

U = unclear risk of bias

n/a = not applicable



Table 4. Treatment Effects

	Cough severity	Cough frequency (subjective)	Cough frequency (objective)	Cough QoL
Pirfenidone in IPF				
Azuma et al 2011	+	n/a	n/a	n/a
Van Manen et al 2017	+	n/a	+	+
Thalidomide in IPF		·		
Horton et al 2012	+	n/a	n/a	+
Acid suppression in IPF				
Kilduff et al 2014	n/a	n/a	-	n/a
Cromolyn in IPF				
Birring et al 2017	-	n/a	+	-
Inhaled steroid in pulmonary sarcoidosis				
Milman et al 1994	-	n/a	n/a	n/a
Du Bois et al 1999	-	n/a	n/a	n/a
Baughman et al 2002	-	n/a	n/a	n/a
Oral cyclophosphamide in SSc-ILD		1		
Theodore et al 2012	-	-	n/a	n/a
Tashkin et al 2017	n/a	+	n/a	-
Mycophenylate in SSc-ILD		l		1
Tashkin et al 2017	n/a	+	n/a	-

- + statistically significant improvement
- no statistically significant difference
- n/a not measured



REFERENCES:

- 1. Van Manen MJ, Birring SS, Vancheri C, et al. Cough in idiopathic pulmonary fibrosis. *European Respiratory Review*. 2016;25(141):278-286.
- 2. Key AL, Holt K, Hamilton A, Smith JA, Earis JE. Objective cough frequency in idiopathic pulmonary fibrosis. *Cough.* 2010;6(1):4.
- 3. Swigris JJ, Stewart AL, Gould MK, Wilson SR. Patients' perspectives on how idiopathic pulmonary fibrosis affects the quality of their lives. *Health and quality of life outcomes.* 2005;3(1):61.
- 4. Ryerson CJ, Abbritti M, Ley B, Elicker BM, Jones KD, Collard HR. Cough predicts prognosis in idiopathic pulmonary fibrosis. *Respirology*. 2011;16(6):969-975.
- 5. Jones RM, Hilldrup S, Hope-Gill BD, Eccles R, Harrison NK. Mechanical induction of cough in Idiopathic Pulmonary Fibrosis. *Cough.* 2011;7(1):2.
- 6. Birring S, Parker D, McKenna S, et al. Sputum eosinophilia in idiopathic pulmonary fibrosis. *Inflammation Research*. 2005;54(2):51-56.
- 7. Doherty M, Mister R, Pearson M, Calverley P. Capsaicin induced cough in cryptogenic fibrosing alveolitis. *Thorax.* 2000;55(12):1028-1032.
- 8. Tobin RW, Pope CE, Pellegrini CA, Emond MJ, Sillery J, Raghu G. Increased prevalence of gastroesophageal reflux in patients with idiopathic pulmonary fibrosis. *American journal of respiratory and critical care medicine.* 1998;158(6):1804-1808.
- 9. Sinha A, Lee KK, Rafferty GF, et al. Predictors of objective cough frequency in pulmonary sarcoidosis. *European Respiratory Journal.* 2016;47(5):1461-1471.
- 10. Tashkin DP, Elashoff R, Clements PJ, et al. Cyclophosphamide versus placebo in scleroderma lung disease. *New England Journal of Medicine*. 2006;354(25):2655-2666.
- 11. Tashkin DP, Roth MD, Clements PJ, et al. Mycophenolate mofetil versus oral cyclophosphamide in scleroderma-related interstitial lung disease (SLS II): a randomised controlled, double-blind, parallel group trial. *The Lancet Respiratory Medicine*. 2016;4(9):708-719.
- 12. Lewis SZ, Diekemper R, Ornelas J, Casey KR. Methodologies for the development of CHEST guidelines and expert panel reports. *Chest.* 2014;146(1):182-192.
- 13. Higgins JP, Altman DG, Gøtzsche PC, et al. The Cochrane Collaboration's tool for assessing risk of bias in randomised trials. *Bmj.* 2011;343:d5928.
- 14. Diekemper R, Ireland B, Merz L. P154 development of the documentation and appraisal review tool (Dart) for systematic reviews. *BMJ Qual Saf.* 2013;22(Suppl 1):61-62.
- 15. Viswanathan M, Ansari MT, Berkman ND, et al. Assessing the risk of bias of individual studies in systematic reviews of health care interventions. 2012.
- 16. Jones J, Hunter D. Consensus methods for medical and health services research. *BMJ: British Medical Journal.* 1995;311(7001):376.
- 17. Bajwah S, Ross JR, Peacock JL, et al. Interventions to improve symptoms and quality of life of patients with fibrotic interstitial lung disease: a systematic review of the literature. *Thorax.* 2013;68(9):867-879.
- 18. Paramothayan NS, Lasserson TJ, Jones P. Corticosteroids for pulmonary sarcoidosis. *The Cochrane Library.* 2005.
- 19. Loveman E, Copley VR, Colquitt J, et al. The clinical effectiveness and costeffectiveness of treatments for idiopathic pulmonary fibrosis: a systematic review and economic evaluation. *Health Technology Assessment.* 2015;19(20):1-336.



- 20. Birring SS, Wijsenbeek MS, Agrawal S, et al. A novel formulation of inhaled sodium cromoglicate (PA101) in idiopathic pulmonary fibrosis and chronic cough: a randomised, double-blind, proof-of-concept, phase 2 trial. *The Lancet Respiratory Medicine*. 2017;5(10):806-815.
- 21. Tashkin DP, Volkmann ER, Tseng C-H, et al. Improved cough and cough-specific quality of life in patients treated for scleroderma-related interstitial lung disease: results of Scleroderma Lung Study II. *Chest.* 2017;151(4):813-820.
- 22. van Manen MJ, Birring SS, Vancheri C, et al. Effect of pirfenidone on cough in patients with idiopathic pulmonary fibrosis. *European Respiratory Journal*. 2017;50(4):1701157.
- 23. Azuma A, Taguchi Y, Ogura T, et al. Exploratory analysis of a phase III trial of pirfenidone identifies a subpopulation of patients with idiopathic pulmonary fibrosis as benefiting from treatment. *Respiratory research.* 2011;12(1):143.
- 24. Taniguchi H, Ebina M, Kondoh Y, et al. Pirfenidone in idiopathic pulmonary fibrosis. *European Respiratory Journal.* 2010;35(4):821-829.
- 25. Ford AC, Forman D, Moayyedi P, Morice AH. Cough in the community: a cross sectional survey and the relationship to gastrointestinal symptoms. *Thorax.* 2006;61(11):975-979.
- 26. Kilduff CE, Counter MJ, Thomas GA, Harrison NK, Hope-Gill BD. Effect of acid suppression therapy on gastroesophageal reflux and cough in idiopathic pulmonary fibrosis: an intervention study. *Cough.* 2014;10(1):4.
- 27. Baughman R, Iannuzzi M, Lower E, et al. Use of fluticasone in acute symptomatic pulmonary sarcoidosis. *Sarcoidosis, vasculitis, and diffuse lung diseases: official journal of WASOG.* 2002;19(3):198-204.
- 28. Du Bois R, Greenhalgh P, Southcott A, Johnson NM, Harris T. Randomized trial of inhaled fluticasone propionate in chronic stable pulmonary sarcoidosis: a pilot study. *European Respiratory Journal*. 1999;13(6):1345-1350.
- 29. Milman N, Graudal N, Grode G, Munch E. No effect of high-dose inhaled steroids in pulmonary sarcoidosis: a double-blind, placebo-controlled study. *Journal of internal medicine.* 1994;236(3):285-290.
- 30. Theodore AC, Tseng C-H, Li N, Elashoff RM, Tashkin DP. Correlation of cough with disease activity and treatment with cyclophosphamide in scleroderma interstitial lung disease: findings from the Scleroderma Lung Study. *Chest.* 2012;142(3):614-621.
- 31. King Jr TE, Bradford WZ, Castro-Bernardini S, et al. A phase 3 trial of pirfenidone in patients with idiopathic pulmonary fibrosis. *New England Journal of Medicine*. 2014;370(22):2083-2092.
- 32. Noble PW, Albera C, Bradford WZ, et al. Pirfenidone in patients with idiopathic pulmonary fibrosis (CAPACITY): two randomised trials. *The Lancet*. 2011;377(9779):1760-1769.
- 33. Richeldi L, du Bois RM, Raghu G, et al. Efficacy and safety of nintedanib in idiopathic pulmonary fibrosis. *New England Journal of Medicine*. 2014;370(22):2071-2082.
- 34. Horton MR, Santopietro V, Mathew L, et al. Thalidomide for the treatment of cough in idiopathic pulmonary fibrosis: a randomized trial. *Annals of internal medicine*. 2012;157(6):398-406.
- 35. Raghu G, Rochwerg B, Zhang Y, et al. An official ATS/ERS/JRS/ALAT clinical practice guideline: treatment of idiopathic pulmonary fibrosis. An update of the 2011 clinical practice guideline. *American journal of respiratory and critical care medicine*. 2015;192(2):e3-e19.



- 36. Shaheen NJ, Crockett SD, Bright SD, et al. Randomised clinical trial: high-dose acid suppression for chronic cough–a double-blind, placebo-controlled study. *Alimentary pharmacology & therapeutics.* 2011;33(2):225-234.
- 37. Faruqi S, Molyneux ID, Fathi H, Wright C, Thompson R, Morice AH. Chronic cough and esomeprazole: A double-blind placebo-controlled parallel study. *Respirology*. 2011;16(7):1150-1156.
- 38. Gibson P, Wang G, McGarvey L, et al. Treatment of unexplained chronic cough: CHEST guideline and expert panel report. *Chest.* 2016;149(1):27-44.
- 39. Vertigan AE, Theodoros DG, Gibson PG, Winkworth AL. Efficacy of speech pathology management for chronic cough: a randomised placebo controlled trial of treatment efficacy. *Thorax.* 2006;61(12):1065-1069.
- 40. Ryan NM, Vertigan AE, Bone S, Gibson PG. Cough reflex sensitivity improves with speech language pathology management of refractory chronic cough. *Cough*. 2010;6(1):5.
- 41. Mitchell SAC, Garrod R, Clark L, et al. Physiotherapy, and speech and language therapy intervention for patients with refractory chronic cough: a multicentre randomised control trial. *Thorax.* 2017;72(2):129-136.
- 42. Ryan NM, Birring SS, Gibson PG. Gabapentin for refractory chronic cough: a randomised, double-blind, placebo-controlled trial. *The Lancet*. 2012;380(9853):1583-1589.
- 43. Vertigan AE, Kapela SL, Ryan NM, Birring SS, McElduff P, Gibson PG. Pregabalin and speech pathology combination therapy for refractory chronic cough: a randomized controlled trial. *CHEST Journal.* 2016;149(3):639-648.
- 44. Morice AH, Menon MS, Mulrennan SA, et al. Opiate therapy in chronic cough. *American journal of respiratory and critical care medicine*. 2007;175(4):312-315.
- 45. Excellence NIoHaC. Palliative Care Cough. *Clinical Knowledge Summaries* 2016, 2017.
- 46. Dowell D, Haegerich TM, Chou R. CDC guideline for prescribing opioids for chronic pain—United States, 2016. *Jama*. 2016;315(15):1624-1645.
- 47. Keller JA, McGovern AE, Mazzone SB. Translating cough mechanisms into better cough suppressants. *Chest.* 2017.
- 48. Morice AH, Millqvist E, Belvisi MG, et al. Expert opinion on the cough hypersensitivity syndrome in respiratory medicine. Eur Respiratory Soc; 2014.
- 49. Hope-Gill BD, Hilldrup S, Davies C, Newton RP, Harrison NK. A study of the cough reflex in idiopathic pulmonary fibrosis. *American journal of respiratory and critical care medicine.* 2003;168(8):995-1002.
- 50. Network TIPFCR. Prednisone, Azathioprine, and N-Acetylcysteine for Pulmonary Fibrosis. *New Engl. J. Med.* 2012;366(21):1968-1977.
- 51. Cundari S, Cavaletti G. Thalidomide chemotherapy-induced peripheral neuropathy: actual status and new perspectives with thalidomide analogues derivatives. *Mini reviews in medicinal chemistry*. 2009;9(7):760-768.
- 52. Ye Q, Chen B, Tong Z, et al. Thalidomide reduces IL-18, IL-8 and TNF-α release from alveolar macrophages in interstitial lung disease. *European Respiratory Journal*. 2006;28(4):824-831.
- 53. National Clinical Guideline C. National Institute for Health and Clinical Excellence: Guidance. *Diagnosis and Management of Suspected Idiopathic Pulmonary Fibrosis: Idiopathic Pulmonary Fibrosis*. London: Royal College of Physicians (UK)

National Clinical Guideline Centre.; 2013.



- 54. Boulet L-P, Coeytaux RR, McCrory DC, et al. Tools for assessing outcomes in studies of chronic cough: CHEST guideline and expert panel report. *Chest.* 2015;147(3):804-814.
- 55. Birring S, Prudon B, Carr A, Singh S, Morgan M, Pavord I. Development of a symptom specific health status measure for patients with chronic cough: Leicester Cough Questionnaire (LCQ). *Thorax.* 2003;58(4):339-343.
- 56. Lechtzin N, Hilliard ME, Horton MR. Validation of the Cough Quality-of-Life Questionnaire in patients with idiopathic pulmonary fibrosis. *Chest.* 2013;143(6):1745-1749.
- 57. Abdulqawi R, Dockry R, Holt K, et al. P2X3 receptor antagonist (AF-219) in refractory chronic cough: a randomised, double-blind, placebo-controlled phase 2 study. *The Lancet.* 2015;385(9974):1198-1205.
- 58. Smith JA, Kitt MM, Morice AH, et al. MK-7264, A P2X3 Receptor Antagonist, Reduces Cough Frequency In Patients With Refractory Chronic Cough: Results From A Randomized, Controlled, Phase 2b Clinical Trial. *B14. CLINICAL TRIALS ACROSS PULMONARY DISEASE*: Am Thoracic Soc; 2017:A7608-A7608.
- 59. Smith JA, Allman D, Badri H, et al. The neurokinin-1 receptor antagonist orvepitant is a novel anti-tussive therapy for chronic refractory cough: results from a phase 2 Study (VOLCANO-1). A101. ADVANCES IN COUGH, DYSPNEA, AND INTERVENTIONAL PULMONARY: American Thoracic Society; 2017:A2672-A2672.
- 60. Bonvini SJ, Birrell MA, Grace MS, et al. Transient receptor potential cation channel, subfamily V, member 4 and airway sensory afferent activation: Role of adenosine triphosphate. *Journal of Allergy and Clinical Immunology.* 2016;138(1):249-261. e212.



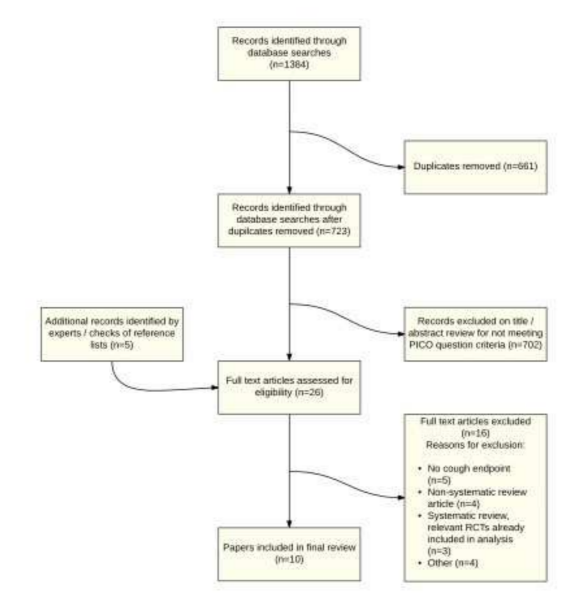
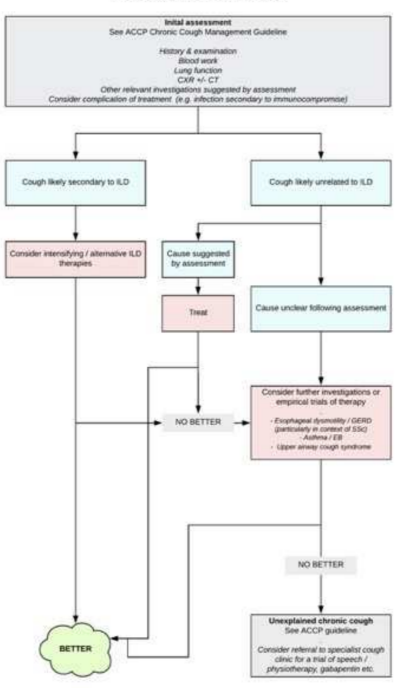


Figure 1. Systematic Review Flow Diagram



Figure 2. A proposed algorithm detailing a management approach to troublesome cough in an ILD patient



Troublesome cough in an ILD patient



CCEPTED MANUSCRIPT SCHEST Online Supplement

e-Appendix 1: Search terms

COUGH_ILD, Sarcoidosis, Bronchiolitis_ Search Strategies

Question 1: In a patient with ILD and cough, when should ILD be considered a cause of cough?

Question 2: Is there evidence of clinically relevant treatment effects for therapies for cough in ILD, Sarcoidosis and Bronchiolitis?

Limits (when available) were generally:

English language Humans Adult (age 19 and older) No editorials, comments or letters No date limits (full date ranges of individual databases)

Systematic Review Searches

PubMed - The subset filter of "AND systematic[sb]" was applied to the results of the PubMed searches (both questions 1 and 2). With this method, **15** systematic reviews or guidelines were identified. Among them were Cochrane reviews (also indexed in PubMed). The rest were systematic reviews or guidelines published by non-Cochrane groups.

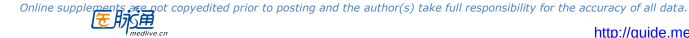
Cochrane Database of Systematic Reviews (Wiley – created in 1995, updated monthly) Additional searches were performed *directly* in the Cochrane Database of Systematic Reviews for both questions 1 and 2, using the same search configurations as for the CINAHL searches.

Of the reviews retrieved for both questions 1 and 2, the majority were focused on children. Only one relevant review (an update) was identified that was not already in the group. Systematic reviews were included in a group folder in the Question 2 (therapy) library.

PubMed Searches: (open access) - produced by the U.S. National Library of Medicine (includes indexed MEDLINE records) – 1950s to present

Question 1:

cough AND ("Lung Diseases, Interstitial" [Mesh:NoExp] OR interstitial lung disease* [tiab] OR "Alveolitis, Extrinsic Allergic"[Mesh:NoExp] OR extrinsic allergic alveolitis[tiab] OR "Anti-Glomerular Basement Membrane Disease"[Mesh] OR anti-glomerular basement membrane disease[tiab] OR "Histiocytosis, Langerhans-Cell"[Mesh] OR Langerhans-cell histiocytosis[tiab] OR "Idiopathic Interstitial Pneumonias"[Mesh] OR idiopathic interstitial pneumonia*[tiab] OR organizing pneumonia[tiab] OR cryptogenic fibrosing alveolitis[tiab] OR non-specific interstitial pneumonia[tiab] OR "Radiation Pneumonitis"[Mesh] OR radiation pneumonitis[tiab] OR "Sarcoidosis, Pulmonary"[Mesh] OR pulmonary sarcoidosis[tiab] OR hypersensitivity pneumonitis[tiab] OR sarcoidosis OR obliterative bronchiolitis[tiab] OR bronchiolitis obliterans[tiab] OR cryptogenic obliterative bronchiolitis[tiab] OR acute bronchiolitis[tiab] OR diffuse pan-bronchiolitis[tiab] OR follicular bronchiolitis[tiab] OR



ACCEPTED MANUSCRIPT SCHEST Online Supplement

aspiration bronchiolitis[tiab] OR }desquamative interstitial pneumonia}[tiab]) AND (etiology OR cause OR epidemiology[mh] OR epidemiology[tiab] OR prevalence OR diagnosis OR assessment[tiab] OR investigation) NOT (editorial[pt] OR letter[pt] OR comment[pt])

Filters: Humans; English; Adult: 19+ years

Question 2:

cough AND ("Lung Diseases, Interstitial" [Mesh:NoExp] OR interstitial lung disease* [tiab] OR "Alveolitis, Extrinsic Allergic"[Mesh:NoExp] OR extrinsic allergic alveolitis[tiab] OR "Anti-Glomerular Basement Membrane Disease"[Mesh] OR anti-glomerular basement membrane disease[tiab] OR "Histiocytosis, Langerhans-Cell"[Mesh] OR Langerhans-cell histiocytosis[tiab] OR "Idiopathic Interstitial Pneumonias"[Mesh] OR idiopathic interstitial pneumonia*[tiab] OR organizing pneumonia[tiab] OR cryptogenic fibrosing alveolitis[tiab] OR non-specific interstitial pneumonia[tiab] OR "Radiation Pneumonitis"[Mesh] OR radiation pneumonitis[tiab] OR "Sarcoidosis, Pulmonary"[Mesh] OR pulmonary sarcoidosis[tiab] OR hypersensitivity pneumonitis[tiab] OR sarcoidosis OR obliterative bronchiolitis[tiab] OR bronchiolitis obliterans[tiab] OR cryptogenic obliterative bronchiolitis[tiab] OR acute bronchiolitis[tiab] OR diffuse pan-bronchiolitis[tiab] OR follicular bronchiolitis[tiab] OR aspiration bronchiolitis[tiab] OR }desquamative interstitial pneumonia}[tiab]) AND (treatment OR therapy OR drug therapy OR thalidomide OR corticosteroids OR cyclophosphamide OR salbutamol OR proton pump inhibitors OR immunosuppressants OR pirfenidone OR nintendanib OR erythromycin OR macrolides OR muscarinic antagonists OR "beta agonist"[tiab]) NOT (editorial[pt] OR letter[pt] OR comment[pt])

Filters: Humans; English; Adult: 19+ years

Scopus Searches (Elsevier) - includes EMBASE records: 1823 to present

Question 1:

TITLE-ABS-KEY(cough AND (

OR {extrinsic allergic alveolitis}²⁸⁻³⁰ OR {anti-glomerular basement membrane disease} OR {Langerhans-cell histiocytosis} OR {idiopathic interstitial pneumonia} OR {organizing pneumonia} OR {cryptogenic fibrosing alveolitis} OR {non-specific interstitial pneumonia} OR {radiation pneumonitis} OR {pulmonary sarcoidosis} OR {hypersensitivity pneumonitis} OR sarcoidosis OR {obliterative bronchiolitis} OR {bronchiolitis obliterans} OR {cryptogenic obliterative bronchiolitis} OR {bronchiolitis obliterans} OR {cryptogenic obliterative bronchiolitis} OR {bronchiolitis obliterans} OR {cryptogenic obliterative bronchiolitis} OR {acute bronchiolitis} OR {diffuse pan-bronchiolitis} OR {follicular bronchiolitis} OR {aspiration bronchiolitis} OR {desquamative interstitial pneumonia}) AND (etiology OR cause OR epidemiology OR prevalence OR diagnosis OR assessment OR investigation)) AND (adult OR adolescent)) AND (LIMIT-TO(DOCTYPE,"ar") OR LIMIT-TO(DOCTYPE,"re") OR LIMIT-TO(DOCTYPE,"cp") OR LIMIT-TO(DOCTYPE,"ip")) AND (LIMIT-TO(LANGUAGE,"English"))



SCHEST Online Supplement

Question 2:

TITLE-ABS-KEY(cough AND ({interstitial lung disease} OR {extrinsic allergic alveolitis} OR {anti-glomerular basement membrane disease} OR {Langerhans-cell histiocytosis} OR {idiopathic interstitial pneumonia} OR {organizing pneumonia} OR {cryptogenic fibrosing alveolitis} OR {non-specific interstitial pneumonia} OR {radiation pneumonitis} OR {pulmonary sarcoidosis} OR {hypersensitivity pneumonitis} OR sarcoidosis OR {obliterative bronchiolitis } OR {bronchiolitis obliterans } OR {cryptogenic obliterative bronchiolitis } OR {acute bronchiolitis} OR {diffuse pan-bronchiolitis} OR {follicular bronchiolitis} OR {aspiration bronchiolitis} OR {desquamative interstitial pneumonia}) AND (treatment OR therapy OR {drug therapy} OR thalidomide OR corticosteroids OR cyclophosphamide OR salbutamol OR {proton pump inhibitors} OR immunosuppressants OR pirfenidone OR nintendanib OR erythromycin OR macrolides OR {muscarinic antagonists} OR {beta agonist}) AND (adult OR adolescent)) AND (LIMIT-TO(DOCTYPE, "ar") OR LIMIT-TO(DOCTYPE,"re") OR LIMIT-TO(DOCTYPE,"cp") OR LIMIT-TO(DOCTYPE,"ip")) AND (LIMIT-TO(LANGUAGE,"English"))

Duplicates found between Scopus and PubMed searches were identified and removed in the EndNote library.

<u>CINAHL Searches</u> (EBSCOhost platform) – Cumulative Index of Nursing and Allied Health Literature - (1981 – present)

Question 1:

cough AND ("interstitial lung disease" OR "extrinsic allergic alveolitis" OR "anti-glomerular basement membrane disease" OR "Langerhans-cell histiocytosis" OR "idiopathic interstitial pneumonia" OR "organizing pneumonia" OR "cryptogenic fibrosing alveolitis" OR "nonspecific interstitial pneumonia" OR "radiation pneumonitis" OR "pulmonary sarcoidosis" OR "hypersensitivity pneumonitis" OR sarcoidosis OR "obliterative bronchiolitis" OR "bronchiolitis obliterans" OR "cryptogenic obliterative bronchiolitis" OR "acute bronchiolitis" OR "diffuse pan-bronchiolitis" OR "follicular bronchiolitis" OR "aspiration bronchiolitis" OR "desquamative interstitial pneumonia") AND (etiology OR cause OR epidemiology OR prevalence OR diagnosis OR assessment OR investigation) Limits: All Adult and English

Question 2:

medlive cn

cough AND ("interstitial lung disease" OR "extrinsic allergic alveolitis" OR "anti-glomerular basement membrane disease" OR "Langerhans-cell histiocytosis" OR "idiopathic interstitial pneumonia" OR "organizing pneumonia" OR "cryptogenic fibrosing alveolitis" OR "nonspecific interstitial pneumonia" OR "radiation pneumonitis" OR "pulmonary sarcoidosis" OR "hypersensitivity pneumonitis" OR sarcoidosis OR "obliterative bronchiolitis" OR "bronchiolitis obliterans" OR "cryptogenic obliterative bronchiolitis" OR "acute bronchiolitis" OR "diffuse pan-bronchiolitis" OR "follicular bronchiolitis" OR "aspiration bronchiolitis" OR "desquamative interstitial pneumonia") AND (treatment OR therapy OR drug therapy OR

pents are not copyedited prior to posting and the author(s) take full responsibility for the accuracy of all data. Online suppler

thalidomide OR corticosteroids OR cyclophosphamide OR salbutamol OR proton pump inhibitors OR immunosuppressants OR pirfenidone OR nintendanib OR erythromycin OR macrolides OR "muscarinic antagonists" OR "beta agonist") *Limits: All Adult and English*

Duplicates found between CINAHL and PubMed searches were identified and removed in the EndNote library.

Cochrane Central Register of Controlled Trials (CENTRAL) (Wiley)

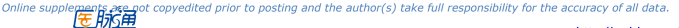
Question 1:

cough AND ("interstitial lung disease" OR "extrinsic allergic alveolitis" OR "anti-glomerular basement membrane disease" OR "Langerhans-cell histiocytosis" OR "idiopathic interstitial pneumonia" OR "organizing pneumonia" OR "cryptogenic fibrosing alveolitis" OR "nonspecific interstitial pneumonia" OR "radiation pneumonitis" OR "pulmonary sarcoidosis" OR "hypersensitivity pneumonitis" OR sarcoidosis OR "obliterative bronchiolitis" OR "bronchiolitis obliterans" OR "cryptogenic obliterative bronchiolitis" OR "acute bronchiolitis" OR "diffuse pan-bronchiolitis" OR "follicular bronchiolitis" OR "aspiration bronchiolitis" OR "desquamative interstitial pneumonia") AND (etiology OR cause OR epidemiology OR prevalence OR diagnosis OR assessment OR investigation)

Question 2:

cough AND ("interstitial lung disease" OR "extrinsic allergic alveolitis" OR "anti-glomerular basement membrane disease" OR "Langerhans-cell histiocytosis" OR "idiopathic interstitial pneumonia" OR "organizing pneumonia" OR "cryptogenic fibrosing alveolitis" OR "nonspecific interstitial pneumonia" OR "radiation pneumonitis" OR "pulmonary sarcoidosis" OR "hypersensitivity pneumonitis" OR sarcoidosis OR "obliterative bronchiolitis" OR "bronchiolitis obliterans" OR "cryptogenic obliterative bronchiolitis" OR "bronchiolitis obliterans" OR "cryptogenic obliterative bronchiolitis" OR "acute bronchiolitis" OR "diffuse pan-bronchiolitis" OR "follicular bronchiolitis" OR "aspiration bronchiolitis" OR "desquamative interstitial pneumonia") AND (treatment OR therapy OR drug therapy OR thalidomide OR corticosteroids OR cyclophosphamide OR salbutamol OR proton pump inhibitors OR immunosuppressants OR pirfenidone OR nintendanib OR erythromycin OR macrolides OR "muscarinic antagonists" OR "beta agonist")

Duplicates found between CENTRAL and other database search results were identified and removed in the EndNote library



http://guide.medlive.cn/

Section CHEST[®] Online Supplement

	Surinder S. Birring, MD	Richard S. Irwin, MD, Master FCCP	Joanne E. Kavanagh, MBChB	Karina Keogh, MD	Kaiser G. Lim, MD	Jay H. Ryu, MD, FCCP
	Financial COI	Financial COI	Financial COI	Financial COI	Financial COI	Financial COI
Suggestions	Description of COI	Description of COI	Description of COI	Description of COI	COI	Description of COI
1. For patients with ILD who present with a troublesome cough, we suggest that patients be assessed for progression of their underlying ILD, or complications from immunosuppressive treatment (e.g. drug side effect, pulmonary infection) and also be considered for further investigation / treatment trials for their cough according to guidelines for acute,						
subacute and chronic cough. (Ungraded Consensus- Based Statement)	none	none	none	none	none	none
2. For patients with IPF, chronic cough and a negative workup for acid gastroesophageal reflux, we suggest that proton pump inhibitor therapy should not be prescribed. (Ungraded						
Consensus- Based Statement) 3. For patients with pulmonary sarcoidosis, we suggest that inhaled corticosteroids should not be routinely prescribed to treat the chronic cough.		none	none	none	none	
(Grade 2C).	none	none	none	none	none	
4. For patients with ILD and refractory chronic cough, we suggest trials of therapies recommended for patients with unexplained chronic cough according to the CHEST guidelines, with treatments such as gabapentin and multimodality speech pathology therapy, or entering into clinical trials if available. (Ungraded Consensus- Based						
Statement)	none	none			none	
5. For patients with chronic cough due to ILD, when alternative treatments have failed and the cough is adversely affecting their quality of life, we suggest that opiates be recommended for symptom control in a palliative care setting with reassessment of the benefits and risks at 1 week and then monthly before continuing. (Ungraded						
Consensus- Based Statement)	none	none	none	none	none	
						-

181250

Online supplements are not copyedited prior to posting and the author(s) take full responsibility for the accuracy of all data.