

1 **Real-life evidence in ERS clinical practice guidelines: from foes to friends**

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17 Throughout the past decades, European Respiratory Society (ERS) Task Forces have produced and published
18 clinical practice guidelines (CPGs), statements, technical standards and other documents to synthesize and
19 summarize bodies of evidence for caregivers and thereby improve healthcare quality in respiratory medicine.
20 Among these various types of documents, only CPGs can propose recommendations for clinical practice. As
21 such, they need to rely on a very strong methodology to limit the risk of recommending suboptimal care.
22 Traditionally, the highest levels of evidence come from randomised controlled trials (RCTs).^{1,2} However, this
23 does not mean that other types of research should be excluded from evidence synthesis as part of guidelines
24 development processes. Indeed, they could usefully complement RCTs, provided that their methods are
25 rigorous, and the results are properly analysed and transparently interpreted.^{3,4} In this editorial, we
26 summarize how real-life evidence could and should be integrated in ERS CPGs.

27 **The development of ERS clinical practice guidelines**

28 The first crucial task of an ERS Task Force developing a CPG is to carefully consider the research questions
29 and outcomes of interest. Only thereafter, a systematic review of the literature as well as an assessment of
30 the quality of evidence can be performed (Figure 1). For the latter, the ERS uses Grading of
31 Recommendations, Assessment, Development and Evaluation (GRADE), an approach adopted and
32 recommended by many organisations including the National Institute for Health and Care Excellence (NICE),
33 the American Thoracic Society (ATS) and the World Health Organization (WHO).⁵ This rigorous method
34 considers a number of factors in addition to risk of bias for assessing the quality of evidence. Moreover, it
35 ensures a transparent linkage between evidence and recommendations when applying the Evidence to
36 Decision (EtD) framework for grading of the strength of a recommendation.⁶

37 The GRADE approach can be used for data originating from RCTs as well as observational studies.⁵ When
38 following the GRADE approach, the developers of guidelines must evaluate the risk of bias, inconsistency,
39 indirectness, imprecision, as well as publication bias to assess the certainty of evidence (Table 1).⁵ This applies
40 for data from both RCTs and observational studies. For the latter, large effect sizes, dose responses and

41 opposing biases may lead to an upgrading of the quality of evidence. This systematic approach that takes
42 every aspect of a published study into account makes it possible to ultimately adjust and grade the quality of
43 evidence as “very low”, “low”, “moderate” or “high”. In a next step, the GRADE EtD framework allows for
44 additional considerations such as balance of benefits and harms, values and preferences, feasibility, equity,
45 acceptability and resource use.⁶

46 This systematic approach makes it possible for the expert panel to transparently draw their final conclusions
47 while taking various aspects and perspectives into consideration, and make recommendations that are
48 supported by the evidence.⁷

49 **Real-life evidence**

50 The historical understanding that RCTs produce the evidence with the highest quality has often been
51 challenged because patient populations in these studies often are selected, not reflecting the patients seen
52 in everyday clinical practice.⁸ Hence, clinically important data from real-life may be missed and not
53 sufficiently emphasized by healthcare professionals and policymakers. A brilliant example of the disparity
54 between patients included in RCTs and real-life cohorts has recently been published by Brown et al. (2018).⁹
55 When comparing data from 342 patients against trial eligibility criteria from 37 RCTs evaluating biological
56 therapies for severe asthma, less than 10% of patients in their real-life cohort were found to be eligible.⁹
57 Similar concerns were reported regarding other major lung diseases, like chronic obstructive pulmonary
58 disease (COPD), lung cancer and bronchiectasis.¹⁰⁻¹⁶ In such cases, the recommendations might not be
59 applicable to most patients.

60 Although the ERS applies a very strict methodological approach and considers both randomized and
61 observational studies, the generalizability of our CPGs might even be improved by the inclusion of real-life
62 evidence from other sources like administrative databases or healthcare registers.³ These sources can
63 complement RCTs by: (1) confirming or challenging their generalizability for different populations or settings,
64 (2) exploring clinically relevant outcomes not available in RCTs, (3) providing safety data, and (4) allowing to

65 explore possible determinants of treatment effects to be further confirmed in prospective RCTs.³ In some
66 cases, when it is not possible to perform an RCT due to, e.g., ethical or feasibility reasons, real-life evidence
67 might even be the only way to generate new data.

68 **The ERS promotes an integrative approach in science and guideline development**

69 Several ERS initiatives were implemented recently to close the gap that often exists between evidence
70 generated from RCTs and real-life. With the ERS Clinical Research Collaborations (CRCs), the respiratory
71 community has the possibility to build international research networks to conduct pan-European pragmatic
72 trials that are generalizable and sufficiently large to impact clinical practice ([https://www.ersnet.org/science-
73 and-research/clinical-research-collaboration-application-programme/](https://www.ersnet.org/science-and-research/clinical-research-collaboration-application-programme/)). Furthermore, the ERS offers
74 investigators the opportunity to promote their research by endorsing pragmatic trials
75 (<https://www.ersnet.org/science-and-research/pragmatic-trials-endorsement/>). Pragmatic trials can be
76 endorsed when they investigate respiratory diseases, meet stringent criteria of quality and are not
77 dependent on a single sponsor from the pharmaceutical industry or another for-profit entity. In addition,
78 CRCs provide an excellent platform to collaboratively develop and use data from healthcare registers.¹⁷
79 Hence, the ERS promotes every type of evidence that can lead to a better understanding of a certain
80 respiratory condition, as long as the highest quality standards are satisfied.³

81 **Conclusion: how to integrate real-life evidence in ERS GCPs**

82 The crucial mission for ERS Task Forces developing CPGs is to appreciate not only data originating from RCTs,
83 but also other sources, to get the best picture of the current evidence and draw solid conclusions.^{3,4} For CPGs,
84 the ERS Guidelines Working Group recommends a thorough process with the selection of a limited number
85 of clinical questions in a PICO (Population, Intervention, Comparison, Outcome) format that can include data
86 both from RCTs and observational studies and that are assessed via a systematic review and the application
87 of GRADE.¹⁸ These questions can be complemented with additional non-comparative questions that are
88 addressed via a narrative review of the literature. The guideline panel then chooses outcomes that are critical

89 or important for clinical decision making and that are relevant for the patients. For the final
90 recommendations, the EtD framework should be systematically applied both for PICO and narrative
91 questions. Real-life evidence that has not been considered in the systematic or narrative reviews should be
92 considered and taken into account in the EtD framework.⁴

93 With this approach, we ensure that ERS CPGs give recommendations that are transparent, trustworthy and
94 clinically relevant both for clinicians and patients.

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98 **Conflicts of interest**

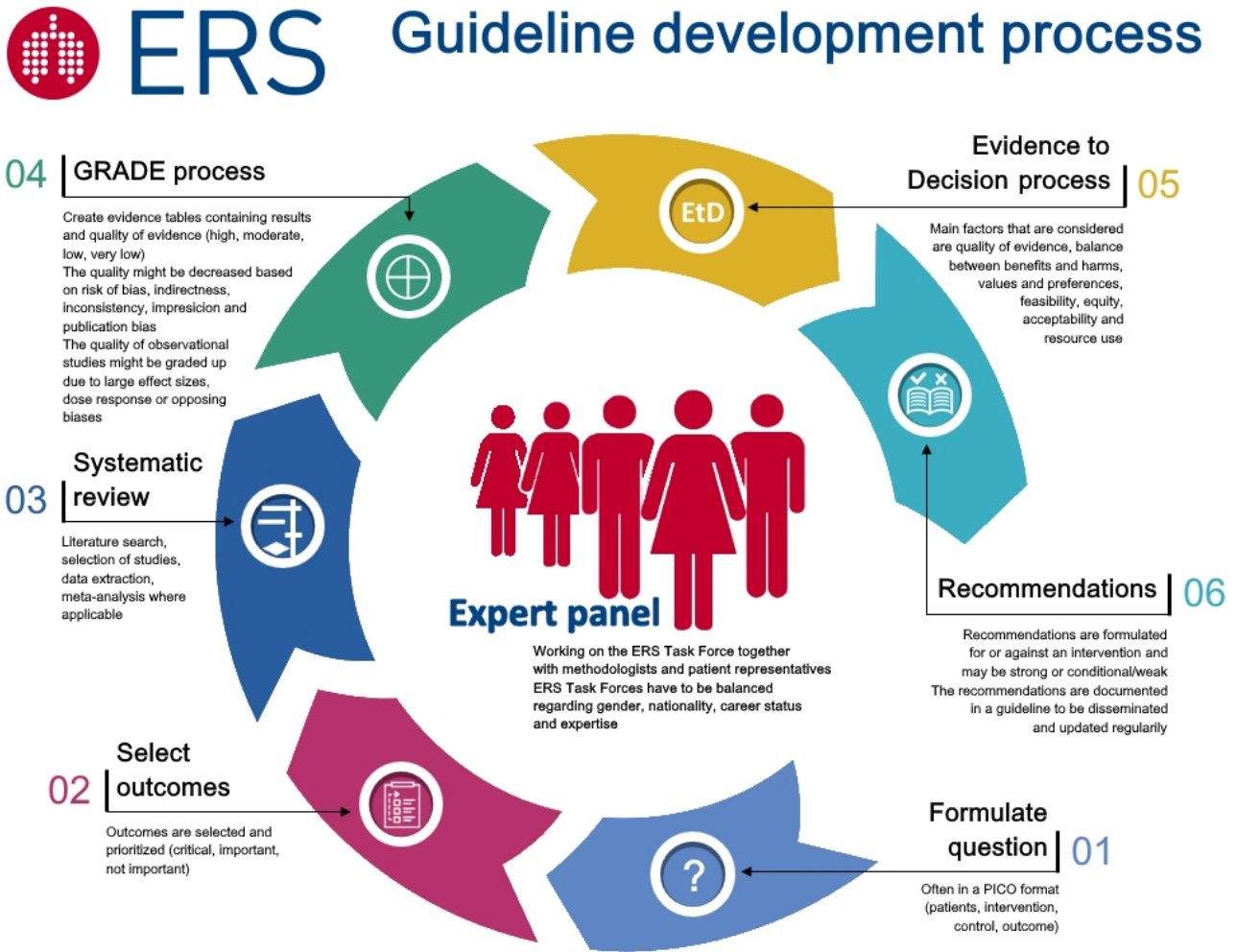
99 **XXX**

100 **References**

- 101 1. Prazma CM, Wenzel SE, Price R, Ortega HG. The impact of duration of oral corticosteroid use on co-
102 morbidities in a severe asthma population treated with mepolizumab. *Am J Respir Crit Care Med*.
103 2015;191(no pagination). [https://www.cochranelibrary.com/central/doi/10.1002/central/CN-](https://www.cochranelibrary.com/central/doi/10.1002/central/CN-01472365/full)
104 01472365/full
- 105 2. Concato J, Shah N, Horwitz RI. Randomized, controlled trials, observational studies, and the
106 hierarchy of research designs. *N Engl J Med*. 2000;342(25):1887-1892.
107 doi:10.1056/NEJM200006223422507
- 108 3. Roche N, Anzueto A, Bosnic Anticevich S, et al. The importance of real-life research in respiratory
109 medicine: manifesto of the Respiratory Effectiveness Group. *Eur Respir J*. 2019;54(3):1901511.
110 doi:10.1183/13993003.01511-2019
- 111 4. Gershon AS, Lindenauer PK, Wilson KC, et al. Informing Healthcare Decisions with Observational

- 112 Research Assessing Causal Effect. An Official American Thoracic Society Research Statement. *Am J*
113 *Respir Crit Care Med*. 2021;203(1):14-23. doi:10.1164/rccm.202010-3943ST
- 114 5. Schünemann H, Brożek J, Guyatt G, Oxman A E. GRADE handbook for grading quality of evidence
115 and strength of recommendations. Updated October 2013. The GRADE Working Group, 2013. The
116 GRADE Working Group. Published October 2013. www.guidelinedevelopment.org/handbook
- 117 6. Alonso-Coello P, Schünemann HJ, Moberg J, et al. GRADE Evidence to Decision (EtD) frameworks: a
118 systematic and transparent approach to making well informed healthcare choices. 1: Introduction.
119 *BMJ*. 2016;353:i2016. doi:10.1136/bmj.i2016
- 120 7. Li SA, Alexander PE, Reljic T, et al. Evidence to Decision framework provides a structured “roadmap”
121 for making GRADE guidelines recommendations. *J Clin Epidemiol*. 2018;104:103-112.
122 doi:10.1016/j.jclinepi.2018.09.007
- 123 8. Sherman RE, Anderson SA, Dal Pan GJ, et al. Real-world evidence - What is it and what can it tell us?
124 *N Engl J Med*. 2016;375(23):2293-2297. doi:10.1056/NEJMs1609216
- 125 9. Brown T, Jones T, Gove K, et al. Randomised controlled trials in severe asthma: selection by
126 phenotype or stereotype. *Eur Respir J*. 2018;52(6):1801444. doi:10.1183/13993003.01444-2018
- 127 10. Halpin DMG, Kerkhof M, Soriano JB, Mikkelsen H, Price DB. Eligibility of real-life patients with COPD
128 for inclusion in trials of inhaled long-acting bronchodilator therapy. *Respir Res*. 2016;17(1):120.
129 doi:10.1186/s12931-016-0433-5
- 130 11. Scichilone N, Basile M, Battaglia S, Bellia V. What Proportion of Chronic Obstructive Pulmonary
131 Disease Outpatients Is Eligible for Inclusion in Randomized Clinical Trials? *Respiration*.
132 2014;87(1):11-17. doi:10.1159/000355082
- 133 12. Kruis AL, Ställberg B, Jones RCM, et al. Primary Care COPD Patients Compared with Large
134 Pharmaceutically-Sponsored COPD Studies: An UNLOCK Validation Study. Schooling CM, ed. *PLoS*
135 *One*. 2014;9(3):e90145. doi:10.1371/journal.pone.0090145
- 136 13. Travers J, Marsh S, Caldwell B, et al. External validity of randomized controlled trials in COPD. *Respir*

- 137 *Med.* 2007;101(6):1313-1320. doi:10.1016/j.rmed.2006.10.011
- 138 14. Chalmers JD, McDonnell MJ, Rutherford R, et al. The generalizability of bronchiectasis randomized
139 controlled trials: A multicentre cohort study. *Respir Med.* 2016;112:51-58.
140 doi:10.1016/j.rmed.2016.01.016
- 141 15. Heuvers ME, Wisnivesky J, Stricker BH, Aerts JG. Generalizability of results from the National Lung
142 Screening Trial. *Eur J Epidemiol.* 2012;27(9):669-672. doi:10.1007/s10654-012-9720-8
- 143 16. Pahu L, Burgel P-R, Roche N, Paillasseur J-L, Chanez P. Randomized controlled trials of
144 pharmacological treatments to prevent COPD exacerbations: applicability to real-life patients. *BMC*
145 *Pulm Med.* 2019;19(1):127. doi:10.1186/s12890-019-0882-y
- 146 17. Brightling C, Genton C, Bill W, et al. ERS clinical research collaborations: Underpinning research
147 excellence. *Eur Respir J.* 2018;52(3). doi:10.1183/13993003.01534-2018
- 148 18. Huang X, Lin J, Demner-Fushman D. Evaluation of PICO as a knowledge representation for clinical
149 questions. *AMIA Annu Symp Proc.* Published online 2006:359-363. Accessed August 4, 2020.
150 <http://www.fpin.org/>
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Factors that can reduce the quality of evidence in RCTs and observational studies				
Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias
Limitations in the study design and execution	Unexplained heterogeneity of results	Indirect comparisons or differences in study populations, interventions or outcomes	Wide confidence intervals due to few patients and few events	Systematic under- or overestimation of a beneficial or harmful effect due to selective publication of studies
Factors that can increase the quality of evidence in observational studies				
Large magnitude of an effect	Dose-response gradient	Plausible residual confounding		
Point estimates for relative risks or hazard ratios way below or above 1	The presence of a dose-response gradient may increase the confidence in the results	The absence of residual confounding would have increased the intervention's effects		