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Rationale and design for the study of rivaroxaban to reduce thrombotic events, hospitalization and death in outpatients with COVID-19: The PREVENT-HD study

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Background COVID-19 is associated with both venous and arterial thrombotic complications. While prophylactic anticoagulation is now widely recommended for hospitalized patients with COVID-19, the effectiveness and safety of thromboprophylaxis in outpatients with COVID-19 has not been established.

Study Design PREVENT-HD is a double-blind, placebo-controlled, pragmatic, event-driven phase 3 trial to evaluate the efficacy and safety of rivaroxaban in symptomatic outpatients with laboratory-confirmed COVID-19 at risk for thrombotic events, hospitalization, and death. Several challenges posed by the pandemic have necessitated innovative approaches to clinical trial design, start-up, and conduct. Participants are randomized in a 1:1 ratio, stratified by time from COVID-19 confirmation, to either rivaroxaban 10 mg once daily or placebo for 35 days. The primary efficacy end point is a composite of symptomatic venous thromboembolism, myocardial infarction, ischemic stroke, acute limb ischemia, non-central nervous system systemic embolization, all-cause hospitalization, and all-cause mortality. The primary safety end point is fatal and critical site bleeding according to the International Society on Thrombosis and Haemostasis definition. Enrollment began in August 2020 and is expected to enroll approximately 4,000 participants to yield the required number of end point events.

Conclusions PREVENT-HD is a pragmatic trial evaluating the efficacy and safety of the direct oral anticoagulant rivaroxaban in the outpatient setting to reduce major venous and arterial thrombotic events, hospitalization, and mortality associated with COVID-19. (Am Heart J 2021;235:12–23.)

COVID-19 has rapidly emerged as the world’s most pressing infectious threat. The novel severe acute respiratory syndrome coronavirus-2 (SARS Co-V-2) responsible for this condition has proven to be readily transmissible, with significant morbidity and a high case fatality rate. SARS Co-V-2 has further demonstrated wide-ranging systemic effects, including significant immunologic, pulmonary, gastrointestinal, cardiac, and neurologic manifestations. A particularly concerning risk that has emerged with COVID-19 is the development of an activated coagulation system associated with macrovascular and microvascular thrombosis and overall poor prognosis. The incidence of venous or arterial thrombotic events in hospitalized patients may be as high as 1 in 6, and up to 1 in 3 in patients requiring intensive care depending on whether surveillance imaging for asymptomatic venous thromboembolism (VTE) is performed. Due to this pronounced hypercoagulable state, attention has focused on antithrombotic treatment to reduce morbidity and mortality in COVID-19. Retrospective analyses suggest lower mortality rates for hospitalized patients with COVID-19 who received prophylactic anticoagulation, compared to those who did not. Preliminary reports from ongoing prospec-

While acknowledging the potential benefit of posthospitalization thromboprophylaxis, expert opinion and guidance statements have disagreed on the need for primary thromboprophylaxis in outpatients with COVID-19 with thrombotic risk factors. The underlying mechanisms of the hypercoagulable state in patients with COVID-19 are not clear. A key question is: when in the course of SARS-Co-V-2 infection does thrombotic risk reach a critical, yet modifiable point? There are data supporting activated thrombin as a key pathogenic driver of pulmonary compromise in COVID-19. Fibrinogen and D-dimer concentrations are often already elevated at the time of hospital admission, and elevated D-dimer concentrations are found in almost half of hospitalized patients with nonsevere disease. Additionally, up to half of venous thromboembolic events in hospitalized patients in one series were diagnosed within the first 24 hours of admission. We hypothesize that the increased risk of thrombotic events, attributable to a thrombotic-inflammatory status associated with reduced mobility, begins prior to severe clinical manifestations of COVID-19, and includes patients who do not require hospitalization. Multiple autopsy series have reported venous thromboembolism and widespread pulmonary microthrombi in decedents with COVID-19, suggesting a role of direct endothelial injury in the development of COVID-19 pulmonary manifestations (Figure 1). Therefore, we hypothesize that intervening to decrease thrombotic risk earlier in the course of COVID-19, especially in patients with known risk factors for thrombosis, will significantly decrease thrombotic complications and reduce disease progression to the point where hospitalization could be avoided.

Direct oral anticoagulants (DOACs) are favored due to their oral administration, selective coagulation factor inhibition, lack of required blood monitoring, and safety profile relative to vitamin K antagonists. Early observations of lower than expected mortality in subjects on DOACs with chronic atrial fibrillation who contract COVID-19 suggested that anticoagulation may benefit patients with COVID-19 in the prehospital setting. DOACs may be a preferred choice over other anticoagulant options for post-hospital discharge from COVID-19 with other indications for therapeutic thromboprophylaxis.

Rivaroxaban, a selective factor Xa inhibitor, has been investigated in a comprehensive cardiovascular development program, establishing risk reduction of venous and arterial thrombotic events in a variety of indications. Rivaroxaban has shown benefit in decreasing the risk of venous thromboembolic events in medically ill inpatients with elevated thrombotic risk and low risk of bleeding, including those with pneumonia and sepsis, starting in-hospital and continuing during the posthospital discharge period. Outpatients with COVID-19 may represent a similar medically ill population that could benefit from thromboprophylaxis with rivaroxaban, but the need for, and the net clinical benefit of, anticoagulation at an early stage of COVID-19 is currently unknown. Given the potential role of factor Xa in the pathogenesis of coronavirus morbidity and mortality, rivaroxaban may have benefits in preventing the progression to severe COVID-19. The PREVENT-HD trial is designed to evaluate the efficacy and safety of prophylactic-dose rivaroxaban in outpatients with symptomatic COVID-19 infection with at least one additional thrombotic risk factor.

Study design and population

PREVENT-HD is a multicenter, randomized, double-blind, placebo-controlled, pragmatic, event-driven phase 3 study to assess the efficacy of rivaroxaban 10 mg once daily in preventing venous and arterial thrombotic events, hospitalization, and death in outpatients with symptomatic COVID-19. The trial is being conducted at large integrated health care delivery networks to facilitate centralized patient recruitment and follow-up, and to enable a pilot study of alternative methods of data collection through integration of a cloud-based data capture tool with electronic medical records (EMR). The study design incorporates several innovative processes intended to reduce exposure of health care providers to SARS Co-V-2, monitor the safety of study participants, and enhance the efficiency of study conduct using advanced informatics integration (Table 1). The study design is shown in Figure 2.

The inclusion criteria are designed to enroll a representative population of symptomatic adult outpatients with COVID-19 with at least one additional risk factor for thrombosis and with a low bleeding risk, and for whom the initial treatment plan does not include hospitalization. Study participants must be at least 18 years of age, have documented COVID-19, have symptoms attributable to COVID-19, and have an additional risk factor for venous or arterial thromboembolic disease, including elevated D-dimer level, thrombophilia, prior venous thromboembolism, history of cancer, coronary artery disease, peripheral artery disease, cerebrovascular disease or ischemic stroke; or a risk factor associated with worse COVID-19 outcomes (age ≥ 60 years of age, body mass index ≥ 35 kg/m², or history of heart failure or diabetes requiring medication). Key exclusion criteria include convalescent infection (eg, positive COVID antibody test or other serology test at least 2 weeks following acute infection), conditions that pose an increased risk of bleeding (eg, bronchiectasis, active cancer undergoing treatment,
Coagulopathy and COVID-19 pathogenesis. Coagulopathy and diffuse pulmonary microthrombi have been documented in COVID-19. While coagulopathy is a known consequence of inflammatory changes, it is unclear if SARS-Co-V-2 independently affects hypercoagulability. Coagulopathy, along with viral endothelial injury, leads to diffuse pulmonary microthrombi which may potentiate pulmonary injury in addition to alveolar damage from SARS-Co-V-2 infection as well as macrothrombotic events. Factor Xa can also play a role in cell entry and infection by SARS-Co-V-2, and therefore viral propagation. Outpatient anticoagulation with rivaroxaban, a specific Factor Xa inhibitor, has the potential to prevent thromboembolic events as well as pulmonary microthrombi and progression of pulmonary insufficiency in COVID-19, reducing the need for hospitalization.

Active gastroduodenal ulcer, significant bleeding in the prior 3 months and use of dual antiplatelet therapy, and recent or planned therapy with medications that significantly impact rivaroxaban concentrations or bleeding risk, including anticoagulants. A complete listing of inclusion and exclusion criteria is provided in Table 2.

Subjects are identified through site specific EMR searches to notify coordinators of potential subjects having a recent positive test result for COVID-19. At some sites, such searches are supplemented with additional inclusion and exclusion criteria verifiable through the EMR. Potential subjects are then called to verify eligibility and to assess interest. Consent is generally obtained electronically, while the site communicates directly by phone or by virtual meeting platforms. For some participants unable to provide consent electronically, the consent is verified using paper forms.

Participants meeting all inclusion and no exclusion criteria may be randomized from the date of the positive COVID-19 test until 14 days later, inclusive. All
participants will be randomized in a 1:1 ratio to receive either rivaroxaban 10 mg or matching oral placebo once daily for 35 days through use of a central computerized interactive voice/web response system that conceals study drug assignment from investigators. Randomization will be further stratified by the time from positive diagnostic of COVID-19 test to randomization, into an early-dosed cohort (1-5 days) and a later-dosed cohort (6-14 days). The protocol also allows capping of enrollment of participants with certain risk factors to enable an adequate assessment of subgroups with particular risk factors. Enrollment began in August 2020 and will continue to an expected total of approximately 4,000 participants randomized to achieve the target number of end point events. The study is being performed in accordance with all local laws and regulations, and with the ethical principles of the Declaration of Helsinki, and the International Council on Harmonization Good Clinical Practice guidelines. The study protocol and informed consent have been reviewed and approved by the responsible health authorities and Institutional Review Boards for all participating study sites.

### Treatment protocol and follow-up procedures

#### Treatment selection

Study drug provided to participants is double-blinded rivaroxaban 10 mg or matching placebo taken once daily. This dose and duration of rivaroxaban was selected for PREVENT-HD primarily based on previous results in medically ill populations,\(^{32,34,38,39}\) suggesting that the 10 mg once daily dose of rivaroxaban is the minimally effective dose needed to reach the trough level to prevent venous thromboembolic events in medically ill populations with high thrombotic risk and low risk of bleeding, including those with pneumonia and sepsis.\(^ {50}\) Furthermore, the 10 mg once daily dose is FDA-approved for VTE prophylaxis following hip and knee arthroplasty and has trough levels approximating the 2.5 mg twice daily dose,\(^ {49}\) which has been previously shown to be effective in preventing thrombotic events in patients with cardiovascular disease.\(^ {53,55}\) Though the 2.5 mg twice daily dose has shown efficacy in preventing both arterial and venous thrombotic events in these studies, the 10 mg once daily dose was favored for the current study given the efficacy demonstrated previously in similar medically ill populations. While there is a dearth of evidence that DOACs reduce the risk of arterial thrombosis in medically ill populations, recent emerging evidence suggests non-hemorrhagic stroke prevention with betrixaban\(^ {41}\) and rivaroxaban\(^ {42}\) in this population. In the MARINER study, a placebo controlled study of extended thromboprophylaxis in medically ill patients with rivaroxaban, 10 mg of rivaroxaban reduced the secondary composite end point of symptomatic VTE, myocardial infarction, non-hemorrhagic stroke, and cardiovascular death by 28% compared with placebo without a significant increase in major bleeding.\(^ {45}\) Doses greater than 10 mg once daily of rivaroxaban have been associated with increased bleeding risk when used for secondary prophylaxis.\(^ {55}\) Therefore, 10 mg once daily dose was selected to optimally balance the prevention of both venous and arterial thrombotic risk with the risk of excess bleeding.

#### Concomitant therapies

The protocol allows treatment with all clinically-recommended therapies according to local practice,
including any prescribed medications intended to inhibit SARS Co-V-2 activity. Drugs that interact with rivaroxaban to either significantly increase or decrease rivaroxaban concentrations are prohibited; such medications include combined P-glycoprotein and strong inhibitors of CYP3A4, or combined P-glycoprotein and strong CYP3A4 inducers. Of importance, dexamethasone, commonly used in the treatment of COVID-19, is not a prohibited medication. Medications that significantly increase the risk of bleeding in the setting of rivaroxaban therapy, such as anticoagulants, dual antiplatelet regimens, or high-dose antiplatelet monotherapy (eg, aspirin >162 mg daily, clopidogrel >75 mg daily, or ticlopidine >250 mg daily), are also prohibited. Single antiplatelet agents including these drugs at lower doses, ticagrelor, and prasugrel are allowed, with the caution that participants should be monitored for increased risk of bleeding. If any prohibited medication is clinically indicated, blinded study drug is to be discontinued while the prohibited medication is taken.

Visit schedule and follow-up

Participants are randomized on study Day 1 and will take blinded study medication once daily through to study Day 35. All study activities will occur remotely, including screening for eligibility, consent, and follow-up. Following randomization, participants are automatically flagged for drug dispensation through an automated system. A centralized drug depot sends study drug directly to the patient’s home by the next day. Delivery is confirmed by telephone contact on Day 3. To further minimize potential study related COVID-19 exposures, subjects receive detailed instructions for safe
self-disposal of any unused study drug rather than being required to return drug to the site. Participants will undergo virtual study visits on Day 1 (defined as Day of randomization), and on Days 3, 14, 35, and 49. At each follow-up contact, participants are assessed for study drug adherence (via questioning start/stop dates and missed doses of study drug), adverse events including bleeding, and any potential study end point events. To aid in site follow-up of subjects, PREVENT-HD will leverage the REDCap Cloud platform to integrate with each site’s local EMR system (Figure 3). This integration will facilitate the transfer (or direct import) of data pertaining to demography, medical history, medications, encounters, diagnostic procedures, blood transfusions, and laboratory values into a parallel clinical database to facilitate the monitoring of subject health status and outcomes. As thromboprophylaxis is now recommended for all hospitalized COVID-19 patients, upon hospitalization, participants will discontinue blinded study medication and will receive standard of care open label thromboprophylaxis per guidelines. Upon discharge, participants may be continued on extended open label thromboprophylaxis if it is the practice of the institution or may go back on blinded study medication if that is not the case.

Pilot study on EMR data reliability

The EMR domains selected for the pilot study were purposely chosen to be generic and applicable to future clinical studies (eg, health care or hospital encounters, demographics, medications, labs, prior medical, or surgical diagnoses). This component of the trial is intended to explore new methods of data acquisition that may expand central monitoring capabilities and improve the efficiency of site monitoring by reducing or eliminating the requirement to perform on-site or remote source data verification. The pilot will also explore whether EMR data can replace manual transcription of study data into the clinical database for selected fields. For this study, all data will also be manually entered by site personnel into a conventional electronic data capture system for regulatory purposes and to allow for comparisons.

Study end points

The primary efficacy end point is a composite including symptomatic VTE, myocardial infarction, ischemic stroke, acute limb ischemia, non-central nervous system (non-CNS) systemic embolization, all-cause hospitalization, and all-cause mortality up to Day 35. Because many of the components of this end point are typically associated with hospitalization, the study end point represents significant morbidities that are meaningful to an outpatient population. Secondary and exploratory efficacy end points are further listed in Table 3 and include emergency department visits and need for supplemental
Data flow in PREVENT-HD. Study data are collected remotely by site staff. Key data flow in daily to weekly from the local hospital electronic medical records (EMR), through REDCap Cloud, to a parallel clinical database. Data from EMR can be used in real time to identify eligible subjects to consent remotely, and to monitor for outcome events in enrolled participants. Site staff conduct virtual follow-up visits by phone or telehealth and enter data from outside the hospital system into electronic case report forms. Participants do not need to leave home through the duration of the study.

**Table 3. PREVENT-HD trial outcomes**

**Primary efficacy outcome**
Time to first occurrence of a composite endpoint of symptomatic venous thromboembolism (VTE), myocardial infarction (MI), ischemic stroke, acute limb ischemia, non-central nervous system (non-CNS) systemic embolization, all-cause hospitalization, and all-cause mortality up to Day 35

**Secondary efficacy outcomes**
- Time to first occurrence of a composite endpoint of symptomatic VTE, MI, ischemic stroke, acute limb ischemia, non-CNS systemic embolization, and all-cause mortality up to Day 35
- Time to first occurrence of all-cause hospitalization up to day 35
- Time to first occurrence of symptomatic VTE up to day 35
- Time to first occurrence of symptomatic VTE up to day 35
- Time to first occurrence of an emergency room (ER) visit up to Day 35
- Time to first occurrence of symptomatic VTE, MI, ischemic stroke, acute limb ischemia, non-CNS systemic embolization, and all-cause hospitalization up to day 35
- Incidence of participants who are hospitalized or dead from any cause on day 35
- Time to all-cause mortality up to day 35

**Exploratory efficacy outcomes**
- World Health Organization [WHO] Research and Development Blueprint: Novel Coronavirus Scale for Clinical Improvement over time
- Time to first occurrence of a component event of the primary efficacy endpoint (MI, ischemic stroke, acute limb ischemia, and non-CNS systemic embolization) up to day 35
- The incidence of participants achieving an oxygen saturation ($O_2$ sat) below 92% on room air at rest or with ambulation up to day 35
- The incidence of participants achieving an $O_2$ sat below 88% on room air at rest or with ambulation up to day 35
- The incidence of participants requiring supplemental oxygen up to day 35
- Time to first occurrence of the use of noninvasive ventilation or high-flow oxygen (WHO 5), intubation and mechanical ventilation (WHO 6), or ventilation and additional organ support (vasopressors, renal replacement therapy [RRT], extracorporeal membrane oxygenation [ECMO]; WHO 7) or all-cause mortality (WHO 8) up to day 35
- The incidence of participants requiring dialysis or having an estimated glomerular filtration rate (eGFR) $<15\text{ mL/min/1.73 m}^2$ on 2 measurements more than 24 hours apart up to day 35
- Time to first occurrence of disseminated intravascular coagulation (DIC) up to day 35
- Time to first occurrence of acute respiratory distress syndrome (ARDS) up to day 35
- The incidence of occurrence of COVID death up to day 35
- Medical Resource Utilization data over time

**Primary safety outcome**
Time to first occurrence of International Society on Thrombosis and Hemostasis (ISTH) critical site and fatal bleeding on treatment [+2 days]

**Secondary safety outcomes**
- Time to first occurrence of ISTH major bleeding on treatment [+2 days]
- Time to first occurrence of nonmajor clinically relevant bleeding on treatment [+2 days]
oxygen therapy. The primary safety end point is fatal and critical-site bleeding according to the International Society on Thrombosis and Haemostasis definition. Additional safety end points will include International Society on Thrombosis and Haemostasis major bleeding and nonmajor clinically relevant bleeding.

All primary end points and most secondary end points will undergo adjudication against trial end point definitions by trained, experienced physicians at each site who remain blinded to treatment assignment. To participate as a physician adjudicator, individuals must undergo additional training on the application of standardized event definitions. Adjudicators are instructed to contact the academic research organization for questions related to complicated cases or unanticipated circumstances, which are then logged as conventions to be applied to future cases across sites. This model of site-level adjudication allows for efficiency gains by taking advantage of full access to local EMR systems for source document review. All end point definitions used in PREVENT-HD are provided in the Appendix. Centralized oversight of local adjudicators is conducted through centralized training, use of conventions, and guidance to local adjudicators through periodic meetings to minimize site-level variability in adjudication.

**Statistical considerations**

Efficacy analyses will be conducted on all randomized patients using the principle of intention-to-treat up to day 35 after randomization; safety analyses will be conducted on all patients that received at least one dose of study drug until 2 days after the last dose of study medication. Treatment assignment will be balanced within a clinical site by block randomization. The randomization will be stratified by the time from COVID-19 positive test to randomization (1-5 days inclusive, 6-14 days inclusive). The primary efficacy analysis will be based on the time from randomization to the first adjudicated occurrence of any component of the primary composite end point up to day 35. The primary efficacy analysis will be done using a stratified log-rank test by the time from COVID-19 positive test to randomization with the treatment as a variable. The primary efficacy outcome will be tested at a 2-sided significance level of 5%, with appropriate apportioning of alpha to account for one planned interim analysis. Secondary end points will be analyzed hierarchically, in the order listed in Table 3, using similar time-to-event analyses described above.

The trial is event-driven such that it would require 333 confirmed first primary end point events to detect an anticipated hazard ratio of 0.70 between the study arms, with 90% power at a 2-sided 5% significance level. It was determined that the study required approximately 4,000 participants in order to have 333 patients experience a component of the primary end point over 35 days with a placebo rate of the primary outcome of 10%. If the blinded, pooled event rate proves to be lower than anticipated, enrollment of up to 5,000 participants will be considered (corresponding to an event rate in the placebo group of 7.8%, all else constant) to achieve the requisite number of events. The estimate for primary outcome event rate was estimated from unpublished data from the Northwell system (Spyropoulos AC, personal communication) suggesting a 10% hospitalization rate in unselected patients, and CDC information suggesting approximately 8% nationally in patients 18 years of age and older. The case fatality rate for unselected patients is approximately 2.3% nationally. It was also assumed that most, but not all thromboembolic events and deaths might occur after hospitalization.

An Independent Data Monitoring Committee (IDMC) is responsible for monitoring the safety of study participants throughout the trial and will perform a minimum of one interim analysis for futility or overwhelming superiority once approximately 50% of the required 333 first primary efficacy events have been observed. The stopping rule for overwhelming superiority uses an O’Brien-Fleming boundary \((Z = 2.96, \alpha = 0.003)\) approach. The IDMC may consider additional factors and request additional interim analyses (with appropriate apportionment of alpha to control the type 1 error rate) to make decisions on trial conduct.

**Study organization**

PREVENT-HD is being conducted at up to approximately 15 sites affiliated with integrated health care delivery networks in the United States. The trial is sponsored by Janssen Research and Development, LLC (Raritan, NJ) and conducted in partnership with CPC Clinical Research (CPC; Aurora, CO), a nonprofit Academic Research Organization that is affiliated with the University of Colorado Anschutz Medical Campus. An Executive Committee (EC) is responsible for oversight of the study with unrestricted access to the necessary data to fulfill this role and will submit the results of the study for publication in a peer-reviewed journal. CPC provides administrative support to the EC, and IDMC, and helps to oversee consistency in site end point confirmation. REDCap Cloud provided oversight for defining the EMR data fields to be collected from each site and for establishing the necessary technical connections for the integration/transfer of EMR data from sites to REDCap Cloud to the study database. No direct funding was provided to support development of the current manuscript. The authors are solely responsible for drafting and editing of the manuscript, and its final contents.

PREVENT-HD is registered on clinicaltrials.gov under number NCT04508023. Major milestones for the PREVENT-HD trial include establishment of the EC in
June 2020, initial protocol finalization in June 2020, first site open for enrollment in August 2020 with the first participant also randomized in August 2020. The first EMR integration was completed in October 2020. The planned interim analysis is anticipated by the second quarter of 2021, and study completion is anticipated by the end of 2021. These timelines will be highly dependent on the dynamics of the pandemic and effective implementation of public health measures and/or widely utilized effective vaccines.

Discussion

The COVID-19 pandemic has posed a unique global public health crisis. As our understanding of this disease evolves, there is increasing recognition of its severity and its association with a myriad of adverse complications including thrombotic events. Notably, thrombotic events have been reported in up to 4.5% of relatively healthy individuals with COVID-19 in the outpatient setting not requiring hospitalization.\(^{46,57}\) Rapidly developing therapeutic strategies to address these risks pose a specific challenge in an environment where traditional randomized trial models are impractical and pose safety issues such as exposure to infected individuals. Such challenges may foster variability in care and a desire to utilize non-randomized data to guide treatment with such approaches associated with important limitations.

The PREVENT-HD trial was designed to answer the hypothesis that initiation of prophylactic-dose rivaroxaban early in the course of COVID-19 infection, prior to hospitalization, will reduce the risk of thrombosis, hospitalization, and death, and to evaluate the safety of such a strategy. It has been designed to answer this hypothesis with the rigor of a double-blind, randomized event-driven design, but with innovative aspects intended to improve feasibility, reduce risk to study participants and study staff, and maximize data transfer and informatics capabilities.

The timeline from the beginning of protocol drafting to first participant visit was only 127 days (Figure 4), a sharp reduction from typical trial start-up timelines.\(^{46}\) This timeline was facilitated by FDA review that was both expedited and accepting of a study design which emphasizes avoidance of in-person contact. The remote electronic consent process, home drug delivery, and virtual follow-up contacts allows participants with this often physically taxing and contagious disease to continue home quarantine and convalescence throughout the trial. Fully remote site monitoring further reduces interpersonal contact at the study sites. The elimination of physical contacts helps to protect the local communities from exposure to the virus and minimizes the burden and risks to a medically fragile participant population with COVID-19. Finally, PREVENT-HD gains further efficiencies by leveraging the infrastructure of integrated health care delivery network, which serves to expand the pool of patients available to recruit from while concentrating start-up activities to fewer sites (eg, contract execution, collection of regulatory documents, and implementation of training). Integration between the local EMR and the clinical database brings key data on prospective and enrolled participants immediately to the attention of the study team. The creation of a parallel dataset of selected EMR data will further enable investigation of the reliability of EMR data and determine if efficiencies can be gained by reducing or eliminating the need for manual transcription of select study data to the primary clinical study database. Site-level adjudication of end point events allows direct access to full EMR records and minimizes the site burdens of medical record collection, redaction, and central submission processing.

In responding to COVID-19, PREVENT-HD also responds to the recognized need to increase efficiency and decrease costs of conducting registration trials.\(^{49,50}\) Many of the design innovations employed in this trial may have application beyond the current infectious pandemic. The long-term goal is to leverage learnings necessitated by the pandemic to transform future clinical trial conduct.

In addition to the innovative approach, PREVENT HD seeks to answer an important scientific question in COVID-19. While most research is currently focused on the role of anticoagulation in hospitalized patients with severe COVID-19, it is possible that decreasing thrombotic risk earlier in the course of disease may prevent some aspects of disease progression and pulmonary injury such as microvascular thrombosis. PREVENT-HD is one of a limited number of placebo-controlled trials registered on clinicaltrials.gov examining antithrombotic interventions in outpatients with COVID-19. A smaller trial \((n = 600)\) is also evaluating rivaroxaban versus placebo in outpatients (NCT04504032), while one larger outpatient primary thromboprophylaxis trial (the OVID trial, NCT04400799) is comparing enoxaparin 40 mg SQ daily versus placebo \((N = 1000)\).\(^{51}\) The National Institute of Health trial [NCT044498273] is randomizing to 1 of 4 arms \((N = 7000)\): apixaban 2.5 mg twice a day, apixaban 5 mg twice a day, low-dose aspirin once daily, or placebo. An additional large randomized, controlled open-label trial of enoxaparin versus no treatment is also under way (the ETHIC trial, NCT04492254).

Of note, 2 observational case-control analyses reported no effect of preadmission exposure to either antiplatelet therapy or anticoagulant therapy prescribed for other clinical indications on presenting acute respiratory distress syndrome, intensive care unit admission rates, or mortality rates for patients admitted with COVID-19.\(^{52,53}\) However, these analyses were of nonrandomized cohorts comprised of patients already hospitalized and prone
Study start-up timeline in PREVENT-HD. To respond to the public health crisis presented by COVID-19, study start-up timelines for PREVENT-HD were accelerated. From first draft of the protocol to first participant enrolled required only 127 days. FDA, Food and Drug Administration; FPI, first participant in; IRB, Institutional Review Board; IND, Investigational New Drug; SA, site activation.

Figure 4

Disclosure
Warren Capell reports research grants from Janssen Research and Development to CPC Clinical Research, during the conduct of the study; grants from Bayer Health Care to CPC Clinical Research, outside the submitted work. Elliot Barnathan is an employee of Janssen Research and Development, LLC, sponsor of the study, owns stock in the company, and reports a patent entitled "Prophylactic treatment of thrombotic events in outpatients" that is pending. Gregory Piazza reports grants from Janssen, during the conduct of the study; grants from BMS, grants from Bayer, grants from Portola, grants from BSC, outside the submitted work. Alex Spyropoulos...
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Supplementary materials
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References


