

American Society of Hematology 2021 L Street NW, Suite 900, Washington, DC 20036 Phone: 202-776-0544 | Fax 202-776-0545 bloodadvances@hematology.org

# ASH living guidelines on use of anticoagulation for thromboprophylaxis in patients with COVID-19: executive summary

Tracking no: ADV-2024-014219R1

Deborah Siegal (University of Ottawa, Canada) Eric Tseng (St. Michael's Hospital, Division of Hematology/Oncology, University of Toronto, Canada) Holger Schünemann (McMaster University, Canada) Pantep Angchaisuksiri (Ramathibodi Hospital, Mahidol University, Thailand) Adam Cuker (University of Pennsylvania, United States) Kathryn Dane (Johns Hopkins Hospital, United States) Maria DeSancho (Weill Cornell Medicine/New York Presbyterian Hospital, United States) David Diuguid (Columbia University Medical Center, United States) Daniel Griffin (Columbia University Medical Center, United States) Frederikus Klok (Leiden University Medical Center, Netherlands) Alfred Lee (Yale University School of Medicine, United States) Ignacio Neumann (Universidad San Sebastian, Chile) Ashok Pai (Kaiser Permanente, United States) Marc Righini (Geneva University Hospital, Switzerland) Kristen Sanfilippo (Washington University School of Medicine in St. Louis, United States) Deirdra Terrell (University of Oklahoma Health Sciences Center, United States) Elie Akl (American University of Beirut, Lebanon) Reyad Al Jabiri (University of Jordan, Jordan) Yazan Al Jabiri (Lincoln Medical and Mental Health Center, United States) Angela Barbara (McMaster University, Canada) Antonio Bognanni (McMaster University, Canada) Imad Bou Akl (American University of Beirut, Lebanon) Mary Boulos (Michael G. DeGroote School of Medicine, McMaster University, Canada) Romina Brignardello-Petersen (McMaster University, Canada) Matthew Chan (McMaster University, Canada) Rana Charide (McMaster University, Canada) Luis Colunga-Lozano (McMaster University, Canada) Karin Dearness (St. Joseph's Healthcare Hamilton, Canada) Andrea Darzi (McMaster University, Canada) Heba Hussein (Faculty of Dentistry, Cairo University, United States) Samer Karam (McMaster University, Canada) Philipp Kolb (Michael G. DeGroote School of Medicine, McMaster University, Canada) Razan Mansour (University of Kansas Medical Center, United States) Gian Paolo Morgano (McMaster University, Canada) Rami Z. Morsi (University of Chicago Medical Center, United States) Giovanna Elsa Ute Muti Schuenemann (University of Milan, Italy) Menatalla Nadim (Faculty of Medicine, Ain Shams University, Egypt) Atefeh Noori (The Michael G. DeGroote National Pain Center, McMaster University, Canada) Binu Philip (McMaster University, Canada) Thomas Piggott (McMaster University, Canada) Yuan Qiu (University of Ottawa Heart Institute, Canada) Yetiani Benitez (McMaster University, Canada) Finn Schünemann (Asklepios Klinik Wandsbek (Hamburg), Germany) Adrienne Stevens (Public Health Agency of Canada; Centre for Immunization Programs, Canada) Karla Solo (London Health Sciences Centre, Canada) Wojtek Wiercioch (McMaster University, Canada) Reem Mustafa (University of Kansas Medical Center, United States) Robby Nieuwlaat (McMaster University, Canada)

#### Abstract:

COVID-19-related critical and acute illness are associated with an increased risk of venous thromboembolism (VTE). These evidence-based recommendations of the American Society of Hematology (ASH) are intended to support patients, clinicians, and other healthcare professionals in decisions about the use of anticoagulation for thromboprophylaxis in patients with COVID-19-related critical illness, acute illness, and those being discharged from the hospital, who do not have suspected or confirmed VTE. ASH formed a multidisciplinary panel, including three patient representatives, and applied a conflicts of interest management policy to minimize potential bias. The Michael G. DeGroote Cochrane Canada and MacGRADE Centres at McMaster University supported the guideline development process, including performing systematic evidence reviews (up to June 2023). The panel prioritized clinical questions and outcomes according to their importance for clinicians and patients. The panel used the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach to assess certainty of the evidence and make recommendations, which were subject to public comment. This is an executive summary of three updated recommendations that have been published which concludes the living phase of the quidelines. For critically ill patients with COVID-19, the panel issued conditional recommendations in favor of (a) prophylactic-intensity over therapeutic-intensity anticoagulation and (b) prophylactic-intensity over intermediate-intensity anticoagulation. For acutely ill patients with COVID-19, conditional recommendations were made in favor of (a) prophylactic-intensity over intermediate-intensity anticoagulation and (b) therapeutic-intensity over prophylactic-intensity anticoagulation. The panel also issued a conditional recommendation against the use of post-discharge extended pharmacologic thromboprophylaxis. These three conditional recommendations were made based on low or very low certainty in the evidence, underscoring the need for additional, high-quality randomized controlled trials in patients with COVID-19-related illness.

#### Conflict of interest: COI declared - see note

**COI notes:** All authors were members of the guideline panel or members of the systematic review team or both. As such, they completed a disclosure of interest form, which was reviewed by ASH and is available as Supplements 1 and 2.

#### Preprint server: No;

Author contributions and disclosures: D.M.S, E.K.T., and R.N. wrote the manuscript. All other authors contributed to critical revisions of the manuscript. All authors approved of the content. Members of the knowledge synthesis team (R.N., R.A.J., Y.A.J., L.E.C.L., K.D., A.J.D., S.G.K., G.P.M., R.Z.M., B.A.P., Y.R.B., K.S., W.W.) searched the literature, extracted data from eligible studies, analyzed the data, and prepared evidence summaries and evidence to decision tables. Panel members (D.M.S., E.K.T., H.J.S., P.A., C.B., A.C., K.D., M.T.D., D.D., D.O.G., F.A.K., A.I.L., I.N., A.P., M.R., K.M.S., D.M.S., M.S., D.R.T., K.T., R.A.M., R.N.) assessed the evidence, voted, and made judgments within the evidence to decision framework, and discussed and issued the recommendations. The methods leadership team (R.N., R.B.P., K.D., A.S., K.S., A.C., E.A.A., W.W., R.A.M., H.J.S.) developed the methods and provided guidance to the knowledge synthesis team and guideline panel. D.M.S., R.A.M., and R.N. were the co-chairs of the panel and led panel meetings.

#### Non-author contributions and disclosures: No;

Agreement to Share Publication-Related Data and Data Sharing Statement:

Clinical trial registration information (if any):

# ASH living guidelines on use of anticoagulation for thromboprophylaxis in patients with COVID-19: executive summary

Short title: ASH guidelines on anticoagulation in COVID-19

Deborah M. Siegal<sup>1\*</sup>, Eric K. Tseng<sup>2\*</sup>, Holger J. Schünemann<sup>3</sup>, Pantep Angchaisuksiri<sup>4</sup>, Adam Cuker<sup>5</sup>, Kathryn Dane<sup>6</sup>, Maria T. DeSancho<sup>7</sup>, David Diuguid<sup>8</sup>, Daniel O. Griffin<sup>9</sup>, Frederikus A. Klok<sup>10</sup>, Alfred Ian Lee<sup>11</sup>, Ignacio Neumann<sup>12</sup>, Ashok Pai<sup>13</sup>, Marc Righini<sup>14</sup>, Kristen M. Sanfilippo<sup>15</sup>, Deirdra R. Terrell<sup>16</sup>, Elie A. Akl<sup>17</sup>, Reyad Al Jabiri<sup>18</sup>, Yazan Al Jabiri<sup>19</sup>, Angela M. Barbara<sup>20</sup>, Antonio Bognanni<sup>21</sup>, Imad Bou Akl<sup>22</sup>, Mary Boulos<sup>23</sup>, Romina Brignardello-Petersen<sup>24</sup>, Matthew Chan<sup>25</sup>, Rana Charide<sup>26</sup>, Luis E. Colunga-Lozano<sup>27</sup>, Karin Dearness<sup>28</sup>, Andrea J. Darzi<sup>29</sup>, Heba Hussein<sup>30</sup>, Samer G. Karam<sup>31</sup>, Philipp Kolb<sup>32</sup>, Razan Mansour<sup>33</sup>, Gian Paolo Morgano<sup>34</sup>, Rami Z. Morsi<sup>35</sup>, Giovanna Muti- Schünemann<sup>36</sup>, Menatalla K. Nadim<sup>37</sup>, Atefeh Noori<sup>38</sup>, Binu A. Philip<sup>39</sup>, Thomas Piggott<sup>40</sup>, Yuan Qiu<sup>41</sup>, Yetiani Roldan Benitez<sup>42</sup>, Finn Schünemann<sup>43</sup>, Adrienne Stevens<sup>44</sup>, Karla Solo<sup>45</sup>, Wojtek Wiercioch<sup>46</sup>, Reem A. Mustafa<sup>47</sup>, and Robby Nieuwlaat<sup>48</sup>

#### \*Co-first authors

- 1. Department of Medicine and the Ottawa Hospital Research Institute, University of Ottawa, Ottawa, ON, Canada; dsiegal@toh.ca; ORCID ID 0000-0003-3806-3245
- 2. St. Michael's Hospital, Division of Hematology/Oncology, University of Toronto, Toronto, ON, Ontario, Canada; eric.tseng@unityhealth.to
- Departments of Health Research Methods, Evidence, and Impact and of Medicine, McMaster University, Hamliton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Institut für Evidence in Medicine, Medical Center & Faculty of Medicine, University of Freiburg, Freiburg, Germany; schuneh@mcmaster.ca, ORCID ID 0000-0003-3211-8479
- 4. Division of Hematology, Department of Medicine, Ramathibodi Hospital, Mahidol University, Bangkok, Thailand; pantep.ang@mahidol.ac.th
- 5. Department of Medicine and Department of Pathology & Laboratory Medicine; adam.cuker@pennmedicine.upenn.edu; ORCID ID 0000-0002-3595-5697
- 6. Department of Pharmacy, The Johns Hopkins Hospital, Baltimore, MD, USA; kdane2@jhmi.edu
- 7. Division of Hematology-Oncology, Department of Medicine, Weill Cornell Medicine, New York Presbyterian Hospital, New York, NY, USA; mtd2002@med.cornell.edu
- 8. College of Physicians & Surgeons of Columbia University, New York, NY, USA; dld6@cumc.columbia.edu
- Department of Medicine, Division of Infectious Diseases, Columbia University, College of Physicians and Surgeons, New York, NY, USA; Optum Tristate, Lake Success, NY, USA; danielgriffinmd@gmail.com
- 10. Department of Medicine Thrombosis and Hemostasis, Leiden University Medical Center, Leiden, the Netherlands; F.A.Klok@lumc.nl
- 11. Section of Hematology, Yale School of Medicine, New Haven, CT, USA; alfred.lee@yale.edu
- 12. School of Medicine, Universidad San Sebastian, Santiago, Chile; ignacio.neumann@gmail.com

- 13. Division of Hematology & Oncology, Kaiser Permanente, Oakland/Richmond, CA, USA; ashok.p.pai@kp.org
- 14. Division of Angiology and Hemostasis, Geneva University Hospitals and Faculty of Medicine, Geneva, Switzerland; marc.righini@hcuge.ch
- 15. Washington University School of Medicine St. Louis, St. Louis, MO, USA; ksanfilippo@wustl.edu; ORCID ID 0000-0002-0433-7845
- 16. Department of Biostatistics and Epidemiology, Hudson College of Public Health, University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA; Dee-Terrell@ouhsc.edu
- 17. Department of Internal Medicine, American University of Beirut, Beirut, Lebanon; ea32@aub.edu.lb
- 18. University of Jordan, Amman, Jordan; reyad.naif@gmail.com
- 19. Washington University in St. Louis, Saint Louis, MO; yazan.aljabirii@gmail.com
- 20. The Canadian Agency for Drugs and Technologies in Health, Ottawa, ON, Canada; barbara@mcmaster.ca
- 21. Department of Health Research Methods, Evidence, and Impact, Michael G. DeGroote Cochrane Canada and GRADE Centres, McMaster University, Hamilton, ON, Canada; bognana@mcmaster.ca
- 22. American University of Beirut's School of Medicine, Beirut, Lebanon; ib08@aub.edu.lb
- 23. University of Toronto, Toronto, ON, Canada
- 24. Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; brignarr@mcmaster.ca
- 25. McMaster University Medical Center, Hamilton, Ontario, Canada; mattchan353@gmail.com
- 26. McMaster University, Department of Health Research Methods, Evidence, and Impact, Hamilton, Ontario, Canada; charider@mcmaster.ca
- 27. Department of Clinical Medicine, Health Science Center, Universidad de Guadalajara, Guadalajara, Jalisco, Mexico; dr.colunga.lozano@gmail.com; ORCID ID 0000-0001-7737-4914
- 28. Library Services, St. Joseph's Healthcare Hamilton, Hamilton, ON, Canada; kdearnes@stjoes.ca, ORCID ID 0000-0002-6854-0156
- 29. Department of Health Research Methods, Evidence, and Impact; Michael G. DeGroote Cochrane Canada & McMaster GRADE centres, McMaster University, Hamilton, ON, Canada; darzia@mcmaster.ca; ORCID ID 0000-0002-2498-1697
- 30. Department of Oral Medicine, Oral Diagnosis, and Periodontology, Faculty of Dentistry, Cairo University, Cairo, Egypt
- 31. McMaster University, Hamilton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; karams1@mcmaster.ca
- 32. McMaster University, Hamilton, ON, Canada; philipp.kolb@medportal.ca
- University of Kansas Medical Center, University of Kansas, Kansis City, KS; razanamansour@gmail.com
- 34. McMaster University, Hamilton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; morganog@mcmaster.ca
- 35. Department of Neurology, University of Chicago, Chicago, IL, USA; Rami.Morsi@uchospitals.edu; ORCID ID 0000-0003-2131-3711
- 36. Department of Emergency Medicine, University of Milan, Milan, Italy; Giovanna.muti@unimi.it
- 37. Department of Clinical Pathology, Faculty of Medicine, Ain Shams University, Cairo, Egypt; menna.nadeem1@gmail.com
- 38. Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; and Hand Program, Division of Plastic, Reconstructive and Aesthetic Surgery, University Health Network, Toronto Western Hospital, University of Toronto, Toronto, Canada

- 39. Department of Health Research Methods, Evidence, and Impact, Michael G. DeGroote Cochrane Canada and GRADE Centres, McMaster University, Hamilton, ON, Canada; abrahab@mcmaster.ca
- 40. Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; and Department of Family Medicine, Queens University, Kingston, ON, Canada
- 41. Department of Cardiac Surgery, University of Ottawa Heart Institute, Ottawa, ON, Canada
- 42. Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; yetiani@gmail.com
- 43. Asklepios Klinik Wandsbek, Hamburg, Germany; finnschuenemann@googlemail.com
- 44. McMaster University, Hamilton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; adrienne.stevens@mcmaster.ca; ORCID ID 0000-0002-6257-4806
- 45. McMaster University, Hamilton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; solok@mcmaster.ca
- 46. McMaster University, Hamilton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; wierciww@mcmaster.ca; ORCID ID 0000-0001-6576-1650
- 47. Department of Internal Medicine, Division of Nephrology, University of Kansas Medical Center, Kansas City, KS, USA; rmustafa@KUMC.edu, ORCID ID 0000-0002-2091-0875
- McMaster University, Hamilton, ON, Canada; Michael G. DeGroote Cochrane Canada and McGRADE Centres; Department of Health Research Methods, Evidence, and Impact, McMaster University, Hamilton, ON, Canada; nieuwlr@mcmaster.ca

Corresponding author: Deborah Siegal, MD, MSc, FRCPC Department of Medicine, University of Ottawa Ottawa Hospital Research Institute 501 Smyth Rd, Box 201A Ottawa, ON, Canada K1H 8L6 Email: dsiegal@toh.ca Telephone: (613)737-8899 ext. 78804

Text word count: 6567 Abstract word count: 265 Number of tables and figures: 3 tables Number of references: 65 Number of supplements: 4

# Abstract

COVID-19-related critical and acute illness are associated with an increased risk of venous thromboembolism (VTE). These evidence-based recommendations of the American Society of Hematology (ASH) are intended to support patients, clinicians, and other healthcare professionals in decisions about the use of anticoagulation for thromboprophylaxis in patients with COVID-19-related critical illness, acute illness, and those being discharged from the hospital, who do not have suspected or confirmed VTE. ASH formed a multidisciplinary panel, including three patient representatives, and applied a conflicts of interest management policy to minimize potential bias. The Michael G. DeGroote Cochrane Canada and MacGRADE Centres at McMaster University supported the guideline development process, including performing systematic evidence reviews (up to June 2023). The panel prioritized clinical questions and outcomes according to their importance for clinicians and patients. The panel used the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach to assess certainty of the evidence and make recommendations, which were subject to public comment. This is an executive summary of three updated recommendations that have been published which concludes the living phase of the guidelines. For critically ill patients with COVID-19, the panel issued conditional recommendations in favor of (a) prophylactic-intensity over therapeutic-intensity anticoagulation and (b) prophylactic-intensity over intermediate-intensity anticoagulation. For acutely ill patients with COVID-19, conditional recommendations were made in favor of (a) prophylactic-intensity over intermediate-intensity anticoagulation and (b) therapeutic-intensity over prophylactic-intensity anticoagulation. The panel also issued a conditional recommendation against the use of post-discharge extended pharmacologic thromboprophylaxis. These three conditional recommendations were made based on low or very low certainty in the evidence, underscoring the need for additional, high-quality randomized controlled trials in patients with COVID-19-related illness.

## Keywords

COVID-19; anticoagulation; practice guidelines; hospital discharge; thromboprophylaxis

# **Summary of recommendations**

#### Background

Venous thromboembolism is an important complication that occurs in acutely and critically ill patients with COVID-19 despite the use of standard thromboprophylaxis regimens.<sup>1</sup> Thrombosis of the microvascular circulation may also contribute to other complications of COVID-19, including respiratory failure.<sup>2,3</sup> Meanwhile, higher-intensity anticoagulation is associated with an increase in bleeding risk among hospitalized patients who have COVID-19.<sup>4</sup> Therefore, there has been broad interest in establishing how anticoagulant regimens may improve clinical outcomes both during hospitalization and following hospital discharge.

These guidelines address the use of anticoagulation for thromboprophylaxis as follows: (1) higher intensity anticoagulation (intermediate- or therapeutic-intensity) compared to standard prophylactic-intensity anticoagulation in critically ill patients with COVID-19, (2) higher intensity anticoagulation (intermediate- or therapeutic-intensity) compared to standard prophylactic-intensity anticoagulation in acutely ill patients with COVID-19, and (3) prophylactic-intensity anticoagulation compared to no anticoagulation in patients discharged after hospitalization for COVID-19. These guidelines are based on systematic reviews of evidence conducted under the direction of the Michael G. DeGroote Cochrane Canada and MacGRADE Centres at McMaster University with international collaborators. The panel followed best practice for guideline development recommended by the Institute of Medicine and the Guidelines International Network (GIN).<sup>5-7</sup> The panel used the GRADE approach to assess the certainty of the evidence and formulate recommendations<sup>8-14</sup>.

**Recommendation 1a.** The American Society of Hematology (ASH) guideline panel *suggests* using prophylactic-intensity over intermediate-intensity anticoagulation in patients with COVID-19-related critical illness who do not have suspected or confirmed venous thromboembolism (VTE) or another indication for anticoagulation (conditional recommendation based on low certainty in the evidence about effects  $\bigoplus \bigoplus \bigcirc$ ).

**Recommendation 1b.** The ASH guideline panel *suggests* using prophylactic-intensity over therapeutic-intensity anticoagulation for patients with COVID-19-related critical illness who

do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus$ 

#### Remarks:

- Patients with COVID-19-related critical illness are defined as those suffering from an immediately life-threatening condition who would typically be admitted to an intensive care unit (ICU) due to COVID-19 infection. Examples include patients requiring hemodynamic support, ventilatory support, and renal replacement therapy.
- An individualized assessment of the patient's risk of thrombosis and bleeding is
  important when deciding on anticoagulation intensity. Risk assessment models
  (RAMs) to estimate thrombotic risk have been validated in hospitalized patients with
  COVID-19 (critically or non-critically ill), with modest prognostic performance. No
  RAMs for bleeding have been validated for patients with COVID-19. The panel
  acknowledges that higher-intensity anticoagulation may be preferred for patients
  judged to be at low bleeding risk and high thrombotic risk.
- At present, there is no direct high-certainty evidence comparing different types of anticoagulants. The selection of a specific agent (e.g., low molecular weight heparin [LMWH], unfractionated heparin [UFH]) may be based on availability, resources required, familiarity, and the aim of minimizing the use of personal protective equipment or exposure to staff to COVID-19-infected patients as well as patientspecific factors (e.g., renal function, history of heparin-induced thrombocytopenia, bleeding risk). LMWH and UFH were used in the identified studies and may be preferred because of a preponderance of evidence with these agents. There are no studies of intermediate- or therapeutic-intensity fondaparinux, argatroban, or bivalirudin in this population.

 These recommendations do not apply to patients who require anticoagulation to prevent thrombosis of extracorporeal circuits such as those on extracorporeal membrane oxygenation (ECMO) or continuous renal replacement therapy (CRRT).

**Recommendation 2a.** The ASH guideline panel *suggests* using prophylactic-intensity over intermediate-intensity anticoagulation for patients with COVID-19-related acute illness who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus$   $\bigcirc \bigcirc \bigcirc$ ).

**Recommendation 2b.** The ASH guideline panel *suggests* using therapeutic-intensity over prophylactic-intensity anticoagulation for patients with COVID-19-related acute illness who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus$   $\bigcirc$  $\bigcirc$ ).

#### Remarks:

- Patients with COVID-19-related acute illness are defined as those with clinical features that would typically result in admission to an inpatient medical ward without requirement for intensive clinical support. Examples include patients with dyspnea or mild-to-moderate hypoxia.
- An individualized assessment of the patient's risk of thrombosis and bleeding is important when deciding on anticoagulation intensity. RAMs to estimate thrombotic risk have been validated in hospitalized patients with COVID-19 (critically or noncritically ill), with modest prognostic performance. No RAMs for bleeding have been validated for patients with COVID-19. The panel acknowledges that lower-intensity anticoagulation may be preferred for patients judged to be at high bleeding risk and low risk of thrombosis.
- At present, there is no direct high-certainty evidence comparing different types of anticoagulants in patients with COVID-19. LMWH or UFH may be preferred because

of a preponderance of evidence with these agents. There are no studies of therapeutic-intensity fondaparinux, argatroban, or bivalirudin in this population.

**Recommendation 3.** The ASH guideline panel *suggests* against using post-discharge outpatient anticoagulant thromboprophylaxis in patients with COVID-19 who are being discharged from hospital and who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus$  ().

#### Remarks:

- An individualized assessment of the patient's risk of thrombosis and bleeding and shared decision-making are important when deciding on whether to use postdischarge thromboprophylaxis.
- The panel acknowledged that post-discharge thromboprophylaxis may be reasonable in patients judged to be at high thrombotic risk and low bleeding risk.

#### Values and preferences

Please refer to the full recommendation reports below and the online Evidence-to-Decision frameworks for considerations regarding values and preferences.

#### Explanations and other considerations

Please refer to the full recommendation reports below and the online Evidence-to-Decision frameworks for explanations and other considerations.

## Interpretation of strong and conditional recommendations

The strength of a recommendation is expressed as either strong ("the guideline panel recommends..."), or conditional ("the guideline panel suggests...") and has the following interpretation<sup>15</sup>:

#### Strong recommendation

- For patients: Most individuals in this situation would want the recommended course of action, and only a small proportion would not.
- For clinicians: Most individuals should follow the recommended course of action. Formal decision aids are not likely to be needed to help individual patients make decisions consistent with their values and preferences.
- For policy makers: The recommendation can be adopted as policy in most situations. Adherence to this recommendation according to the guideline could be used as a quality criterion or performance indicator.
- For researchers: The recommendation is supported by credible research or other convincing
  judgments that make additional research unlikely to alter the recommendation. On occasion, a
  strong recommendation is based on low or very low certainty in the evidence. In such instances,
  further research may provide important information that alters the recommendations.

#### Conditional recommendation

- For patients: The majority of individuals in this situation would want the suggested course of action, but many would not. Decision aids may be useful in helping patients to make decisions consistent with their individual risks, values, and preferences.
- For clinicians: Recognize that different choices will be appropriate for individual patients and that you must help each patient arrive at a management decision consistent with their values and preferences. Decision aids may be useful in helping individuals to make decisions consistent with their individual risks, values, and preferences.
- For policy makers: Policy-making will require substantial debate and involvement of various stakeholders. Performance measures about the suggested course of action should focus on whether an appropriate decision-making process is duly documented.
- For researchers: This recommendation is likely to be strengthened (for future updates or adaptation) by additional research. An evaluation of the conditions and criteria (and the related judgments, research evidence, and additional considerations) that determined the conditional (rather than strong) recommendation will help identify possible research gaps.

# Introduction

## Aims and specific objectives of these guidelines

The ASH guidelines on thromboprophylaxis in patients with COVID-19 were created as living guidelines that were updated through living systematic reviews, as new evidence emerged throughout the course of the global pandemic. More background on this methodology and approach can be found in the original ASH guideline on thromboprophylaxis in patients with COVID-19.<sup>15</sup>

Using living guideline methods, these recommendations were initially published individually as new evidence was published. The first guideline (Recommendations 1 and 2, regarding critically ill and acutely ill patients with COVID-19) was published in February 2021. There have been four subsequent updates including the addition of Recommendation 3 regarding post-discharge thromboprophylaxis<sup>16-19</sup>.

The present manuscript is an executive summary of all updated ASH guideline panel recommendations (summarized in Table 1) representing the conclusion of the living guideline phase. All recommendations and updates to these living guidelines are also accessible at the ASH COVID-19 anticoagulation webpage<sup>20</sup>.

# Description of the health problem

The COVID-19 pandemic has had a significant public health impact with substantial global morbidity and mortality.<sup>21</sup> Patients who develop COVID-19-related acute or critical illness may develop hypercoagulability, thrombosis, and coagulopathy which is marked by elevated fibrinogen, D-dimer concentrations, and inflammatory markers.<sup>22,23</sup> Vascular endothelial dysfunction (endotheliopathy) may also occur which can contribute to systemic hypercoagulability and microvascular thrombosis.<sup>24</sup>

Thrombosis is an important complication of patients hospitalized with COVID-19-related acute or critical illness. Early cohort studies in predominantly unvaccinated patients reported VTE in 7.9% and 22.7% of patients in these clinical contexts, respectively despite the use of standard pharmacological thromboprophylaxis.<sup>1</sup>

Additionally, based on the high observed incidence of VTE during hospitalization for COVID-19, there is concern that COVID-19 patients may have a higher risk of VTE after discharge than non-COVID-19

patients. However, estimates of post-discharge VTE in COVID-19 patients generally range from 0.5% to 1.5%, comparable to the incidence of VTE after hospitalization for non-COVID-19 illnesses.<sup>25-27</sup> Nevertheless, there has been ongoing interest in establishing whether extended thromboprophylaxis is beneficial in these patients. There are no risk assessment models (RAMs) that have been specifically derived and prospectively validated in COVID-19 patients, although non-COVID RAMs such as IMPROVE-DD have been externally validated in retrospective cohorts of hospitalized COVID-19 patients, and the COVID-TE score was derived specifically in COVID-19 patients with concomitant malignancy.<sup>28-30</sup>

The present manuscript is an executive summary encompassing three updated recommendations on the use of anticoagulant therapy for patients admitted with COVID-19-related critical illness, acute illness, and patients discharged from hospital that concludes the living phase of the guidelines.

## **Description of the target populations**

The target populations in this guideline include patients with COVID-19 with critical illness, acute illness, and those discharged from acute care hospitals. These groups are described in Table 2.

# **Methods**

This updated executive summary includes three recommendations which were developed as part of ASH's living guidelines effort regarding the use of anticoagulant thromboprophylaxis in hospitalized patients with COVID-19. These recommendations have been previously published separately as standalone recommendations or updates.<sup>16–19</sup> The living phase (i.e., continuous review and updating) is concluded. Going forward, ASH will maintain these guidelines through regular review and scheduled revision.. For all recommendations, we followed the same methods as reported in publications to date, and important methodological aspects and updates are highlighted below.

The initial and updated recommendations were created as follows:

Recommendation	Population	Anticoagulation	First version <sup>*</sup>	Update <sup>*</sup>	Update for
		intensities being			Executive
		compared			Summary <sup>*</sup>

1a	Critically ill COVID-19 patients	Prophylactic vs Intermediate	Oct-2020	Jun-2021	Apr-2024
1b	Critically ill COVID-19 patients	Prophylactic vs Therapeutic	Oct-2020	Apr-2022	Apr-2024
2a	Acutely ill COVID-19 patients	Prophylactic vs Intermediate	Oct-2020	Jul-2022 <sup>#</sup>	Apr-2024
2b	Acutely ill COVID-19 patients	Prophylactic vs Therapeutic	Oct-2020	Mar-2022	Apr-2024
3	COVID-19 patients being discharged	Prophylactic vs none	Aug-2021	Aug-2022	Apr-2024

\* Dates on which the recommendations were approved by the ASH Guideline Oversight Subcommittee and the ASH Committee on Quality.

# Date when public commenting was closed on the ASH website.

This executive summary includes final versions of all recommendations as approved by the ASH Guideline Oversight Subcommittee and the ASH Committee on Quality in April 2024. For the executive summary we have applied the following important aspects:

 Guideline funding and management of conflicts of interest: Supplement 1 provides updated "Participant Information Forms" for all panel members, detailing financial and non-financial interests, as well as the ASH conflict of interest policies agreed to by each individual. Supplement 2 provides the updated complete Participant Information Forms of researchers on the systematic review team who contributed to these guidelines.

- Evidence review and development of recommendation: New EtD frameworks were created for all recommendations including new evidence and considerations. The systematic review to identify comparative anticoagulation studies for the entire guideline was updated until June 2023. During the project, the initial guideline's literature search strategy (Supplement 3) was modified to add search terms for antiplatelet agents for the guideline question on post-discharge anticoagulation, and the protocol (Supplement 4) was modified to focus on inclusion of only randomized controlled trials. The systematic review to identify baseline risk studies for important outcomes for all guideline questions was updated until June 2023 and the methods remained the same throughout the project (search strategy and protocol previously published<sup>15</sup>)
- Criteria to update living systematic reviews and recommendations: Due to the rapid emergence
  of a wealth of research studies related to this topic, the systematic reviews were periodically
  updated and recommendations were reconsidered if new evidence could potentially lead to
  important changes in baseline risk estimates, intervention effect estimates, certainty of the
  evidence, or to ensure face validity especially to include important trials.
- Decision thresholds: To support judgements about whether the magnitude of an effect estimate was trivial, small, moderate, or large, as well as for determining imprecision of effect estimates, we used decision thresholds for all outcomes considered in the final reported recommendations in the executive summary. Thresholds were calculated using the outcome-specific utility value and results from a decision threshold survey that included the members of the panel. The decision threshold values that were used for each recommendation are reported in the footnotes of the online Evidence Profiles.
- Document review: Draft recommendations were reviewed by all members of the panel, and made available online from December 1, 2023, to December 22, 2023 for external review by stakeholders including allied organizations, other medical professionals, patients, and the public. One individual submitted a response that did not require changes to the recommendations. In April 2024, the ASH Guideline Oversight Subcommittee and the ASH Committee on Quality approved that the defined guideline development process was followed, and in May 2024, the officers of the ASH Executive Committee approved submission of the executive summary manuscript for publication under the imprimatur of ASH.

For more information on how these guidelines should be used by patients, clinicians, policy makers, and researchers, we refer readers to the description in the initial guideline publication from February 2021,<sup>15</sup> as well as the user guide to ASH clinical practice guidelines.<sup>31</sup>

# Recommendations

## Patients with COVID-19 related critical illness

Question: Should direct oral anticoagulants, low molecular weight heparin, unfractionated heparin, fondaparinux, argatroban, or bivalirudin at intermediateintensity or therapeutic-intensity vs prophylactic-intensity be used for patients with COVID-19 related critical illness who do not have suspected or confirmed VTE or another indication for anticoagulation?

**Recommendation 1a.** The American Society of Hematology (ASH) guideline panel *suggests* using prophylactic-intensity over intermediate-intensity anticoagulation in patients with COVID-19-related critical illness who do not have suspected or confirmed venous thromboembolism (VTE) or another indication for anticoagulation (conditional recommendation based on low certainty in the evidence about effects  $\bigoplus \bigoplus \bigcirc \bigcirc$ ).

Recommendation 1b. The ASH guideline panel *suggests* using prophylactic-intensity over therapeutic-intensity anticoagulation for patients with COVID-19-related critical illness who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus$   $\bigcirc \bigcirc \bigcirc$ ).

Remarks:

 Patients with COVID-19-related critical illness are defined as those suffering from an immediately life-threatening condition due to COVID-19 infection who would typically be admitted to an intensive care unit (ICU). Examples include patients requiring hemodynamic support, ventilatory support, and renal replacement therapy.

- An individualized assessment of the patient's risk of thrombosis and bleeding is important when deciding on anticoagulation intensity. Risk assessment models (RAMs) to estimate thrombotic risk have been validated in hospitalized patients with COVID-19 (critically or non-critically ill), with modest prognostic performance. No RAMs for bleeding have been validated for patients with COVID-19. The panel acknowledges that higher-intensity anticoagulation may be preferred for patients judged to be at low bleeding risk and high thrombotic risk.
- At present, there is no direct high-certainty evidence comparing different types of anticoagulants. The selection of a specific agent (e.g., low molecular weight heparin [LMWH], unfractionated heparin [UFH]) may be based on availability, resources required, familiarity, and the aim of minimizing the use of personal protective equipment or exposure to staff to COVID-19-infected patients as well as patientspecific factors (e.g., renal function, history of heparin-induced thrombocytopenia, bleeding risk). LMWH and UFH were used in the identified studies. LMWH or UFH may be preferred because of a preponderance of evidence with these agents. There are no studies of intermediate- or therapeutic-intensity fondaparinux, argatroban, or bivalirudin in this population.
- These recommendations do not apply to patients who require anticoagulation to prevent thrombosis of extracorporeal circuits such as those on extracorporeal membrane oxygenation (ECMO) or continuous renal replacement therapy (CRRT).

### Summary of the evidence

The now-expired, first iteration of Recommendation 1 published in February 2021 compared therapeutic-intensity or intermediate-intensity with prophylactic-intensity anticoagulation in patients with COVID-19 related critical illness. However, with the publication of new evidence this recommendation was split into two recommendations comparing intermediate-intensity versus prophylactic-intensity anticoagulation (Recommendation 1a, first published in October 2021) and a

separate recommendation comparing therapeutic-intensity versus prophylactic-intensity anticoagulation (Recommendation 1b, first published in September 2022).

#### **Recommendation 1a**

The EtD framework for Recommendation 1a was updated as of November 2023. Three randomized controlled trials were identified that provided evidence related to this question regarding the effects of intermediate-intensity compared with prophylactic-intensity anticoagulation on multiple critical outcomes among all-cause mortality, pulmonary embolism (PE) deep venous thrombosis (DVT), ischemic stroke, major bleeding, intracranial hemorrhage (ICH), multiple organ failure, ST-elevation myocardial infarction (STEMI), limb amputation, IMV, length of hospital admission and length of ICU admission.<sup>32-34</sup> Two of the trial groups provided unpublished data on request for selected outcomes. The overall certainty of the evidence of effects was very low. Depending on the outcome, this was primarily due to extremely serious imprecision and/or serious risk of bias (see Evidence Profile and EtD framework online at: https://guidelines.ash.gradepro.org/profile/blz3F60WNWs).

Based on the panel's thresholds for effect sizes, intermediate-intensity anticoagulation may reduce allcause mortality (OR: 0.92, 95% CI: 0.62 to 1.37; corresponding to 16 fewer [from 85 fewer to 67 more] deaths per 1,000 patients), and may reduce PE (OR: 0.55, 95% CI: 0.12 to 2.62; corresponding to 34 fewer [from 68 fewer to 103 more] PEs per 1,000 patients), but the evidence was very uncertain. Intermediate-intensity anticoagulation likely results in little to no effect on length of hospital admission (mean difference: 0.39 days fewer [from 1.82 days fewer to 1.04 days more]), and may not reduce length of ICU admission (mean difference: 0.09 days fewer [from 1.83 days fewer to 1.65 days more]). Intermediate-intensity anticoagulation may have little to no effect on DVT (OR: 0.93, 95% CI: 0.23 to 3.80; corresponding to 3 fewer [from 31 fewer to 99 more] DVTs per 1,000 patients), but the evidence was very uncertain. In terms of potential harms, intermediate-intensity anticoagulation may result in little to no difference in major bleeding (OR: 1.50, 95% CI: 0.63 to 3.58; corresponding to 16 more [from 12 fewer to 78 more] major bleeding events per 1,000 patients), but the evidence was very uncertain. Intermediate-intensity anticoagulation may have little to no effect on all other critical outcomes, but the evidence was very uncertain. No effects could be determined for multiple organ failure and limb amputation.

#### **Recommendation 1b**

The EtD framework for Recommendation 1b was last updated as of September 2023. Seven randomized controlled trials were identified that provided evidence related to this question regarding the effects of therapeutic-intensity compared with prophylactic-intensity anticoagulation on the same multiple critical outcomes.<sup>33,35-40</sup> Unpublished data were provided on request for selected outcomes by two trial groups. The overall certainty of the evidence of effects was very low. Depending on the outcome, this was primarily due to very serious imprecision, serious risk of bias and/or serious indirectness (see Evidence Profile and EtD framework online at: https://guidelines.ash.gradepro.org/profile/IHYtm7MSFLE).

Based on the panel's thresholds for effect sizes, therapeutic-intensity anticoagulation may reduce allcause mortality (OR: 0.90, 95% CI: 0.70 to 1.17; corresponding to 21 fewer [from 66 fewer to 33 more] deaths per 1,000 patients), PE (OR: 0.40, 95% CI: 0.26 to 0.61; corresponding to 45 fewer [from 56 fewer to 29 fewer] PEs per 1,000 patients), and invasive mechanical ventilation (OR: 0.82, 95% CI: 0.57 to 1.20; corresponding to 33 fewer [from 84 fewer to 34 more] IMV per 1,000 patients), but the evidence was very uncertain. Therapeutic-intensity anticoagulation may result in little to no difference in ischemic stroke (OR: 0.75, 95% CI: 0.32 to 1.77; corresponding to 5 fewer [from 14 fewer to 15 more] ischemic strokes per 1,000 patients) and STEMI (OR: 0.83, 95% CI: 0.33 to 2.10; corresponding to 2 fewer [from 6 fewer to 10 more] STEMI per 1,000 patients). Therapeutic-intensity anticoagulation may have little to no effect on DVT (OR: 0.73, 95% CI: 0.42 to 1.24; corresponding to 11 fewer [from 23 fewer to 9 more] DVTs per 1,000 patients), but the evidence was very uncertain. In terms of potential harms, therapeuticintensity anticoagulation may increase major bleeding (OR: 1.78, 95% CI: 1.00 to 3.18; corresponding to 25 more [from 0 to 67 more] major bleedings per 1,000 patients), may result in little to no difference in length of hospital admission (mean difference: 1.32 days more [from 0.02 days more to 2.61 days more]), but the evidence for the latter was very uncertain. Therapeutic-intensity anticoagulation may have little to no effect on all other critical outcomes, but the evidence was very uncertain. No effects could be determined for ICH.

#### **Conclusions for this recommendation**

Regarding intermediate-intensity anticoagulation, although the panel judged the overall certainty of evidence to be very low for both desirable and undesirable effects, the panel judged that the trivial-tosmall benefits do not outweigh the trivial harms of intermediate-intensity anticoagulation. Regarding therapeutic-intensity anticoagulation, although the panel judged the overall certainty of evidence to be very low for both desirable and undesirable effects, the panel judged that the small-to-moderate harms would outweigh the small benefits of therapeutic-intensity anticoagulation. Other factors considered in the EtD framework did not importantly affect this assessment for the recommendations. The panel therefore suggested prophylactic-intensity rather than intermediate-intensity and therapeutic-intensity anticoagulation in patients with COVID-19-related critical illness, as utilized in critically ill non-COVID-19 patients.<sup>41-45</sup> This guideline did not address the use of therapeutic- versus intermediate-intensity anticoagulation in patients with COVID-19 related critical illness as this PICO question was not prioritized by the panel.

The panel noted for both Recommendations 1a and 1b that an individualized decision is important for each patient based on an assessment of thrombosis and bleeding risk. Dose adjustment of prophylacticintensity anticoagulation for extremes of body weight or renal impairment may also be considered.<sup>46-50</sup> This recommendation does not apply to thrombotic complications related to extracorporeal circuits. While high rates of circuit-related thrombosis during extracorporeal membrane oxygenation (ECMO) and continuous renal replacement therapy (CRRT) have been reported in patients with COVID-19, this outcome was not prioritized as critical for this question.<sup>51</sup>

## Patients with COVID-19 related acute illness

Question: Should direct oral anticoagulants, low molecular weight heparin, unfractionated heparin, fondaparinux, argatroban, or bivalirudin at intermediateintensity or therapeutic-intensity vs prophylactic-intensity be used in patients with COVID-19-related acute illness who do not have suspected or confirmed VTE or another indication for anticoagulation?

**Recommendation 2a.** The ASH guideline panel *suggests* using prophylactic-intensity over intermediate-intensity anticoagulation for patients with COVID-19-related acute illness who do not have suspected or confirmed VTE or another indication for anticoagulation

(conditional recommendation based on very low certainty in the evidence about effects  $\oplus$ 

**Recommendation 2b.** The ASH guideline panel *suggests* using therapeutic-intensity over prophylactic-intensity anticoagulation for patients with COVID-19-related acute illness who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus$ 

#### Remarks:

- Patients with COVID-19-related acute illness are defined as those with clinical features that would typically result in admission to an inpatient medical ward without requirement for intensive clinical support. Examples include patients with dyspnea or mild-to-moderate hypoxia.
- An individualized assessment of the patient's risk of thrombosis and bleeding is important when deciding on anticoagulation intensity. RAMs to estimate thrombotic risk have been validated in hospitalized patients with COVID-19 (critically or noncritically ill), with modest prognostic performance. No RAMs for bleeding have been validated for patients with COVID-19. The panel acknowledges that lower-intensity anticoagulation may be preferred for patients judged to be at high bleeding risk and low risk of thrombosis.
- At present, there is no direct high-certainty evidence comparing different types of anticoagulants in patients with COVID-19. LMWH or UFH may be preferred because of a preponderance of evidence with these agents. There are no studies of therapeutic-intensity fondaparinux, argatroban, or bivalirudin in this population.

## Summary of the evidence

The now-expired, first iteration of Recommendation 2 published in February 2021 compared therapeutic-intensity or intermediate-intensity with prophylactic-intensity anticoagulation in patients with COVID-19 related acute illness. However, with the publication of new evidence this recommendation was split into two recommendations comparing intermediate-intensity versus prophylactic-intensity anticoagulation (Recommendation 2a) and a separate recommendation comparing therapeutic-intensity versus prophylactic-intensity anticoagulation (Recommendation 2b, first published in September 2022).

#### **Recommendation 2a**

The EtD framework for Recommendation 2a was updated as of November 2023. Three randomized controlled trials were identified that provided evidence related to this question regarding the effects of intermediate-intensity compared with prophylactic-intensity anticoagulation on multiple critical outcomes among all-cause mortality, PE, DVT, ischemic stroke, major bleeding, ICH, multiple organ failure, STEMI, limb amputation, IMV, and ICU admission.<sup>32,52,53</sup> One of the trial groups provided unpublished data on request for selected outcomes. The overall certainty of the evidence of effects was very low. This was primarily due to extremely serious imprecision and for some outcomes risk of bias (see Evidence Profile and EtD framework online at:

https://guidelines.ash.gradepro.org/profile/cZ63B6hzUMI).

Based on the panel's thresholds for effect size, intermediate-intensity anticoagulation may increase allcause mortality (OR: 1.49, 95% CI: 0.82 to 2.72; corresponding to 41 more [from 16 fewer to 129 more] deaths per 1,000 patients) and multiple organ failure (OR: 1.53, 95% CI: 0.25 to 9.40; corresponding to 24 more [from 36 fewer to 277 more] deaths per 1,000 patients) but the evidence is very uncertain. Intermediate-intensity anticoagulation may have little to no effect on all other critical outcomes, including major bleeding, but the evidence was very uncertain. No effects could be determined for DVT, ICH, and limb amputation.

#### **Recommendation 2b**

The EtD framework for Recommendation 2b was updated as of September 2023. Nine randomized controlled trials were identified that provided evidence related to this question regarding the effects of

therapeutic-intensity compared with prophylactic-intensity anticoagulation on the same multiple critical outcomes.<sup>35,52,54-60</sup> Three of the trial groups provided unpublished data on request for selected outcomes. The overall certainty of the evidence of effects was low. This was primarily due to imprecision and risk of bias (see Evidence Profile and EtD framework online at:

https://guidelines.ash.gradepro.org/profile/noIMdHDZo6Y).

Based on the panel's thresholds for effect sizes, therapeutic-intensity anticoagulation may reduce allcause mortality (OR: 0.80, 95% CI: 0.55 to 1.16; corresponding to 26 fewer [from 41 fewer to 16 more] deaths per 1,000 patients) and probably results in little difference (low absolute risk reduction) in PE (OR: 0.53, 95% CI: 0.33 to 0.83; corresponding to 12 fewer [from 17 fewer to 4 fewer] PEs per 1,000 patients), DVT (OR: 0.58, 95% CI: 0.30 to 1.08; corresponding to 3 fewer [from 6 fewer to 1 more] DVTs per 1,000 patients), and IMV (OR: 0.76, 95% CI: 0.59 to 0.96; corresponding to 15 fewer [from 26 fewer to 3 fewer] IMV per 1,000 patients). Therapeutic-intensity anticoagulation may not reduce ICU hospitalization and STEMI and may have little to no effect on ischemic stroke, multiple organ failure, and limb amputation, but the evidence was very uncertain.

In terms of potential harms, therapeutic-intensity anticoagulation probably results in little difference in major bleeding (OR: 1.92, 95% CI: 1.10 to 3.36; corresponding to 1 more [from 12 more to 29 more] major bleedings per 1,000 patients) and may have little to no effect on ICH (OR: 2.12, 95% CI: 0.22 to 20.37; corresponding to 1 more [from 1 fewer to 19 more] ICH per 1,000 patients), although the evidence was very uncertain for the latter.

## **Conclusions for this recommendation**

Regarding Recommendation 2a, the panel judged that the balance of effects probably favors the comparison (prophylactic-intensity anticoagulation) based on the trivial desirable effects, trivial undesirable effects, possibly important uncertainty or variability in how much people value the outcomes, and the overall very low certainty of the available data. Other factors considered in the EtD framework did not importantly affect this assessment for the recommendations.

Regarding Recommendation 2b, the panel judged that the balance of effects probably favors the intervention (therapeutic-intensity anticoagulation) based on the small desirable effects, owing to additive trivial effects on multiple independent outcomes, trivial undesirable effects, possibly important uncertainty or variability in how much people value the outcomes, and the overall low certainty of the

available data. Other factors considered in the EtD framework did not importantly affect this assessment for the recommendations.

The panel noted for both Recommendations 2a and 2b that an individualized decision is important for each patient based on an assessment of thrombosis and bleeding risk. Dose adjustment of prophylacticintensity anticoagulation for extremes of body weight or renal impairment may also be considered.<sup>46-50</sup> This guideline did not address the use of therapeutic- versus intermediate-intensity anticoagulation in patients with COVID-19 related acute illness as this PICO question was not prioritized by the panel.

# Patients being discharged from hospital after COVID-19

Question: Should prophylactic-intensity direct oral anticoagulants, low molecular weight heparin, unfractionated heparin, or fondaparinux vs. no anticoagulation be used for post-discharge outpatient thromboprophylaxis in patients with COVID-19 who are being discharged from the hospital and who do not have suspected or confirmed VTE or another indication for anticoagulation?

**Recommendation 3.** The ASH guideline panel *suggests* against using post-discharge outpatient anticoagulant thromboprophylaxis in patients with COVID-19 who are being discharged from the hospital and who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects  $\bigoplus \bigcirc \bigcirc \bigcirc$ ).

#### Remarks:

- An individualized assessment of the patient's risk of thrombosis and bleeding and shared decision-making are important when deciding on whether to use postdischarge thromboprophylaxis.
- The panel acknowledged that post-discharge thromboprophylaxis may be reasonable in patients judged to be at high thrombotic risk and low bleeding risk.

## Summary of the evidence

The now-expired, first iteration of Recommendation 3 published in January 2022 compared postdischarge outpatient prophylactic-intensity anticoagulation with no anticoagulation in patients being discharged following hospital for COVID-19 related illness.

The EtD framework for Recommendation 3 was updated as of June 2023. Two randomized controlled trials were identified that provided evidence related to this question regarding the effects of postdischarge outpatient prophylactic-intensity anticoagulation compared with no anticoagulation on multiple critical outcomes among all-cause mortality, pulmonary embolism (PE) deep venous thrombosis (DVT), ischemic stroke, major bleeding, STEM, and readmission.<sup>61,62</sup> One of the trial groups provided unpublished data on request for selected outcomes. The overall certainty of the evidence of effects was low. This was primarily due to extremely serious imprecision and for some outcomes risk of bias (see Evidence Profile and EtD framework online at: https://guidelines.ash.gradepro.org/profile/hzJoT4NBkkk).

Based on the panel's thresholds for effect sizes, post-discharge outpatient prophylactic-intensity anticoagulation probably results in little to no difference for DVT (OR: 0.51, 95% CI: 0.08 to 3.29; corresponding to 1 fewer [from 3 fewer to 7 more] DVTs per 1,000 patients), and may result in little to no difference for all-cause mortality (OR: 0.84, 95% CI: 0.37 to 1.89; corresponding to 3 fewer [12 fewer to 16 more] deaths per 1,000 patients), PE (OR: 0.66, 95% CI: 0.08 to 5.44; corresponding to 2 fewer [6 fewer to 30 more] PEs per 1,000 patients), STEMI (OR: 0.14, 95% CI: 0.01 to 2.74; corresponding to 5 fewer [from 6 fewer to 10 more] STEMI per 1,000 patients), and readmission (OR: 0.20, 95% CI: 0.01 to 4.15; corresponding to 25 fewer [31 fewer to 86 more] STEMI per 1,000 patients). Post-discharge anticoagulation may have little to effect on ischemic stroke (OR: 2.99, 95% CI: 0.12 to 73.55; corresponding to 4 more [from 2 fewer to 126 more] ischemic strokes per 1,000 patients), but the evidence is very uncertain.

In terms of potential harms, post-discharge anticoagulation may have little to no effect on major bleeding in patient with COVID-19 (OR: 1.99, 95% CI: 0.18 to 22.04; corresponding to 3 more [from 2 fewer to 59 more] major bleeds per 1,000 patients), and probably has little to effect on major bleeding in other patients being discharged (indirect evidence - OR: 2.09, 95% CI: 1.33 to 3.27; corresponding to 4 more [from 2 fewer to 125 more] per 1,000 patients).

## **Conclusions for this recommendation**

The panel judged the benefits of post-discharge outpatient thromboprophylaxis to be trivial in terms of absolute effects on all critical outcomes. This judgment was based primarily on the low baseline risk estimates for thrombotic events after hospital discharge. Meanwhile, the risk of major bleeding was also judged to be of trivial magnitude, based on low baseline risk estimates along with indirect evidence from non-COVID patients.<sup>63-65</sup> Of note, patients with high bleed risk characteristics were excluded from the MICHELLE trial (e.g. recent bleeding, recent major surgery, known coagulopathy or bleeding diathesis, prior intracranial hemorrhage, recent gastroduodenal ulcer, thrombocytopenia active cancer) and the ACTIV-4c trial (e.g. recent intracranial bleed, stroke or neurosurgery, recent major surgery, inherited or acquired bleeding disorder, thrombocytopenia).

On balance, the panel judged that the undesirable potential major bleeding complications outweighed the potential benefits, particularly considering the low baseline risk of post-discharge VTE. The panel emphasized the importance of an individualized decision for each patient based on an assessment of thrombosis and bleeding risk. This thrombosis risk assessment may include the use of externally validated RAMs such as the IMPROVE-DD risk score, which was used in the MICHELLE trial to identify patients at potentially higher thrombotic risk for study inclusion.<sup>28,60</sup> No risk assessment models for bleeding have been validated in patients with COVID-19.

# Conclusions: what others are saying and where we go from here

At the onset of the COVID-19 pandemic, the ASH living guidelines were created to answer urgent questions in a time of rapidly evolving evidence and clinical experience. The living phase (i.e., continuous review and updating) is concluded. Going forward, ASH will maintain these guidelines through regular review and scheduled revision. It was noted by the panel that the included trials primarily enrolled patients early in the COVID-19 pandemic and that the applicability of these results to the current phase of the pandemic are unclear due to potential differences in the patient population, baseline rates of VTE, and illness severity related to evolution of viral variants, prior infection, and use of non-anticoagulant therapies (corticosteroids, vaccination, antiviral therapies, monoclonal antibodies) which have contributed to improvements in the burden and severity of COVID-19 disease.

## **Other guidance**

Four years after the onset of the the pandemic, multiple guideline documents on the use of anticoagulation in patients with COVID-19 are available. These other guidance documents include the 2022 CHEST (American College of Chest Physicians [ACCP]) COVID-19 guidelines update, the 2024 International Society on Thrombosis and Haemostasis (ISTH) 2023 ISTH update of the 2022 ISTH guidelines for antithrombotic treatment in COVID-19, National Institutes of Health (NIH) COVID-19 treatment guidelines and European Society of Cardiology (ESC) guidance for the diagnosis and management of cardiovascular disease during the COVID-19 pandemic<sup>66-69</sup>.

Major methodologic differences between the current ASH guidelines and these other documents include use of high-quality systematic reviews and EtD frameworks, which increase transparency, along with use of marker states and decision thresholds to estimate the relative importance to patients as key outcomes of treatment. The present ASH guideline is also unique in its "living" format, though other guidance documents may also be updated.

Amongst critically ill patients, the guidance documents from ASH, ACCP, ISTH, and NIH uniformly suggest prophylactic-intensity anticoagulation (as opposed to intermediate- or therapeutic-dose anticoagulation) for patients without suspected or confirmed VTE.

Meanwhile, in acutely ill COVID-19 patients, most guidance documents suggest or recommend that therapeutic-intensity anticoagulation be considered in preference to prophylactic-intensity anticoagulation. The NIH is more specific in recommending that therapeutic-dose heparin be used for patients who have an elevated D-dimer, who are on low-flow oxygen, and who have low bleeding risk. The ISTH guidelines recommend that therapeutic LMWH or UFH is beneficial in preference to intermediate- or prophylactic-dose LMWH or UFH in select non-critically ill patients.<sup>66</sup> ESC guidance endorses anticoagulation at standard-dose prophylactic doses for hospitalized patients with COVID-19.<sup>69</sup>

Finally, regarding post-discharge thromboprophylaxis, these other guidance documents also do not recommend the routine use of post-discharge pharmacological thromboprophylaxis. However, in the absence of high-quality evidence, they generally suggest that an individualized decision be made, balancing the patient's thrombosis and bleeding risk factors at the time of discharge, and that thromboprophylaxis may be considered for select patients. The CHEST 2020 guideline suggests that post-discharge thromboprophylaxis would only result in net clinical benefit if the risk of symptomatic VTE were found to be above 1.8% within 35 to 42 days after release from the hospital<sup>67</sup>, while the

updated 2022 CHEST guideline did not comment specifically on post-discharge thromboprophylaxis.<sup>70</sup> The 2023 ISTH guideline suggests that post-discharge thromboprophylaxis with prophylactic dose rivaroxaban may be considered for approximately 30 days to reduce the risk of VTE after hospitalization for COVID-19, particularly in patients with persistent VTE risk factors that may include a high IMPROVE risk score, or high D-dimer.<sup>66</sup>

There are several ongoing trials in a variety of settings that may have implications for patients with COVID-19.<sup>71</sup> This includes studies of primary thromboprophylaxis with direct oral anticoagulants in non-hospitalized outpatients (e.g., PREVENT-HD [NCT04508023], HERO-19 [NCT04359246]), and anticoagulation in hospitalized non-ICU patients (e.g., XACT [NCT04640181], FREEDOM COVID-19 [NCT04512079]). Ongoing studies in critically ill COVID-19 patients also include novel therapeutic approaches including the use of nebulized heparin (e.g., CHARTER-MT [NCT04397510]) and fibrinolytics for acute respiratory distress syndrome (e.g., STARS [NCT04357730]), TRISTARDS [NCT04640194]).

## **Future research priorities**

Based on gaps in evidence identified during the guideline development process, the panel identified the following research priorities in this patient population:

- Large, high-quality randomized controlled trials to increase the certainty of the evidence on health effects,
- Studies examining the impact of non-anticoagulant interventions (e.g., vaccines, corticosteroids, antiviral therapies, antiplatelet therapies, anti-cytokine therapies, monoclonal antibody therapies) on thrombotic risk,
- Studies examining the impact of different viral variants on thrombotic risk
- Further development and validation of risk assessment models for thrombosis and bleeding in prospective cohorts of patients with COVID-19 during and after hospitalization,
- Studies examining the impact of anticoagulant therapy on thrombosis and bleeding according to social determinants of health

# Limitations of these guidelines

The limitations of these guidelines are inherent in the low to very low certainty of the evidence we identified for the research questions. This relates to risk of bias, as well as imprecision which may also relate to heterogeneity in study designs, patient characteristics, and outcome measurements used.

In addition, non-anticoagulant treatments administered to hospitalized patients with COVID-19 (e.g., corticosteroids, anti-cytokine therapies, ventilatory support), patient characteristics, viral variants and immunity have changed over the course of the pandemic. It remains uncertain how advancements in clinical care may impact the baseline risk of VTE in-hospital and after hospital discharge. Evidence collected earlier in the pandemic and included in our systematic reviews may not fully reflect the baseline risk of VTE or the effect of thromboprophylaxis in the current phase of the pandemic, due to the impact of vaccination, prior infection, viral variants, and other non-antithrombotic therapies on COVID-19 disease course and severity, and baseline VTE risk.

# Updating or adapting recommendations locally

Adaptation of these guidelines will be necessary in many circumstances. These adaptations should be based on the associated EtD frameworks.<sup>12</sup>

# **Acknowledgments**

The authors acknowledge Rob Kunkle, Eddrika Russell, Deion Smith, Natale DiFlorio, Kendall Alexander and Meghin Brooks for their overall coordination of the guideline panel. The authors acknowledge the investigators of the HEP-COVID, BEMICOP, HEP-COVID, and RAPID trials for sharing unpublished data regarding selected prioritized outcomes for the guidelines. The authors thank Susan Kahn and Jennifer Davila for their participation in the panel and previous contributions to recommendations and publications. DRT was supported by a career development award from the National Institutes of Health, National Heart, Lung and Blood Institute, award number 1K01HL135466. DMS is supported by a Tier 2 Canada Research Chair in Anticoagulant Management of Cardiovascular Disease.

# **Authorship contributions**

D.M.S, E.K.T., and R.N. wrote the manuscript. All other authors contributed to critical revisions of the manuscript. All authors approved of the content. Members of the knowledge synthesis team (R.N., R.A.J., Y.A.J., L.E.C.L., K.D., A.J.D., S.G.K., G.P.M., R.Z.M., B.A.P., Y.R.B., K.S., W.W.) searched the literature, extracted data from eligible studies, analyzed the data, and prepared evidence summaries and evidence to decision tables. Panel members (D.M.S., E.K.T., H.J.S., P.A., C.B., A.C., K.D., M.T.D., D.D., D.O.G., F.A.K., A.I.L., I.N., A.P., M.R., K.M.S., D.M.S., M.S., D.R.T., K.T., R.A.M., R.N.) assessed the evidence, voted, and made judgments within the evidence to decision framework, and discussed and issued the recommendations. The methods leadership team (R.N., R.B.P., K.D., A.S., K.S., A.C., E.A.A., W.W., R.A.M., H.J.S.) developed the methods and provided guidance to the knowledge synthesis team and guideline panel. D.M.S., R.A.M., and R.N. were the co-chairs of the panel and led panel meetings.

# **Disclosures of conflicts of interest**

All authors were members of the guideline panel or members of the systematic review team or both. As such, they completed a disclosure of interest form, which was reviewed by ASH and is available as Supplements 1 and 2.

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# **Tables**

Table 1. Recommendations.

Recommendation	Remarks		
<b>Recommendation 1a.</b> The American Society of Hematology (ASH) guideline panel suggests using prophylactic-intensity over intermediate-intensity anticoagulation in patients with COVID-19- related critical illness who do not have suspected or confirmed venous thromboembolism (VTE) (conditional recommendation based on low certainty in the evidence about effects $\bigoplus \bigoplus \bigcirc$ ).	illness are defined as those suffering from an immediately life-threatening condition who would typically be admitted to an intensive care unit (ICU) due to COVID-19 infection. Examples include patients requiring hemodynamic support, ventilatory support, and renal replacement therapy. This does not include patients admitted to the ICU for		
Recommendation 1b. The ASH guideline panel suggests using prophylactic-intensity over therapeutic-intensity anticoagulation in patients with COVID-19-related critical illness who do not have suspected or confirmed venous thromboembolism (VTE) (conditional recommendation based on very low certainty in the evidence about effects ⊕).	<ul> <li>other reasons who incidentally test positive for COVID-19.</li> <li>An individualized assessment of the patient's risk of thrombosis and bleeding is important when deciding on anticoagulation intensity. Risk assessment models (RAMs) to estimate thrombotic risk have been validated in</li> </ul>		

	<ul> <li>hospitalized patients with COVID-19 (critically or non-critically ill), with modest prognostic performance. No RAMs for bleeding have been validated for patients with COVID-19. The panel acknowledges that higher-intensity anticoagulation may be preferred for patients judged to be at low bleeding risk and high thrombotic risk.</li> <li>At present, there is no direct high- certainty evidence comparing different types of anticoagulants. The selection of a specific agent (e.g., low molecular weight heparin [LMWH], unfractionated heparin [UFH]) may be based on availability, resources required, familiarity, and the aim of minimizing the use of personal protective equipment or exposure to staff to COVID-19-infected patients as well as patient-specific factors (e.g., renal function, history of heparin- induced thrombocytopenia, concerns about gastrointestinal tract absorption). LMWH and UFH were used in the identified studies and may be preferred because of a preponderance of evidence with these agents. There are no studies of intermediate- or therapeutic-intensity fondaparinux, argatroban, or bivalirudin in this population.</li> <li>These recommendations do not apply to patients who require anticoagulation to prevent thrombosis of extracorporeal circuits such as those on extracorporeal membrane oxygenation (ECMO) or continuous renal replacement therapy (CRRT).</li> </ul>
<b>Recommendation 2a.</b> The ASH guideline panel suggests using prophylactic-intensity over intermediate-intensity anticoagulation in patients with COVID-19-related acute illness who do not have suspected or confirmed VTE (conditional recommendation based on very low certainty in the evidence about effects $\bigoplus$ ).	<ul> <li>Patients with COVID-19-related acute illness are defined as those with clinical features that would typically result in admission to an inpatient medical ward without requirement for intensive clinical support. Examples include patients with dyspnea or mild-to-moderate hypoxia.</li> <li>An individualized assessment of the patient's risk of thrombosis and bleeding</li> </ul>
suggests using therapeutic-intensity over	is important when deciding on

prophylactic-intensity anticoagulation in patients with COVID-19-related acute illness who do not have suspected or confirmed VTE (conditional recommendation based on very low certainty in the evidence about effects ⊕).	<ul> <li>anticoagulation intensity. RAMs to estimate thrombotic risk have been validated in hospitalized patients with COVID-19 (critically or non-critically ill), with modest prognostic performance. No RAMs for bleeding have been validated for patients with COVID-19. The panel acknowledges that lower-intensity anticoagulation may be preferred for patients judged to be at high bleeding risk and low risk of thrombosis.</li> <li>At present, there is no direct high- certainty evidence comparing different types of anticoagulants in patients with COVID-19. LMWH or UFH may be preferred because of a preponderance of evidence with these agents. There are no studies of therapeutic-intensity fondaparinux, argatroban, or bivalirudin in this population.</li> </ul>
<b>Recommendation 3.</b> The ASH guideline panel suggests against using post-discharge outpatient anticoagulant thromboprophylaxis in patients with COVID-19 who are being discharged from hospital and who do not have suspected or confirmed VTE or another indication for anticoagulation (conditional recommendation based on very low certainty in the evidence about effects $\bigoplus \bigcirc \bigcirc$ ).	<ul> <li>An individualized assessment of the patient's risk of thrombosis and bleeding and shared decision-making are important when deciding on whether to use post-discharge thromboprophylaxis.</li> <li>The panel acknowledged that post-discharge thromboprophylaxis may be reasonable in patients judged to be at high thrombotic risk and low bleeding risk.</li> </ul>

## Table 2. Descriptions of target populations.

Target population	Definition
Critically ill	Patients with COVID-19 who develop respiratory or cardiovascular
	failure normally requiring advanced clinical support in the ICU or CCU,
	but could include admission to another department of the ICU/CCU was
	over capacity. ICU/CCU capacity and admission criteria could vary
	according to the specific setting. This does not include patients admitted
	to the ICU for other reasons who incidentally test positive for COVID-19.
Acutely ill	Patients with COVID-19 who require hospital admission, generally to an
	inpatient medical ward, without intensive clinical support (i.e., not in the
	IC/CCU), but may be treated in other settings if the hospital is over
	capacity. Hospital capacity and admission criteria may vary according to

	the specific setting. Some observational studies informing the baseline	
	risk of critical outcomes reported on all patients hospitalized with	
	COVID-19 in aggregate had fewer than 20% in the ICU without	
	separating their outcomes. Such patients were labeled as acutely ill.	
Post-discharge	Patients discharged from acute care hospital following COVID-19-related	
	critical illness or acute illness.	