Hormonal intervention for the treatment of veterans with COVID-19 requiring hospitalization (HITCH): a multicenter, phase 2 randomized controlled trial of best supportive care vs best supportive care plus degarelix: study protocol for a randomized controlled trial

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Abstract

Background: Therapeutic targeting of host-cell factors required for SARS-CoV-2 entry is an alternative strategy to ameliorate COVID-19 severity. SARS-CoV-2 entry into lung epithelium requires the TMPRSS2 cell surface protease. Pre-clinical and correlative data in humans suggest that anti-androgenic therapies can reduce the expression of TMPRSS2 on lung epithelium. Accordingly, we hypothesize that therapeutic targeting of androgen receptor signaling via degarelix, a luteinizing hormone-releasing hormone (LHRH) antagonist, will suppress COVID-19 infection and ameliorate symptom severity.

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Methods: This is a randomized phase 2, placebo-controlled, double-blind clinical trial in 198 patients to compare efficacy of degarelix plus best supportive care versus placebo plus best supportive care on improving the clinical outcomes of male Veterans who have been hospitalized due to COVID-19. Enrolled patients must have documented infection with SARS-CoV-2 based on a positive reverse transcriptase polymerase chain reaction result performed on a nasopharyngeal swab and have a severity of illness of level 3–5 (hospitalized but not requiring invasive mechanical ventilation). Patients stratified by age, history of hypertension, and severity are centrally randomized 2:1 (degarelix: placebo). The composite primary endpoint is mortality, ongoing need for hospitalization, or requirement for mechanical ventilation at 15 after randomization. Important secondary endpoints include time to clinical improvement, inpatient mortality, length of hospitalization, duration of mechanical ventilation, time to achieve a normal temperature, and the maximum severity of COVID-19 illness. Exploratory analyses aim to assess the association of cytokines, viral load, and various comorbidities with outcome. In addition, TMPRSS2 expression in target tissue and development of anti-viral antibodies will also be investigated.

Discussion: In this trial, we repurpose the FDA approved LHRH antagonist degarelix, commonly used for prostate cancer, to suppress TMPRSS2, a host cell surface protease required for SARS-CoV-2 cell entry. The objective is to determine if temporary androgen suppression with a single dose of degarelix improves the clinical outcomes of patients hospitalized due to COVID-19.

Trial registration: ClinicalTrials.gov NCT04397718. Registered on May 21, 2020

Keywords: COVID-19, SARS-CoV-2, TMPRSS2, Androgen receptor, Androgen suppression, Coronavirus, Hormone therapy, Anti-androgen

Background

SARS-CoV-2 recognizes host cell membrane proteins and relies upon their enzymatic activity to infect host cells. Recognition of the host surface protein ACE2 facilitates attachment of the virus to the host cell. However, entry of the virus into the host cell requires catalytic cleavage of the Viral Spike (S) protein (a process termed S protein priming) by the host cell membrane protein TMPRSS2 [1]. Thus, TMPRSS2 is required for viral entry and infection. TMPRSS2 is expressed within the nasal mucosa, respiratory sinuses, buccal mucosa, tracheal epithelium, bronchial epithelium, lung type 2 pneumocytes, and alveolar macrophages [2–4]. In addition to the aero-digestive tract, TMPRSS2 is highly expressed on prostate, kidney, and pancreatic epithelia [4].

The transcriptional regulation of the TMPRSS2 gene has been extensively characterized, most rigorously within the prostate. The TMPRSS2 gene is located on chromosome 21 and is under the control of the androgen receptor (AR). Binding of androgens (e.g., testosterone or dihydrotestosterone) to the AR results in homodimerization and translocation of the AR to the nucleus, where it binds to its cognate androgen response element in the regulatory regions of its target genes, and thereby regulates gene expression [5, 6]. It is firmly established that suppression of AR transcriptional activity through reduction in circulating androgens or direct antagonism of AR-androgen binding using AR competitive antagonists reduces expression and protein levels of TMPRSS2 within the prostate, as well as within prostate cancers [7].

Our team queried publicly available gene expression databases to identify FDA-approved drugs that downregulate TMPRSS2. Notably, anti-androgenic compounds and estrogens were among the strongest and most consistent downregulators of TMPRSS2 expression, while androgens consistently led to upregulated TMPRSS2 gene expression [8]. In other words, these studies support the notion that the AR induces TMPRSS2 expression, whereas estrogen receptor (ER) transcriptional activity is associated with suppression of TMPRSS2 expression.

Expression of TMPRSS2 also appears to be hormonally regulated within the lung and bronchial cells [9]. Androgens enrich AR binding at the TMPRSS2 enhancer and upregulate expression of TMPRSS2 in human lung derived cells, in a fashion similarly to that found in the prostate [10]. Moreover, AR, TMPRSS2, and ACE2 are co-expressed in human lung epithelial cells including alveolar and bronchial epithelial cells [9]. In mice, testosterone suppression reduces expression of both TMPRSS2 and ACE2 in lung bronchial cells, which was reversible upon testosterone repletion [9]. In vitro, anti-androgens reduced SARS CoV 2 infection in cultured human cells [9].

Based on the aforementioned data, it is hypothesized that drugs that interfere with AR driven transcription will reduce TMPRSS2 and ACE2 expression. FDA-approved drugs that block AR signaling include the GnRH analogs that reduce pituitary release of luteinizing hormone (LH), thereby potently suppressing testosterone production, and anti-androgens that interfere with
binding of androgens to the AR. These drugs are predominantly used in the treatment of prostate cancer, have well known and generally tolerable side effect profiles, and exhibit reversibility of their biologic effects. We hypothesized that androgen suppression will reduce TMPRSS2 expression in the target cells of SARS-CoV-2 and reduce the severity of COVID-19. This general hypothesis is supported by population level data from Italy that suggest a protective effect from androgen suppressive therapy against COVID-19 in prostate cancer patients [11].

Our objective is to investigate if temporary androgen suppression that reduces expression of TMPRSS2 will reduce severity of COVID-19 disease due to SARS-CoV-2 infection. Our specific hypothesis is that the LHRH antagonist degarelix reduces the composite endpoint of mortality, need for extended hospitalization, and/or need for invasive mechanical ventilation in patients hospitalized with COVID-19.

Methods

Study design

This is a randomized phase 2, placebo-controlled, double-blind clinical trial in 198 patients testing the efficacy of degarelix plus best supportive care versus placebo plus best supportive care for improvement in the clinical outcomes of male patients who have been hospitalized due to COVID-19 (see Fig. 1 for Schema, Fig. 2 for SPIRIT). Enrolled patients must have documented infection with SARS-CoV-2 based on a positive reverse transcriptase polymerase chain reaction result performed on a nasopharyngeal swab and have a severity of illness of level 3–5 (see Table 1) [12]. Stratification is by age (<65 versus ≥ 65), history of hypertension, and disease severity score (3 versus 4 or 5), and centrally randomized 2:1 (degarelix: placebo). Best supportive care includes all treatments that would be applied irrespective of patient enrollment and includes but is not limited to supplemental oxygen, antibiotics, vasopressor support, peritoneal or hemodialysis, antibiotics, and intravenous fluids. The treatment landscape for COVID-19 is rapidly evolving. Accordingly, the clinical trial allows for updates to the allowed treatments. Remdesivir and convalescent plasma are allowed. New treatments will have a drug utilization report assessment prior to use to identify overlapping toxicities prior to use of a new treatment. The development of novel treatments and interventions.

Fig. 1 Trial schema
for COVID-19 is monitored by site investigators as well as other study team members. Appropriate changes to allowed treatments and interventions are instituted accordingly. Patients randomized receive either a one-time dose of degarelix 240 mg (a 30 day depot) or placebo administered subcutaneously in the periumbilical area. Patients, investigators, and treating physicians are blinded to treatment assignment (i.e., double blinding) and randomization is accomplished centrally Unblinding for emergency situations is managed through a 24 h a day emergency call service managed by the VA CSP. Patients are followed for up to 60 days.

Participants
The trial is open at the Greater Los Angeles, Manhattan, Brooklyn, Puget Sound, Phoenix, Central Arkansas, Long Beach, Miami, Philadelphia, Charleston, Memphis, St Louis, Dallas, and Houston Veterans Healthcare Systems. Inclusion and exclusion criteria are listed in Table 2. The trial is exclusive to males ≥ 18 who manifest symptoms and a disease severity that warrant hospitalization for supportive care but not invasive mechanical ventilation. In this patient population, the virus itself still drives the severity of the disease, whereas a hyperinflammatory response is thought to mediate ARDS, SIRS, and respiratory failure that underlie the severity of COVID-19 in critically ill, intubated ICU patients. Female patients are excluded in this study because female androgen suppression reduces endogenous estrogenic activity and thereby may induce TMPRSS2 expression, which in turn could exacerbate COVID-19.

Study drug
Degarelix is an FDA-approved drug indicated for the treatment of prostate cancer. One loading dose of 240 mg given as two injections of 120 mg, a concentration of 40 mg/mL, administered subcutaneously in the abdominal region serves as a 30-day depot. As a LHRH antagonist, degarelix acts at the level the pituitary to rapidly reduce LH secretion, thereby decreasing testosterone production within the testes. A rapid reduction in circulating total testosterone to ≤ 50 ng/dl is achieved in the majority of patients within 2–3 days. Testosterone production recovers after the 30-day depot is cleared.

Study endpoints
Primary, secondary, and exploratory endpoints are listed in Table 3. Briefly, the primary endpoint is a composite of mortality, need for ongoing hospitalization, or requirement for mechanical ventilation (including extracorporeal membrane oxygenation) at day 15 after randomization. This endpoint was selected because it is a clinically meaningful outcome for hospitalized patients. The secondary endpoints include time to clinical improvement, inpatient mortality, length of hospitalization, duration of intubation for mechanical ventilation, time to achieve a normal temperature, and the maximum severity of COVID-19 illness. Exploratory endpoints assess the association of D-dimer, IL-6, LDH, ferritin, total WBC, absolute neutrophil count, absolute lymphocyte count, testosterone, viral load (in nasopharyngeal swabs and blood), and various demographic and concurrent

Table 1 7-category ordinal scale of clinical status of hospitalized influenza patients

| Category | Description |
|----------|-------------|
| 1        | Not hospitalized with resumption of normal activities. |
| 2        | Not hospitalized, but unable to resume normal activities |
| 3        | Hospitalization, not requiring supplemental oxygen |
| 4        | Hospitalization, requiring supplemental oxygen |
| 5        | Hospitalization, requiring nasal high-flow oxygen therapy and/or non-invasive mechanical ventilation |
| 6        | Hospitalization, requiring extracorporeal membrane oxygenation and/or invasive mechanical ventilation |
| 7        | Death |

Fig. 2 SPIRIT figure. Asterisk indicates if patient is still hospitalized
Table 2 Inclusion and exclusion criteria

Inclusion criteria
1. Male Veterans admitted to a VA hospital.
2. Age ≥ 18
3. Hospitalized on an acute care ward with a diagnosis of COVID-19 contributing to hospitalization.
4. Positive RT-PCR assay for SARS-CoV-2 on a nasopharyngeal swab sample.
5. Severity of illness of level 3, 4, or 5 on the influenza severity scale at the time of randomization.
6. The subject (or legally acceptable representative if applicable) must provide written informed consent for the trial.

Exclusion criteria
1. History of severe hypersensitivity to degarelix or any component of their respective formulation
2. History of congenital long QT syndrome or known history of prolonged QT interval OR Fridericia correction formula (QTcF) > 500 ms on electrocardiogram performed at screening.
3. Planned discharge within 24 h of treatment initiation.
4. Subject is planning to conceive or father children within the projected duration of the study, starting with the screening visit through 120 days after the last dose of study treatment.
5. Ongoing usage of a class IA or class III antiarrhythmic agent. At least 5 half-lives must elapse since any prior use of a class IA or III antiarrhythmic agent prior to administration of study drug.
6. Baseline electrolyte abnormalities of grade 3 or higher (based on CTCAE v5.0 criteria). Patients may be included if baseline electrolyte abnormalities are corrected to grade 2 or lower prior to study drug administration.
7. Myocardial infarction in the past 6 months, severe or unstable angina, or New York Heart Association (NYHA) class III or IV heart disease.
8. Known psychiatric or substance abuse disorder that would interfere with the requirements of the trial.
9. Child-Pugh Class C liver disease.
10. Use of any of the following hormonal agents:
   - Androgen receptor antagonists or agonists within 4 weeks of study enrollment,
   - Ketoconazole or abiraterone acetate within 2 weeks of study enrollment,
   - Estrogens or progestins within 2 weeks of study enrollment,
   - Herbal products that contain hormonally active agents within 2 weeks of study enrollment.
   - Any prior use of an LHRH analog unless a serum total testosterone measured within 30 days of study enrollment is ≥ 150 ng/dL.
   - Other hormonal agents listed in Appendix B within one day of study enrollment.
11. Unwilling or unable to comply with the study protocol.
12. Any condition, which in the opinion of the investigator, would preclude participation in the trial.

comorbidities with outcome. Additionally, for patients enrolled at the Greater Los Angeles, pharmacologic suppression of TMPRSS2 expression in the nasopharynx is also assessed via nasal swab, and the presence of antiviral antibodies in blood are assessed over time.

Safety evaluations/monitoring
Safety monitoring is conducted at scheduled study time points and through spontaneous reports from participants to evaluate adverse events (AEs). Adverse events of special interest include cardiac arrhythmias and thromboembolic events that may result from androgen suppression. Specifically, cardiac arrhythmias and thromboembolic complications of grades 3–5 (CTCAE v5.0). Non-serious adverse events related to the study intervention are reported to the sponsor through the electronic data capture (EDC) system. Expedited reporting of AEs of special interest (grades 3–5) and serious adverse events (SAEs) undergo daily review and reports generated for regular planned teleconference calls among the study investigators. Additionally, the VA Clinical Sciences Research and Development (CSR&D) centralized Data Monitoring Committee (DMC) will receive and review study reports monthly. Predetermined toxicity thresholds have been established as quality tolerance limits for statistical analysis. Stopping rules are applied if unbalanced toxicity signals are detected in the active arm at an α-level of 0.01.

Statistical considerations
We assumed an effect size for the primary endpoint of 42%. A total of 186 evaluable patients (i.e., 124 evaluable patients in the degarelix group and 62 evaluable patients in the placebo group) have 90% power of detecting the expected effect size using a two-sided two proportion test with a significance level of 0.05. Based on an assumed 5% attrition rate, 198 patients will be required (i.e., 132 in the degarelix group and 66 in the placebo group). The primary analysis of the study will be performed on the primary endpoint on the intention to treat population. Secondary endpoint analyses will be adjusted for multiplicity with a α-level of 0.0083 for each endpoint. The endpoints and specific statistical methods used for each are listed in Table 3. A mid-term interim analysis of the primary endpoint is planned after half of the patients complete the study. If the mid-term analysis of the primary endpoint indicates that the null hypothesis can be rejected with a boundary value of 2.77 at an α-level of 0.006 or accepted with a boundary value of 0.44 based on O’Brien and Fleming criteria, the study will be recommended for trial termination for efficacy or futility. Additionally, once the toxicity markers reach a threshold of 25%, a statistical analysis of the toxicity will be performed. If the one group has a significantly higher rate of toxicity compared to the placebo group at an α-level of 0.01, then results will be reported to the DMC for a recommendation for trial termination.
Table 3 Endpoints and data analysis plan

| Endpoint analyses                                                                 | Statistical methods                          | SAS procedures |
|----------------------------------------------------------------------------------|----------------------------------------------|----------------|
| **Primary endpoint**                                                              |                                              |                |
| A composite endpoint of mortality, hospital stay rate, and ECMO or mechanical intubation at 15 days after randomization | Pearson chi-square tests or Fisher exact tests Logistic regression adjusted for age, hypertension, and COPD | PROC FREQ, PROC LOGISTIC |
| **Secondary endpoints**                                                           | 1. Log-rank test, Kaplan-Meier curves       | 1. PROC LIFETEST |
|                                                                                 | Cox regression adjusted for age, hypertension, and COPD | PROC PHREG |
|                                                                                 | Logistic regression adjusted for age, hypertension, and COPD | PROC FREQ |
|                                                                                 | Wilcoxon test; quantile regression           | PROC GENMOD |
|                                                                                 | Wilcoxon test; Quantile regression            | PROC NPAR1WAY |
|                                                                                 | Log-rank test, Kaplan-Meier curves           | PROC LIFETEST |
|                                                                                 | Cox regression adjusted for age, hypertension, and COPD | PROC PHREG |
|                                                                                 | Pearson chi-square test, Cochran–Armitage trend test | PROC FREQ |
|                                                                                 | Proportional odds logistic regression         | PROC LOGISTIC |
| **Exploratory analyses**                                                          | 1. TIME TO EVENT DATA                        | 1. PROC LIFETEST |
|                                                                                 | Log-rank tests, Kaplan-Meier curves, and Cox regressions | PROC PHREG |
|                                                                                 | Pearson chi-square tests and logistic regression | PROC LOGISTIC |
|                                                                                 | Interval data                                | PROC TTEST |
|                                                                                 | Student t/Wilcoxon tests, linear or quantile regressions | PROC NPAR1WAY |
|                                                                                 | Longitudinal data                            | PROC Mixed |
|                                                                                 | Mixed-effect model repeated measure          |                |
| **AE/SAE analyses**                                                               | For All AE/SAE                               | PROC FREQ, SAS Macros |
|                                                                                 | Incidence rate estimation and testing         |                |
|                                                                                 | Pearson chi-square tests or Fisher exact tests |                |
| **Other analyses**                                                                | 1. STUDENT t or Wilcoxon tests, Pearson chi-square or Fisher exact tests | 1. PROC TTEST, PROC FREQ |
|                                                                                 | Admission characteristics, Pearson chi-square test | PROC NPAR1WAY |
|                                                                                 | Admission status, Pearson chi-square test    | PROC TTEST |
|                                                                                 | Admission status, Pearson chi-square test    | PROC NPAR1WAY |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |
|                                                                                 | Admission status, Pearson chi-square test    | PROC PHREG |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |
|                                                                                 | Admission status, Pearson chi-square test    | PROC FREQ |

**Trial status**

The current protocol version number is 5.5, February 9, 2021. Enrollment began July 6, 2020. Estimated date of recruitment completion is July 6, 2021.

**Discussion**

Pre-clinical and correlative data in humans suggest that anti-androgenic therapies can reduce the expression of TMPRSS2 on lung epithelium. Accordingly, we hypothesized that therapeutic targeting of androgen receptor signaling will suppress viral infection and thereby ameliorate the severity of symptomatic COVID-19. The purpose of this trial is to determine if temporary androgen suppression improves the clinical outcomes of patients who are hospitalized to an acute care ward due to COVID-19.

Degarelix was selected among the numerous FDA-approved drugs that target the AR analogs due to its rapid effect on circulating testosterone, safety profile, and availability. In contrast, LHRH agonists (e.g., leupro-lide) achieve testosterone suppression over 2–3 weeks and induce a transient increase in serum testosterone, which could exacerbate the severity of the infection. The potent androgen receptor antagonists (e.g., enzalutamide and apalutamide) achieve steady state in the serum over approximately 4 weeks. Of all the FDA-approved agents that inhibit AR signaling, degarelix exhibits the most rapid and robust effects. Degarelix has over a decade of post-marketing safety information collected from its use in patients with prostate cancer. Documented side effects are often predictable in terms of their onset, duration, and severity. These side effects include fatigue, hot flashes, reduced libido, reduced fertility, and result directly from testosterone suppression. These can reduce quality of life but are not life threatening and are reversed upon testosterone recovery. Most of the more severe side effects associated with androgen suppression occur after long term use. These include potential worsening of preexisting cardiovascular disease, reduced bone mineral density, loss of lean muscle mass, hypercoagulability, and prolongation of the QT/QTc interval. Most patients are expected to recover testosterone within a few months after the single depot of degarelix, limiting the risks associated with long term androgen suppression. For the purposes of safety monitoring, we consider any grade 3 or higher CTCAEv5 toxicities...
associated with hypercoagulation or arrhythmias to be of special interest given the baseline elevated risks of hypercoagulability in hospitalized patients with COVID-19 and the risks of torsade de pointes associated with COVID-19 QT/QTc intervals. We also chose to exclude patients with congenital long QT syndrome or known history of prolonged QT interval and require a screening electrocardiogram for QT interval corrected by the Fridericia correction formula (QTcF) > 500 ms.

Pharmacologic inhibition of TMPRSS2 with camostat mesilate prevents SARS-CoV-2 entry into cultured human lung cells [1]. However, a randomized trial of camostat mesilate versus placebo in patients hospitalized for COVID19 demonstrated no clinical improvement, ICU admission, or mortality [13].

In addition to our group, others are testing AR antagonism as a therapeutic strategy for hospitalized patients with COVID-19. The COVIDENZA trial led by the University of Gothenburg is an ongoing randomized trial of enzalutamide plus standard of care versus standard of care in hospitalized patients with COVID-19 [14]. A trial in the outpatient setting recently reported that androgen treatment accelerated viral clearance as compared to placebo [15]. Whether or not AR antagonism should be considered as a therapeutic strategy for patients hospitalized with COVID-19 will depend on the results of this and related trials.

**Abbreviations**

ACE2: Angiotensin-converting enzyme 2; AR: Androgen receptor; COPD: Chronic obstructive pulmonary disease; CSRD: Clinical Sciences Research and Development; DMC: Data Monitoring Committee; ECMO: Extracorporeal membrane oxygenation; ER: Estrogen receptor; FDA: Food and Drug Administration; GrnRH: Gonadotropin-releasing hormone; LDH: Lactate dehydrogenase; LHRH: Luteinizing hormone-releasing hormone; SARS-CoV-1: Severe acute respiratory syndrome coronavirus 1; SARS-CoV-2: Severe acute respiratory syndrome coronavirus 2; Spike protein; TMPRSS2: Transmembrane protease, serine 2; WBC: White blood cell count

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**Authors’ contributions**

MR, NN, XW, MM, and DG developed the concept. MR and NN wrote the trial with assistance from all other authors. ZM developed the statistical plan. The author(s) read and approved the final manuscript.

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**Availability of data and materials**

Data sharing is not applicable to this protocol manuscript as no datasets were generated or analyzed.

**Declarations**

**Ethics approval and consent to participate**

This trial was approved by the VA Central IRB on 5/1/2020 (VA Central IRB Study 2020-011). This manuscript does not contain any individual persons' data.

**Consent for publication**

The authors declare that they have no competing interests. There are no financial and non-financial competing interests to disclose related to this article.

**Competing interests**

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