## The COVID-NMA Project: Building an Evidence Ecosystem for the COVID-19 Pandemic

Isabelle Boutron, MD, PhD; Anna Chaimani, PhD; Joerg J. Meerpohl, MD; Asbjørn Hróbjartsson, MD, PhD, MPhil; Declan Devane, PhD; Gabriel Rada, MD; David Tovey, MBChB; Giacomo Grasselli, MD; and Philippe Ravaud, MD, PhD\* for the COVID-NMA consortium

ven before the coronavirus disease 2019 (COVID-19) pandemic, the ability of the evidence synthesis model to meet the needs of stakeholders was challenged (1, 2). There are too many low-quality systematic reviews that mainly address pairwise comparisons and are rarely updated, resulting in redundancies and gaps. Producing high-quality, up-to-date systematic reviews requires substantial time and resources. In addition, although evidence synthesis is directly affected by the quality of primary research, interaction is limited between the evidence generation and synthesis communities. These issues have been highlighted and exacerbated by the COVID-19 pandemic, where stakeholders urgently need relevant, accessible, up-to-date, and trustworthy syntheses of high-quality evidence to inform their decisions. Thousands of randomized controlled trials (RCTs) have been initiated during the pandemic, and their results are frequently rushed to publication or communicated through non-peer-reviewed preprints. The situation is further complicated by changes in the questions of interest and trial components (such as standard of care) as the pandemic develops (3).

To tackle COVID-19, we developed and implemented a previously proposed model (4, 5) to address the challenges and help to connect evidence generation, synthesis, and decision making. Rather than focusing on 1 specific treatment or comparison, the COVID-NMA project provides a living mapping of all trials and a comprehensive living synthesis of all available trial evidence evaluating the effect of interventions for the prevention or treatment of COVID-19 (Figure). We developed a master protocol (6) and subprotocols dedicated to specific questions, which are discussed and agreed on by a steering committee.

Every week, we screen the COVID-19 database produced by the World Health Organization's International Clinical Trials Registry Platform to identify eligible RCTs. The living mapping produced provides a description of all registered RCTs. The data retrieved and extracted can be explored through interactive data visualizations to identify research gaps and help prioritize and improve future trials.

We are also conducting a living systematic review based on a living protocol (6) that is scalable to stakeholders' evolving needs. All changes in the protocol (for example, primary study design and outcomes) are discussed by a steering committee and reported transparently. As part of the living process, we do a systematic search daily, collect data as soon as we identify any trial that has published results or is available in preprint,

and assess risk of bias fully using the Cochrane Risk of Bias Tool, version 2.0 (7). We provide the descriptive data online and produce forest plots of appropriately pooled data with GRADE (Grading of Recommendations Assessment, Development and Evaluation) summary-of-findings tables and evidence profiles. We have developed a tool to automatically identify new versions or publication of preprints. We contact trialists at the outset (that is, trial registration) to request information (protocol) and inform them of the outcomes (consistent with the core outcome sets developed by the COMET [Core Outcome Measures in Effectiveness Trials] initiative [8, 9]) that should be reported to enable their trial to be incorporated into the meta-analyses. When results are available, we systematically request from trial authors any missing data and update the reviews accordingly. We have established robust quality control processes in collaboration with the Cochrane Bias Methods Group. Collectively, COVID-NMA data are used to conduct systematic reviews on specific questions, meta-analyses of individual participant data (IPD), and network meta-analyses and to support the guideline development process and health decision making. Our databases can also be shared to allow guideline developers to do their own analyses.

To improve research planning, we monitor trials' quality related to outcomes, completeness of reporting (that is, adherence to some CONSORT [Consolidated Standards of Reporting Trials] items), risk of bias, and data sharing (intended and realized). As a feedback loop, we provide trialists and funders the results of this monitoring to increase the value of COVID-19 trials research. We also send automatic e-mails to investigators of completed trials to encourage them to post results on registries (10) and share IPD, and we have developed a secure process to enable them to do this at no cost.

Our collaborative project involves an international consortium of 85 persons, including methodologists, clinicians, and statisticians. On 31 August 2020, our research mapping identified 1686 registered RCTs, of which 944 are recruiting. Overall, 54% have fewer than 100 participants. We have screened more than 42 000 records and reported detailed data for 45 RCTs, with forest plots for all comparisons. We have contacted

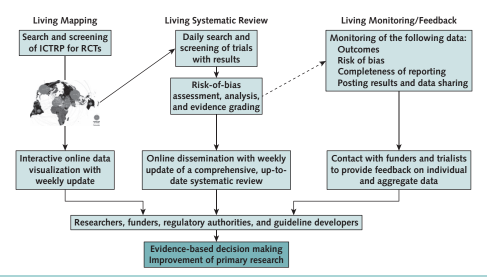
See also:

Related article

This article was published at Annals.org on 15 September 2020.

<sup>\*</sup> For members of the COVID-NMA consortium, see the Appendix (available at Annals.org).

Figure. Process of the COVID-NMA project.



The project aims to provide an up-to-date mapping of trials; a comprehensive, critical, up-to-date synthesis of all available trial-based evidence about the efficacy and safety of interventions for the prevention or treatment of coronavirus disease 2019; and a living monitoring on trial planning, conduct, and reporting. ICTRP = International Clinical Trials Registry Platform; RCT = randomized controlled trial.

about 1000 investigators of ongoing trials and requested missing data from 45 authors.

This new approach is creating challenges and threats. First, sustainability is an issue as the crisis continues. We developed COVID-NMA with the support of many volunteers from various countries who were available during the containment period but must now return to normal activities. As the amount of data increases, we need to move to a long-term and sustainable structure with a website that is more accessible and useful to end users. The resources necessary to maintain this model are critical because the volume of evidence is increasing, the scope is expanding at end users' request (for example, new focus on vaccine trials), and new sources (clinical study reports) or new types of data (such as IPD) are becoming available. We need funders to provide long-term funding for this platform. This would be far more cost-effective than funding a disparate and uncoordinated series of systematic reviews on narrow research questions.

Second, some cultural issues exist. The success of this approach depends entirely on the acceptance of and engagement with this model by stakeholders, in particular funders and trialists. Some may be reluctant to add new outcomes, adhere to reporting guidelines, or share IPD because this involves change in culture, as well as time and effort. We hope that the urgency associated with the COVID-19 pandemic, combined with external pressure, may help to overcome these barriers.

Governance of the project is an important consideration. We must ensure that volunteers and researchers involved in the platform receive the appropriate reward and recognition for their contributions. We are developing transparent processes for both the researchers involved and the users of the data, and our work is overseen by an independent steering committee.

Overall, the present crisis unmasks the shortcomings of the current synthesis model and provides a

strong impetus for change and improvement. We hope COVID-NMA plays a role in this work.

From Université de Paris, Centre of Research Epidemiology and Statistics (CRESS), Inserm, and Cochrane France, Paris, France (I.B., A.C., D.T., P.R.); Institute for Evidence in Medicine, Medical Center, and Faculty of Medicine, University of Freiburg, and Cochrane Germany, Cochrane Germany Foundation, Freiburg, Germany (J.J.M.); Centre for Evidence-Based Medicine Odense, University of Southern Denmark, and Odense University Hospital, Odense, Denmark (A.H.); Evidence Synthesis Ireland, Cochrane Ireland, and HRB Trials Methodology Research Network, National University of Ireland, Galway, Ireland (D.D.); Epistemonikos Foundation and UC Evidence Center, Cochrane Chile Associated Center, Pontificia Universidad Católica de Chile, Santiago, Chile (G.R.); Cochrane France, Paris, France; and Fondazione IRCCS Ca' Granda Ospedale Maggiore Policlinico and University of Milan, Milan, Italy (G.G.).

**Note:** The protocol can be accessed at https://zenodo.org/record/3744599. All data are shared on our website, https://covid-nma.com.

Financial Support: This work received some funding from the Agence Nationale de la Recherche, the World Health Organization, Cochrane France, Centre of Research in Epidemiology and Statistics, Centre d'Epidémiologie Clinique (GHU Cochin, Hôtel Dieu, Assistance Publique Hôpitaux de Paris, and Université de Paris), Federal Ministry of Health (Germany), and the Centre National de la Recherche Scientifique.

**Disclosures:** Disclosures can be viewed at www.acponline.org /authors/icmje/ConflictOfInterestForms.do?msNum=M20-5261.

Corresponding Author: Isabelle Boutron, MD, PhD, Centre d'épidémiologie clinique, Hôpital Hôtel-Dieu, 1 place du Par-

2 Annals of Internal Medicine Annals.org

vis Notre-Dame, 75004 Paris, France; e-mail, isabelle.boutron @aphp.fr.

Current author addresses and author contributions are available at Annals.org.

Ann Intern Med. doi:10.7326/M20-5261

## References

- 1. Boutron I, Créquit P, Williams H, et al. Future of evidence ecosystem series: 1. Introduction evidence synthesis ecosystem needs dramatic change. J Clin Epidemiol. 2020;123:135-142. [PMID: 32145367] doi:10.1016/j.jclinepi.2020.01.024
- 2. Gurevitch J, Koricheva J, Nakagawa S, et al. Meta-analysis and the science of research synthesis. Nature. 2018;555:175-182. [PMID: 29517004] doi:10.1038/nature25753
- 3. Guan WJ, Ni ZY, Hu Y, et al; China Medical Treatment Expert Group for Covid-19. Clinical characteristics of coronavirus disease 2019 in China. N Engl J Med. 2020;382:1708-1720. [PMID: 32109013] doi:10.1056/NEJMoa2002032
- 4. Créquit P, Boutron I, Meerpohl J, et al. Future of evidence ecosystem series: 2. Current opportunities and need for better tools and

- methods. J Clin Epidemiol. 2020;123:143-152. [PMID: 32145369] doi: 10.1016/j.jclinepi.2020.01.023
- 5. Ravaud P, Créquit P, Williams HC, et al. Future of evidence ecosystem series: 3. From an evidence synthesis ecosystem to an evidence ecosystem. J Clin Epidemiol. 2020;123:153-161. [PMID: 32147384] doi:10.1016/j.jclinepi.2020.01.027
- 6. Boutron I, Chaiman A, Meerpohl JJ, et al. Interventions for preventing and treating COVID-19: protocol for a living mapping of research and a living systematic review. Zenodo. 2020. doi:10.5281/zenodo.3744599
- 7. Sterne JAC, Savovic J, Page MJ, et al. RoB 2: a revised tool for assessing risk of bias in randomised trials. BMJ. 2019;366:l4898. [PMID: 31462531] doi:10.1136/bmj.l4898
- 8. WHO Working Group on the Clinical Characterisation and Management of COVID-19 infection. A minimal common outcome measure set for COVID-19 clinical research. Lancet Infect Dis. 2020;20: e192-e197. [PMID: 32539990] doi:10.1016/S1473-3099(20)30483-7
- 9. **COMET Initiative.** Core outcome set developers' response to COVID-19. 7 July 2020. Accessed at www.comet-initiative.org/Studies /Details/1538 on 31 August 2020.
- 10. Maruani A, Boutron I, Baron G, et al. Impact of sending email reminders of the legal requirement for posting results on Clinical Trials.gov: cohort embedded pragmatic randomized controlled trial. BMJ. 2014;349:q5579. [PMID: 25239625] doi:10.1136/bmj.q5579

Annals.org Annals of Internal Medicine 3

**Current Author Addresses:** Drs. Boutron, Chaimani, and Ravaud: Centre d'épidémiologie clinique, Hôpital Hôtel-Dieu, 1 place du Parvis Notre-Dame, 75004 Paris, France.

Dr. Meerpohl: Institute for Evidence in Medicine, Medical Center - University of Freiburg,

Breisacher Strasse 86, 79110 Freiburg, Germany.

Dr. Hróbjartsson: Centre for Evidence-Based Medicine Odense (CEBMO), University of Southern Denmark and Odense University Hospital, Kløvervæget 10, 13th Floor, Gate 112, 5000 Odense C, Denmark.

Dr. Devane: School of Nursing and Midwifery, Aras Moyola, NUI Galway, Galway, Ireland.

Dr. Rada: Holanda 895, Providencia, Santiago de Chile, Chile. Dr. Tovey: 46 Dalmore Road, London SE21 8HB, England.

Dr. Grasselli: Fondazione IRCCS Ca' Granda Ospedale Maggiore Policlinico, Via Francesco Sforza 35, 20122 Milan, Italy.

**Author Contributions:** Conception and design: I. Boutron, A. Chaimani, J.J. Meerpohl, A. Hróbjartsson, D. Devane, D. Tovey, G. Grasselli, P. Ravaud.

Analysis and interpretation of the data: I. Boutron, J.J. Meerpohl, D. Devane, G. Rada, D. Tovey, P. Ravaud.

Drafting of the article: I. Boutron, A. Chaimani, J.J. Meerpohl, D. Devane, D. Tovey, P. Ravaud.

Critical revision of the article for important intellectual content: I. Boutron, J.J. Meerpohl, A. Hróbjartsson, D. Devane, G. Grasselli.

Final approval of the article: I. Boutron, A. Chaimani, J.J. Meerpohl, A. Hróbjartsson, D. Devane, G. Rada, D. Tovey, G. Grasselli, P. Ravaud.

Provision of study materials or patients: G. Rada.

Statistical expertise: A. Chaimani.

Obtaining of funding: J.J. Meerpohl, P. Ravaud.

Administrative, technical, or logistic support: I. Boutron, G. Rada, P. Ravaud.

Collection and assembly of data: I. Boutron, D. Devane, G. Rada, D. Tovey, P. Ravaud.

## Appendix: Members of the COVID-NMA Consortium

Names are reported in alphabetical order.

Members of the COVID-NMA consortium who contributed to this work but did not author it: Solaf Alawadhi<sup>1,2</sup>, Sihem Amer-Yahia<sup>3</sup>, Chiara Arienti<sup>4</sup>, David Auber<sup>5</sup>, Camila Ávila<sup>6</sup>, Aïda Bafeta<sup>2</sup>, Fulvia Baldassarre<sup>7</sup>, Rita Banzi<sup>8</sup>, Julien Barnier<sup>9</sup>, Julia Baudry<sup>10</sup>, Hanna Bergman<sup>11</sup>, Claudia Bollig<sup>12</sup>, Hillary Bonnet<sup>1,13</sup>, Marinette Bouet<sup>14</sup>, Mohand Boughanem<sup>15</sup>, Brian Buckley<sup>11</sup>, Guillaume Cabanac<sup>15</sup>, Sarah Charpy<sup>2</sup>, David Chavalarias<sup>17</sup>, Yaolong Chen<sup>18</sup>, Astrid Chevance<sup>2</sup>, Sarah Cohen-Boulakia<sup>19</sup>, Elise Cogo<sup>11</sup>, Françoise Conil<sup>20</sup>, Emmanuel Coquery<sup>20</sup>, Mauricia Davidson<sup>2,16</sup>, Laura De Nale<sup>16</sup>, Elise Diard<sup>16</sup>, Taoufiq Dkaki<sup>15</sup>, Bastien Doreau<sup>14</sup>, Merwan El Asri<sup>19</sup>, Theodoros Evrenoglou<sup>1,16</sup>, Alice Fabbri<sup>22</sup>, Robin Featherstone<sup>23</sup>, Gilles Feron<sup>24</sup>, Gabriel Ferrand<sup>16</sup>, Leopold Fezeu<sup>10</sup>, Mathilde Fouet<sup>25</sup>, Joly Ghanawi<sup>26</sup>, Lina Ghosn El Chall<sup>16</sup>, Carolina Graña<sup>2,16</sup>, François Grolleau<sup>1</sup>, Benoit Groz<sup>19</sup>, Mohand-Saïd Hacid<sup>20</sup>, Candyce Hamel<sup>11</sup>, Camilla Hansen<sup>22</sup>, Nicholas Henschke<sup>11</sup>, Ameer Hohlfeld<sup>28</sup>, Chantal Julia<sup>10</sup>, Dimitris Mavridis<sup>29</sup>, Brice Meyer<sup>14</sup>, Silvia Minozzi<sup>31</sup>, Jose G. Moreno<sup>15</sup>, Nivantha Naidoo<sup>2</sup>, Van Thu Nguyen<sup>16</sup>, Theodora Oikonomidi<sup>1,16</sup>, Matthew Page<sup>32</sup>, Jennifer Petkovic<sup>11</sup>, Elizabeth Pienaar<sup>27</sup>, Olivier Pierre<sup>2</sup>, Katrin Probyn<sup>11</sup>, Fiona Quirke<sup>33</sup>, Pierre Ripoll<sup>20</sup>, Carolina Riveros<sup>2,16</sup>, Philippe Rivière<sup>20</sup>, Marie Sauvant<sup>14</sup>, Jelena Savovic<sup>35</sup>, Christine Schmucker<sup>30</sup>, Yanina Sguassero<sup>11</sup>, Jonathan Sterne<sup>36</sup>, Farouk Toumani<sup>14</sup>, Gemma Villanueva<sup>11</sup>, Romain Vuillemot<sup>20</sup>, Jun Xia<sup>36</sup>, Xuan Yu<sup>18</sup>, Emina Zoletic<sup>1,2</sup>, and Pierre Zweigenbaum<sup>38</sup>.

Members of the COVID-NMA consortium who authored this work: Isabelle Boutron<sup>1,2,16</sup>, Anna Chaimani<sup>1,16</sup>, Declan Devane<sup>21</sup>, Giacomo Grasselli<sup>26</sup>, Asbjørn Hróbjartsson<sup>22</sup>, Joerg J. Meerpohl<sup>12,29</sup>, Gabriel Rada<sup>6,33</sup>, Philippe Ravaud<sup>1,2,16</sup>, and David Tovey<sup>16</sup>.

- 1. Université de Paris, France
- 2. Centre of Research in Epidemiology and StatisticS (CRESS UMR1153), Methods team, Inserm, France
- 3. Laboratoire d'Informatique de Grenoble (LIG), CNRS, France
- 4. IRCCS Fondazione Don Carlo Gnocchi, Italy
- 5. Laboratoire Bordelais de Recherche en Informatique (LaBRI), Université Bordeaux I, France
- 6. Epistemonikos Foundation, Chile
- 7. McMaster University, Canada
- 8. Center for Health Regulatory Policies, Istituto di Ricerche Farmacologiche Mario Negri IRCCS, Italy
- 9. Centre Max Weber, CNRS, France
- 10. Centre of Research in Epidemiology and Statistics (CRESS UMR1153), Eren team, France
- 11. Cochrane Response, United Kingdom
- 12. Cochrane Germany, Cochrane Germany Foundation, Freiburg, Germany
- 13. Bordeaux Pharmacoepi ADERA, France
- 14. Laboratoire d'Informatique, de Modélisation et d'Optimisation des Systèmes (LIMOS), CNRS, Université Clermont Auvergne, France
- 15. Université Toulouse 3 Paul Sabatier Institut de Recherche en Informatique de Toulouse -IRIT UMR 5505, France
- 16. Cochrane France
- Institut des Systèmes Complexes de Paris IDF (ISC-PIF), CNRS, France
- 18. WHO Collaborating Centre for Guideline Implementation and Knowledge Translation & Chinese GRADE Centre, Lanzhou University, China
- 19. Laboratoire de recherche en Informatique (LRI), CNRS, Université Paris-Saclay, France
- Laboratoire d'InfoRmatique en Image et Systèmes d'information (LIRIS), CNRS, Université Claude Bernard Lyon 1, France

- 21. Evidence Synthesis Ireland, Cochrane Ireland and HRB-Trials Methodology Research Network, National University of Ireland, Galway, Ireland
- 22. Centre for Evidence Based Medicine Odense (CEBMO), University of Southern Denmark and Odense University Hospital, Denmark
- 23. Cochrane Editorial and Methods Department, Cochrane Central
- 24. French National Research Institute for Agriculture, Food and Environment (INRAE), France
- 25. Service de Neurochirurgie, Hôpital d'Instruction des Armées Percy (HIA), France
- 26. The Collaborative Approach to Meta-Analysis and Review of Animal Data from Experimental Studies (CAMARADES), Centre for Clinical Brain Sciences, University of Edinburgh, Scotland
- 27. Department of Anesthesia, Intensive Care and Emergency, Fondazione IRCCS Ca' Granda Ospedale Maggiore Policlinico, Department of Pathophysiology and Transplantation, University of Milan, Italy
- 28. Cochrane South Africa, South African Medical Research Council, South Africa
- 29. Department of Primary Education, University of Ioannina, Greece
- 30. Institute for Evidence in Medicine, Medical Center & Faculty of Medicine, University of Freiburg, Freiburg, Germany

- 31. Cochrane Review Group on Drugs and Alcohol; International GRADE Working Group; Department of Epidemiology, Lazio Regional Health Service, Italy
- 32. Research Methodology Division, School of Public Health and Preventive Medicine, Monash University, Australia
- 33. Health Research Board-Trials Methodology Research Network (HRB-TMRN), NUI Galway, Ireland
- 34. UC Evidence Center, Cochrane Chile Associated Center, Pontificia Universidad Católica de Chile, Santiago, Chile
- 35. Population Health Sciences, Bristol Medical School, University of Bristol, UK; NIHR CLAHRC West, University Hospitals Bristol and Weston NHS Foundation Trust, United Kingdom
- 36. Bristol Medical School, Bristol Population Health Science Institute, University of Bristol, United Kingdom
- 37. Nottingham Ningbo GRADE Centre, The Nottingham China Health Institute, the University of Nottingham Ningbo, China
- 38. Laboratoire d'Informatique pour la Mécanique et les Sciences de l'Ingénieur (LIMSI), CNRS, France

Annals of Internal Medicine Annals.org