

RECOVER-AUTONOMIC: A Platform Protocol for Evaluation of Interventions for Autonomic Dysfunction in Post-Acute Sequelae of SARS-CoV-2 Infection (PASC)

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STATEMENT OF COMPLIANCE

The trial will be carried out in accordance with International Conference on Harmonization Good Clinical Practice (ICH GCP) and the following:

• United States (US) Code of Federal Regulations (CFR) applicable to clinical studies (45 CFR Part 46, 21 CFR Part 50, 21 CFR Part 56, 21 CFR Part 312, and/or 21 CFR Part 812)

National Institutes of Health (NIH)-funded investigators and clinical trial site staff who are responsible for the conduct, management, or oversight of NIH-funded clinical trials have completed Human Subjects Protection and ICH GCP Training.

The protocol, informed consent form(s), recruitment materials, and all participant materials will be submitted to the central Institutional Review Board (IRB) for review and approval. Approval of both the protocol and the consent form(s) must be obtained before any participant is enrolled. Any amendment to the protocol will require review and approval by the IRB before the changes are implemented to the study. In addition, all changes to the consent form will be IRB-approved; a determination will be made regarding whether a new consent needs to be obtained from participants who provided consent, using a previously approved consent form.

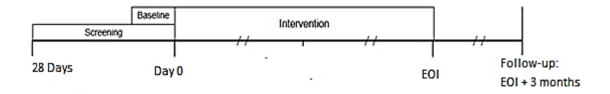
1.1 SYNOPSIS

	,				
Title:	RECOVER-AUTONOMIC: A Platform Protocol for Evaluation of				
	Interventions for Autonomic Dysfunction in Post-Acute Sequelae of SARS-				
	CoV-2 Infection (PASC)				
Study Description:	This study is a platform protocol designed to be flexible so that it is suitable for a wide range of settings within health care systems and in community settings where it can be integrated into COVID-19 programs and subsequent treatment plans.				
	This protocol is a prospective, multi-center, multi-arm, randomized, controlled platform trial evaluating various interventions for use in the treatment of autonomic dysfunction symptoms, including cardiovascular complications and postural orthostatic tachycardia syndrome (POTS), in PASC participants. The interventions tested will include non-pharmacologic care and pharmacologic therapies with study drugs.				
	The hypothesis is that some of the autonomic dysfunction symptoms are immune-mediated, so immunotherapy and other applicable therapies will result in improvement in autonomic symptoms.				
Objectives:	Primary:				
	Evaluate the effect of study intervention versus control on orthostatic intolerance				

	Secondary:		
	Evaluate the effect of study intervention versus control on symptom-specific outcome measures		
	 3. Describe the effect of study intervention versus control on performance-based and vital sign outcome measures 4. Evaluate the effect of study intervention versus control on quality of life 		
	 Compare the effect of study intervention versus control on specifi tracked measurements 		
	Characterize the safety and tolerability of study intervention for treatment of PASC		
	Exploratory:		
	7. Evaluate the effect of study intervention versus control on autonomic function tests		
	Evaluate the effect of study intervention versus control on relevant biomarkers		
Study Population:	Each study appendix will enroll approximately 180-200 evaluable adult participants who experience autonomic dysfunction symptoms of PASC. This is expected to be sufficient to conclude whether there is a meaningful evidence of intervention benefit. Overall enrollment will depend on the number of screen failures, the number of study drug appendices that are added, the ability to pool their control groups for analysis, and adjustments to sample size based on-study data.		
	The goal is to have a diverse population, including the underserved communities and racial/ethnic populations frequently underrepresented in clinical research.		
Phase:	Phase 2		
Description of	Participants will be recruited from various acute COVID-19 trials and		
Sites/Facilities Enrolling			
Participants:	communities. Approximately up to 75 sites in the US will participate.		
Description of Study Intervention:	Each study appendix describes a different intervention and control.		
Participant Duration:	Duration of intervention will vary depending on the specific study		
	appendix. Participants will have one follow-up visit approximately 3		
	months after the end of intervention.		

1.2 SCHEMA

Figure 1. Platform Protocol Study Schema



1.3 KEY ROLES

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2 INTRODUCTION

2.1 STUDY RATIONALE

Post-Acute Sequelae of SARS-CoV-2 infection (PASC), also known as Long COVID, is a chronic condition present in up to 80% of SARS-CoV-2-infected, hospitalized patients and 40% to 70% of non-hospitalized patients with the novel coronavirus disease 2019 (COVID-19).¹⁻⁴ The personal and global impact of these long-term symptoms from SARS-CoV-2 infection can be debilitating, and the number of PASC patients is escalating. With the increasing number of people infected with SARS-CoV-2, an urgent and unmet clinical need exists to better understand the pathophysiology of PASC and to develop targeted therapeutics to resolve the disease more rapidly and restore patients' health.

This study aims to investigate various interventions for autonomic symptoms in PASC. If successful, this trial will rapidly provide the foundational evidence to enable providers to identify and treat and ameliorate symptoms in PASC patients.

2.2 BACKGROUND

In 2019, COVID-19 emerged in Wuhan, China. A month later, the Chinese Center for Disease Control and Prevention identified a new beta-coronavirus (SARS coronavirus 2, or SARS-CoV-2) as the etiological agent.⁵ The clinical manifestations of COVID-19 range from asymptomatic infection or mild, transient symptoms to severe viral pneumonia with respiratory failure, and other serious, life-threatening complications.

COVID-19 has led to the death of more than 6 million people worldwide; however, this disease has affected even more lives through often-debilitating symptoms lingering long after acute SARS-CoV-2 infection. Post-Acute Sequelae of SARS-CoV-2 infection (PASC), also known as PASC, is a chronic condition present in up to 80% of infected, hospitalized patients and 40% to 70% of non-hospitalized patients. PASC affects nearly every organ system, with more than 200 individual symptoms, ranging from new-onset anxiety, depression, and cognitive difficulties to shortness of breath, dizziness, and arrhythmias. Moreover, PASC can occur regardless of severity of acute COVID-19, and it impacts across socioeconomic, racial and ethnic, and age strata. These prolonged symptoms open the door for substantial short- and long-term individual and societal costs, including healthcare costs and inability to work. Prolonged symptoms have kept individuals out of work, which has exacerbated poverty in the underserved, historically minoritized populations and worsening a decades-long mental health crisis. Considering these costs, identification of safe and effective methods to treat and prevent the occurrence of PASC represents an urgent, unmet public health need.

To address this need, the NIH has launched the RECOVER initiative across the nation (RECOVER: Researching COVID to Enhance Recovery) to better understand the disease. The RECOVER Initiative brings together patients, caregivers, clinicians, community leaders, and scientists from across the nation to understand, prevent, and treat PASC. The RECOVER Consortium represents and supports researchers who are leading studies on PASC at more than 200 sites around the country. These studies have a diverse group of participants, including adults, pregnant people, and children. Data from the RECOVER initiative, as well as existing literature, have highlighted 3 predominant symptom clusters, including exercise intolerance, cognitive dysfunction, and autonomic dysfunction that are frequently reported but also of substantial importance to patients. In addition, data have identified substantial heterogeneity in symptomology and presentation, even among those within a specific symptom cluster; and the lack of concordance between objective findings and reported symptoms in many cases.^{7,8}

Autonomic multisystem dysfunction including cardiovascular dysregulation, and postural tachycardia syndrome (POTS), are often seen in PASC patients. Postural orthostatic tachycardia syndrome (POTS) is an abnormal increase in heart rate (HR) that occurs when transitioning from lying down to standing up. Among COVID-19 survivors, 9% to 16% experience POTS-like symptoms, such as tachycardia, orthostatic intolerance, fatigue, and cognitive impairment, and 2% to 14% develop POTS.⁹

Immune therapies, such as intravenous immunoglobulin (IVIG) have been used to treat several viral infections. A substantial proportion of POTS is thought to be immune-mediated. A number of case reports show improvement with immune modulating therapy, particularly intravenous immunoglobulin (IVIG), in patients with various post-COVID conditions.^{10,11}

Medications that are used to treat chronic heart failure with elevated HR are also potential treatment options for improving tachycardia symptoms in patients with PASC and POTS.

Apart from pharmacologic therapy, the current foundation of treatment for POTS, and for POTS in patients with PASC, is non-pharmacologic therapy, including volume expansion, behavioral and rehabilitation therapy. Specifically, based on small studies and pathophysiologic rationale, patients are treated with volume expansion with encouragement of fluid and salt intake, abdominal binders, and physical therapy directed at orthostatic symptoms. An important question is whether such therapy, if applied in a systematic and intense way with the assistance of a care coordinator, would be more effective than usual care in which these treatments may be recommended but without assurance of adherence. Thus, there is interest to assess the effect of coordinated non-pharmacologic, behavioral and rehabilitation care versus usual care.

Evidence in the current literature indicates that non-pharmacologic therapy in combination with immune therapies and cardiac modulators have promise in treatment of PASC-related autonomic symptoms. Randomized trial evidence is needed to determine if these treatments provide meaningful benefit. If successful, this trial program will rapidly provide the foundational evidence to enable providers to identify and treat patients with PASC and POTS to ameliorate these symptoms.

2.3 RISK/BENEFIT ASSESSMENT

2.3.1 KNOWN POTENTIAL RISKS

Potential risks of this study include those associated with the specific study intervention (refer to Appendices for details), blood draws, nasal swabs, performance measures, and loss of confidentiality.

Risks associated with blood draws include momentary discomfort and/or bruising. Infection, excess bleeding, clotting, or fainting are also possible, although unlikely.

Risks associated with nasal swabs include mild irritation, insignificant local pain, and minor bleeding.

Risks associated with the 6-minute walk test include tiredness, shortness of breath, or palpitations during the test. The active stand test and full autonomic testing may cause some participants to faint or feel lightheaded or weak.

The following risks are associated with the autonomic function tests:

- The head-up tilt (HUT) test may cause lightheadedness and fainting due to a drop in blood pressure. However, there is no risk of falling from the table as the subject is restrained by heavy straps and the table will be lowered horizontally to relieve any dizziness.
- The Valsalva maneuver may lead to chest pain, hypotension, fainting, and abnormal heart rhythms.
- Deep breathing test may lead to shortness of breath and dizziness
- Skin biopsy is a minimally invasive procedure and is a generally safe procedure. Risks of obtaining a tissue biopsy are bleeding, bruising, scarring, soreness, redness, and infection at the tissue punch

biopsy site. For the procedure, lidocaine with epinephrine is administered subcutaneously at the biopsy site. At this dose, there is minimal risk of systemic toxicity. Potential risks include allergic reactions and anaphylactic shock. The transcranial doppler is a form of non-invasive medical imaging. The risks associated with it are minimal and include discomfort or pressure from the probe.

There is also a risk of loss of confidentiality. However, coding all participant data with a unique identification number will minimize risk to loss of participant confidentiality.

2.3.2 KNOWN POTENTIAL BENEFITS

Participants may benefit from improvement in PASC symptoms.

Society in general and future patients infected with SARS-CoV-2 may benefit from the study's results, which will provide a better understanding of the benefit/risk of non-pharmacologic and pharmacologic treatment options for autonomic dysfunction symptoms, including POTS, in PASC patients.

2.3.3 ASSESSMENT OF POTENTIAL RISKS AND BENEFITS

Potential benefits may outweigh potential risks. Autonomic dysfunction in PASC is a significant health issue and can have consequences that impact quality of life. These consequences are extremely important for the individual and also impact the society at large.

3 OBJECTIVES AND ENDPOINTS

The objectives, outcome measures, and endpoints for the study are listed in Table 1, below. Further details on outcome measures and endpoints are provided in Section Error! Reference source not found. and the Statistical Analysis Plan (SAP).

While the primary outcome is OHQ/OIQ, the overarching goal is to determine if the interventions result in improved symptoms and quality of life. This will be evaluated by a number of questionnaires including COMPASS-31 and PROMIS-29, as well as tests of functional status such as 6-minute walk test. As a phase 2 trial, the total evidence from the primary and secondary outcomes will provide the most complete assessment of the impact of the interventions.

Table 1. Summary of study objectives, endpoints, and outcome measures.

OBJECTIVES	OUTCOME MEASURES	ENDPOINTS		
Primary				

OBJECTIVES	OUTCOME MEASURES	ENDPOINTS	
Evaluate the effect of study intervention versus control on orthostatic intolerance	Orthostatic Hypotension Questionnaire (OHQ)/ Orthostatic Intolerance Questionnaire (OIQ)	Change from baseline to end of intervention (EOI)	
Secondary			
Evaluate the effect of study intervention versus control on symptom-specific outcome measures	Composite Autonomic Symptoms Score 31 (COMPASS-31) Malmo POTS Symptom Score (MAPS)	Change from baseline to EOI	
Describe the effect of study intervention versus control on performance-based and vital sign outcome measures	Active Stand Test Blood pressure (BP) and heart rate (HR) measurements 6-min Walk Test	Change from baseline to EOI	
Evaluate the effect of study intervention versus control on quality of life	PROMIS-29 + 2 Questionnaire	Change from baseline to EOI	
Compare the effect of study intervention versus control on specific tracked measurements	Wearable device measuring accelerometry (step count and HR)	Change from baseline to EOI	
Characterize the safety and tolerability of study intervention for treatment of PASC	Adverse events, including serious adverse events (SAEs) and events of special interest (ESIs)	Proportion of participants who experience individual SAEs and the proportion who experience any one or more SAEs Incidence of SAEs leading to discontinuation	

OBJECTIVES	OUTCOME MEASURES	ENDPOINTS	
		Incidence of ESIs	
Exploratory			
Evaluate the effect of study intervention versus control on autonomic function tests	Autonomic function tests at specialized sites: • Head-up tilt (HUT) test • Valsalva maneuver with a pressure of 40 mmHg for 15 seconds • Deep breathing test • Skin biopsy (if applicable as per Appendix) • Transcranial doppler (TCD), if available	Change from baseline to EOI	
Describe the effect of study intervention versus control on performance-based symptom measures	Vanderbilt Orthostatic Symptoms Score (VOSS)	Change from baseline to EOI	
Describe the effect of study intervention versus control on PASC symptoms	PASC Symptom Questionnaire	Change from baseline to EOI	

OBJECTIVES	OUTCOME MEASURES	ENDPOINTS
Evaluate the effect of study intervention versus control on relevant biomarkers	 Viral biomarkers Autonomic function biomarkers Immune biomarkers Inflammatory biomarkers Endothelial function biomarkers Cognitive function (CCL11) 	Change in biomarker(s) from baseline to EOI

4 STUDY DESIGN

4.1 PLATFORM PROTOCOL DESIGN

This platform protocol is a prospective, multi-center, multi-arm, randomized, controlled trial evaluating treatment of autonomic dysfunction PASC symptoms in outpatients previously infected with SARS-CoV-2. This study is a platform protocol designed to be flexible so that it is suitable for a wide range of settings within healthcare systems and in community settings where it can be integrated into COVID-19 testing programs and subsequent treatment plans. The platform protocol will enroll participants meeting criteria for autonomic dysfunction symptoms. Each appendix will describe a study intervention that is sized to meet the platform protocol objectives. Participants will be randomized to pre-specified factorial combinations of a study drug and non-pharmacologic intervention along with their controls that are actively enrolling at the time of randomization. Interventions may be added or removed according to adaptive design and/or emerging evidence. Refer to the protocol Appendices for further information on each study intervention and study design.

Patient-reported outcomes (PROs) of symptoms and functional status and objective assessments will be done. Periodic clinic visits will be done during the intervention period and a follow-up assessment will be done 3 months after EOI.

4.2 SCIENTIFIC RATIONALE FOR STUDY DESIGN

A randomized trial design is used to compare the intervention to control under a platform protocol. In addition, where applicable, double-blind will be maintained. A factorial design is used in each study intervention in combination with a non-pharmacologic care, where the non-pharmacologic care is not blinded due to its practical implementation.

Evidence in the current literature indicates that non-pharmacologic therapy and medications have promise in treatment of PASC-related cardiovascular symptoms. Randomized trial evidence is needed to determine if these treatments provide meaningful benefit.

Additional pharmacologic therapies can be added to this platform protocol as new drugs are available or as the understanding of PASC is improved and favorable drug properties are discovered.

Apart from pharmacologic therapy, the current foundation of treatment for POTS, is non-pharmacologic therapy, including volume expansion, behavioral and rehabilitation therapy. An important question is whether such therapy, if applied in a systematic and intense way with the assistance of a care coordinator, would be more effective than usual care in which these treatments may be recommended but without assurance of adherence. Thus, this study design was selected to determine the effect of coordinated non-pharmacologic, behavioral and rehabilitation care versus usual care. It may be worthwhile to also explore the interaction effect of non-pharmacologic treatment concurrently with pharmacologic therapy.

4.3 JUSTIFICATION FOR DOSE

The duration (and dose if applicable) of the intervention will vary depending on the intervention. Refer to the appendices for details.

4.4 END OF STUDY DEFINITION

The End of Study (EOS) will occur when all participants have completed EOI and Follow-up visit.

5 STUDY POPULATION

5.1 INCLUSION CRITERIA

In order to be eligible to participate in this study, an individual must meet all of the following criteria:

- 1. \geq 18 years of age at the time of enrollment
- 2. Previous suspected, probable, or confirmed SARS-CoV-2 infection, as defined by the Pan American Health Organization^{12¢}
 - ^{\$\phi\$} Enrollment of participants with suspected or probable SARS-CoV-2 infection will only be allowed if they occurred before May 1, 2021 and be limited to no more than 10% of the total sample size per Study Drug Appendix. Refer to the Manual of Procedures (MOP) for details.

Suspected case of SARS-CoV-2 infection - Three options, A through C:

- A. Meets the clinical OR epidemiological criteria.
 - a. Clinical criteria: Acute onset of fever AND cough (influenza-like illness) OR Acute onset of ANY THREE OR MORE of the following signs or symptoms: fever, cough, general, weakness/fatigue, headache, myalgia, sore throat, coryza, dyspnea, nausea, diarrhea, anorexia.
 - Epidemiological criteria: Contact of a probable or confirmed case or linked to a COVID-19 cluster; or
- B. Presents with acute respiratory infection with history of fever or measured fever of ≥ 38°C; and cough; with onset within the last 10 days; and who requires hospitalization); or
- C. Presents with no clinical signs or symptoms, NOR meeting epidemiologic criteria with a positive professional use or self-test SARS-CoV-2 antigen-Rapid Diagnostic Test.

Probable case of SARS-CoV-2 infection, defined as meets clinical criteria above AND is a contact of a probable or confirmed case or is linked to a COVID-19 cluster.

Confirmed case of SARS-CoV-2 infection - Two options, A through B:

- A. A person with a positive nucleic acid amplification test, regardless of clinical criteria OR epidemiological criteria; or
- B. Meeting clinical criteria AND/OR epidemiological criteria (See suspect case A). With a positive professional use or self-test SARS-CoV-2 Antigen-Rapid Diagnostic Test.
- 3. Moderate, self-identified autonomic symptoms (defined as COMPASS-31 >25) following a SARS-CoV-2 infection that has persisted for at least 12 weeks and is still present at the time of consent
- 4. OHQ/OIQ, question 1 score >2

5.2 EXCLUSION CRITERIA

An individual who meets any of the following criteria will be excluded from participation in this study:

- 1. Known pregnancy, breast-feeding, or contemplating pregnancy during the study period
- 2. Known active acute SARS-CoV-2 infection ≤ 4 weeks from enrollment
- 3. Known renal failure (eGFR <20ml/1.73 m²)
- 4. Known atrial fibrillation or significant cardiac arrhythmia
- 5. Known cardiovascular conditions such as heart failure (Class 3-4), severe valvular disease, symptomatic ischemic coronary artery disease, revascularization for PAD/CAD within the past 6 months
- 6. Clinically significant atherosclerotic disease, defined as history of stroke or myocardial infarction or revascularization 6 months prior to enrollment and/or current symptomatic angina
- 7. Existing uncontrolled hypertension
- 8. History of significant hypercoagulability disorders
- 9. Active or recent thrombosis

- 10. Inability to comply with the protocol
- 11. Less than 2 weeks since discontinuation/weaning of beta-blockers and calcium channel blockers
- 12. Any condition that the investigator believes makes it inappropriate to treat with study intervention, including severe debility.
- 13. History of POTS or significant dysautonomia before COVID-19 index infection.
- 14. Enrolled in another clinical trial or another study intervention appendix in this platform protocol *Participants may enroll in the trial or re-enroll in the trial for a different study intervention appendix if they have completed a 30-day washout period. For those who wish to re-enroll, re-enrollment can only occur after efficacy has been determined for the appendix in which they were previously enrolled.

Exclusion criteria specific to each intervention are listed in the Appendices.

5.3 LIFESTYLE CONSIDERATIONS

Participants of child-bearing age must agree to use an effective method of contraception during study drug administration and for at least 7 days after their final dose of study drug. Effective methods include any of the following: abstinence, partner vasectomy, bilateral tubal ligation, intrauterine device, progestin implants, or barrier (condom, diaphragm, cervical cap) plus spermicide.

5.4 SCREEN FAILURES

Screen failures are defined as participants who consent to participate in the clinical trial but are not subsequently randomly assigned to a study intervention or entered in the study. A minimal set of screen failure information is required to ensure transparent reporting of screen failure participants, to meet the Consolidated Standards of Reporting Trials (CONSORT) publishing requirements and to respond to queries from regulatory authorities. Minimal information includes demography, screen failure details, and eligibility criteria.

5.5 STUDY DEFINITION OF ENROLLMENT

For this study, enrollment is defined as signing consent and completing randomization.

5.6 STRATEGIES FOR RECRUITMENT AND RETENTION

The RECOVER Clinical Trial Data Coordinating Center (CT-DCC) will use an integrated strategy of coordinating with community organizations, the public, and clinical trial sites to identify and retain study participants. To ensure a diverse population is enrolled, strategies from prior successful initiatives will be refined and utilized. The study team will develop a comprehensive communication strategy involving print and social media, as well as leveraging existing organizational structures where possible, to educate the public on concerns about PASC and opportunities for clinical trial participation. Interested

members of the public will be provided with information to contact a local site for potential participation.

Participants can be recruited and identified through participating site outreach. Site investigators, or their designee, may contact eligible participants to introduce the study and discuss study participation.

Participants may be recruited from other ongoing COVID-19 trials if they opted-in to be contacted about future research opportunities. To support participant referral to actively enrolling trials, a series of invitation algorithms based on appendix-specific inclusion/exclusion criteria and participant entered data may be used. Automatic invitations will be generated for participants who appear eligible based on trial interest, demographics, and medical history. Once participants accept the invitation and adequate consent is obtained, their information will be shared with the applicable study team.

Patient advocates that represent a diverse PASC community will be engaged in the study at every step. Patient advocates will serve as consultants to inform study design, protocol development, and recruitment and retention strategies.

During active study, study sites will maintain close connections with study participants.

6 STUDY INTERVENTION

6.1 STUDY INTERVENTION(S) ADMINISTRATION

6.1.1 STUDY INTERVENTION DESCRIPTION

See appendices for pharmacologic interventions and Section 6.6 for non-pharmacologic intervention.

6.1.2 DOSING AND ADMINISTRATION

See appendices.

6.2 PREPARATION/HANDLING/STORAGE/ACCOUNTABILITY

6.2.1 ACQUISITION AND ACCOUNTABILITY

Receipt and use of any pharmacologic study intervention (i.e. study drug) will be handled, tracked by the sites, and stored in a safe and secure location to which only the investigator and designated personnel have access. Use of the study drug will be tracked by the sites. When the study intervention is a pill or tablet, the participants will be asked to bring the packaging to every clinic visit and sites will monitor adherence through pill/tablet count. Participants will also be asked to bring any unused study drug with them to the clinic during their EOI visit. Sites may also contact the participants at regular intervals during the study intervention period to monitor compliance.

See appendices for further details.

6.2.2 FORMULATION, APPEARANCE, PACKAGING, AND LABELING

See appendices.

6.2.3 PRODUCT STORAGE AND STABILITY

As applicable, upon receipt, study intervention should be stored according to the instructions specified on the label. See appendices.

6.2.4 PREPARATION

See Appendices.

6.3 MEASURES TO MINIMIZE BIAS: RANDOMIZATION AND BLINDING

Since the route and duration of administration of each study intervention differs, the controls also differ. To achieve blinding and an equitable randomization probability, the following process will be used.

In each Appendix trial, each participant will be assigned with equal probability to one of the factorial combinations based on 2 factors: (1) a study intervention/control and (2) non-pharmacologic intervention/control if the participant is eligible for the study intervention. If more study drugs are added with the same administration route and the same inclusion and exclusion criteria with a shared control, the level of the study intervention factor may be m+1 level where m is the number of active study interventions along with one shared control. In that case, the study design will be an (m+1) x 2 factorial design.

Sites will be informed to which appendix participants are randomized, but not whether they are allocated to the active study intervention arm or control arm within that appendix. The participants and investigators will be blinded throughout the study for the study intervention, but not for the non-pharmacologic intervention. Due to the nature of non-pharmacologic intervention, blinding is not possible.

6.3.1 UNBLINDING

The participant, treating clinicians, and study personnel will remain blinded to study intervention versus control assignment until after the database is locked and blinded analysis is completed. Only the biostatistical team who is preparing closed interim reports will be unblinded. Specifically, pharmacologic intervention/control will be dispensed with packaging and labeling that would blind treatment assignment. Unblinding will occur only if required for participant safety or treatment, at the request of the treating clinician.

Refer to the MOP for further details.

6.4 STUDY INTERVENTION ADHERENCE

Participants will be notified of the importance of completing all aspects of their intervention.

6.5 CONCOMITANT THERAPY

For this protocol, a prescription medication is defined as a medication that can be prescribed only by a properly authorized/licensed clinician. Medications to be reported in the Case Report Form (CRF) are relevant concomitant prescription medications, over-the-counter medications, and supplements. Baseline concomitant medications will include all concomitant therapies taken by the participant within 14 days of informed consent.

No medications or standard treatments for POTS will be withheld from study participants.

Refer to the Appendices for study-specific information on concomitant medications.

6.6 NON-PHARMACOLOGIC CARE

All participants will receive coordinated non-pharmacologic care or usual non-pharmacologic care (control) for a duration of 3 months. This will be concurrent to study drug intervention (refer to Appendices).

Coordinated non-pharmacologic care involves a systematic and intensified program of volume expansion through high salt diet (6-10 grams of salt per day), water intake (at least 2 liters [64 ounces] per day), abdominal binder, physical rehabilitation, weekly monitoring of BP, motivation, education, and assisted care through a care coordinator. The care-coordinators will be required to complete a training program, and details will be provided in the MOP.

All participants receiving coordinated non-pharmacologic care will receive a kit of equipment needed. Additional details are in the MOP.

The care coordinator will call participants in coordinated non-pharmacologic care arm weekly during the 3-month non-pharmacologic intervention period to provide education and support on lifestyle modifications. Participants will be asked to maintain a weekly log of non-pharmacological care (average dietary intake of salt and water, physical rehabilitation) and provide a summary to the care-coordinators during the weekly calls.

To abort lightheadedness episodes, participants will be encouraged to sleep with their head elevated (6-12 inches of elevation) and drink 16 ounces of water (about ½ liter) in the morning while in bed before standing. In addition, practicing physical countermeasures (such as leg crossing, buttock clenching,

squatting, squeezing a stress ball repeatedly, or clasping and pulling the fists apart) will be encouraged with the goal of further helping to prevent lightheadedness. Participants will also be informed to avoid triggering events such as prolonged standing in one position, hot environments (hot showers), sudden standing after waking in the morning, prolonged fasting, excessive alcohol intake and straining (during urination or bowel movements).

7 PARTICIPANT DISCONTINUATION/WITHDRAWAL

7.1 PARTICIPANT DISCONTINUATION FROM STUDY DRUG

Discontinuation from study intervention does not mean discontinuation from the study, and remaining study procedures should be completed as indicated by the study protocol. If a clinically significant finding is identified (such as changes from baseline) after enrollment, the investigator or qualified designee will determine if any change in participant management is needed.

An investigator may discontinue a participant from the study intervention at their discretion, for any reason including, but not limited to, one of the following:

- Significant study intervention non-compliance
- If any clinical AE, laboratory abnormality, or other medical condition or situation occurs such that continued participation in the study or study drug would not be in the best interest of the participant
- Confirmed new case of acute SARS-CoV-2

Participants will be followed for safety even if study intervention is discontinued. The reason for participant discontinuation from study intervention will be recorded on the CRF. Participants who are discontinued from the study intervention, but who are not withdrawn from the study will continue to be followed for all study procedures.

Participants who discontinue the study drug, but do not withdraw consent, will be followed for safety for at least 28 days after the final study drug administration. Participants will continue to receive non-pharmacologic care even if study drug intervention is discontinued. Participants may be replaced with additional randomized participants, but will not be specifically matched with the withdrawn participant.

7.2 PARTICIPANT WITHDRAWAL FROM THE STUDY

Participants are free to withdraw consent from participation in the study at any time upon request. The study team will attempt to determine a reason for withdrawal; however, participants are not obligated to provide a reason for withdrawal. If obtained, the reason for withdrawal will be recorded on the CRF. No further study procedures will be performed and no further data will be collected from the participant following study withdrawal. All of the data collected up until the time of consent withdrawal will be maintained in the study database and will be used as the participant's data are evaluable for analysis.

7.3 LOST TO FOLLOW-UP

Participants will be considered lost to follow-up if they fail to return for any scheduled visit *and* if they are unable to be contacted after multiple attempts and methods by the study site staff and/or a central search company.

The following actions must be taken if a participant fails to return to the clinic for a required study visit:

- The site will attempt to contact the participant and reschedule the missed visit and counsel the
 participant on the importance of maintaining the assigned visit schedule and ascertain if the
 participant wishes to and/or should continue in the study
- Before a participant is deemed lost to follow-up, the investigator or designee will make every effort to regain contact with the participant or next of kin (where possible, telephone calls and, if necessary, a certified letter to the participant's last known mailing address or local equivalent methods). These contact attempts should be documented in the participant's study file.
- Should the participant continue to be unreachable, after exhausting all methods, he or she will be considered to have withdrawn from the study with a primary reason of lost to follow-up.

7.4 HALTING AND STOPPING RULES

The sponsor has engaged Duke Clinical Research Institute (DCRI) Safety Surveillance to oversee real-time serious adverse event (SAE) collection, evaluation, and expedited regulatory reporting for this study. An independent DCRI Safety Medical Monitor will be responsible for evaluating causality assessment and expectedness of site reported SAEs compared to the product label. Details of this process can be found in the study-specific Safety Management Plan.

Participant safety will be reviewed by RECOVER Data and Safety Monitoring Board (DSMB) that is composed of individuals with the appropriate expertise. Members of the DSMB will be independent from the study conduct and free of conflict of interest, and measures will be in place to minimize perceived conflict of interest. The DSMB will meet as stated in their charter to approve protocols, assess safety and efficacy data, and at appropriate intervals to meet requirements for the Interim Analyses on each arm of the study. The DMSB will operate under the rules of an approved charter that will be written and reviewed at the organizational meeting of the DSMB. At this time, each data element that the DSMB needs to assess will be clearly defined. The DSMB will provide its input to the sponsor.

This DSMB may recommend that the study may be temporarily suspended or prematurely terminated if there is sufficient reasonable cause. Written notification, documenting the reason for recommendation of study suspension or termination, will be provided by the sponsor to study participants, site investigators, the central IRB, and the US Food and Drug Administration (FDA). Study participants will be contacted, as applicable, and be informed of changes to study visit schedule.

A participant's study drug will be permanently discontinued if:

- The participant requests to stop study drug
- The investigator believes that, for safety reasons or tolerability reasons, it is in the best interest of the participant to discontinue study drug

 The participant has a serious (i.e. SAE) or severe (Grade 3 or above) anaphylactic or allergic reaction that is determined to be related to the study medication
 Permanent discontinuation of study drug should be considered for patients with side effects possibly related to study drug such as thrombosis, worsening renal function, aseptic meningitis, hemolytic anemia, and transfusion related lung injury.

The overall trial would be paused, pending DSMB review for:

- Two confirmed anaphylactic reactions determined to be related to study drug
- Two episodes of thrombosis, including deep venous thrombosis and/or embolic events, determined to be related to study drug
- Two episodes of renal failure that is not otherwise explained by the subject's co-morbidities
- Two episodes of aseptic meningitis syndrome
- Two episodes of hemolytic anemia requiring transfusion
- Two episodes of transfusion related acute lung injury determined to be related to study drug
- Two deaths determined to be related to study drug
- Two of any other SUSARS

The overall trial will be stopped if:

 Sponsor determines that the study should be stopped based upon recommendations from DSMB and NIH (e.g. based on unblinded assessment of risk and benefit, including imbalance of serious or severe allergic reactions, anaphylaxis, deaths, and SUSARs).

Circumstances that may warrant termination or suspension may also include protocol compliance and data quality issues:

- Insufficient compliance to protocol requirements
- Data that are not sufficiently complete and/or evaluable

Please refer to the Appendices for any further specific halting and stopping rules.

Study may resume once concerns about safety, protocol compliance, and data quality are addressed, and satisfy the sponsor, IRB and/or FDA.

No formal stopping rules for efficacy will be implemented in this study because it is unlikely that there will be such a large benefit detected given the modest sample size and short duration. As above, study intervention may be suspended for serious safety concerns, after considering the balance of efficacy and safety. The DSMB recommendations will be considered by the sponsor, NIH, and the study Principal Investigator prior to making any decisions regarding study continuation or discontinuation.

No ESIs are recorded as part of the master protocol, but ESIs will be documented in each study appendix depending upon the study intervention (Refer to Appendices).

8 STUDY ASSESSMENTS AND PROCEDURES

8.1 SCHEDULE OF PROCEDURES

Table 2. Schedule of Study Procedures: Platform Protocol

	Screening	Baseline/ Randomization ⁶	Study Intervention Period ¹	End of Intervention (EOI) Clinic Visit	Early Termination Clinic Visit	Follow-up Visit
PROCEDURE	Day -28 to -7	Day -10 to -7	Day 0 to EOI	Refer to Appendix		EOI + 3 months (± 2 weeks)
Platform Protocol Eligibility Criteria	X					
Informed consent	X					
Demographics	X					
Medical History ⁹	X					
Concomitant Medications	X	X			X	
Randomization		X				
Coordinated Non- pharmacologic Care ²			Refer to Appendix-	Refer to Appendix-		
Appendix Eligibility Criteria	X	X	specific study procedures	specific study procedures		
Nasal Swab for SARS-CoV-2 rapid antigen test		X				
Clinic visits, as per specific study Appendix					X	X
OHQ/OIQ	X	X			X	X
COMPASS-31	X				X	X
Malmo POTS Symptom Score (MAPS)		X			X	X

Vanderbilt Orthostatic Symptoms Score	X			X	X
Active Stand Test	X			X	X
6-minute Walk Test	X			X	X
Modified Depaul Symptom Questionnaire- Post Exertional Malaise (DSQ- PEM) ³ (look back period of 7 days)	X	Refer to Appendix- specific study procedures	Refer to Appendix- specific study procedures		
DSQ-PEM ³ (look back period "since last visit")	X			X	X
PROMIS-29 + 2 Questionnaire	X			X	X
PASC Symptom Questionnaire	X			X	X
Blood Collection ⁴	X			X	X
Stool Collection	X			X	
Skin Biopsy ⁷	X			X	
Autonomic function tests at specialized sites	X			X	
Pregnancy Test	X				
Height and weight	X				
Study Drug Administration					
Distribution of wearable ⁸	X				

Safety Assessment of AEs including SAEs and ESIs	X		X	X
Adherence ⁵			X	

Abbreviations: PASC-Post-acute sequelae of SARS-CoV-2 infection, OHQ— Orthostatic hypotension questionnaire; OIQ-Orthostatic intolerance questionnaire; AE-Adverse Events; SAE — Serious Adverse Events; ESI — Events of Special Interest ¹A detailed schedule of activities for the Study Intervention Period is provided in each Appendix.

8.2 SCREENING (DAY -28 TO -7)

Information about the study will be presented to potential participants and questions will be asked to determine potential eligibility. Study-specific screening procedures will begin only after informed consent is obtained.

It is required to have completed the Screening prior to the Baseline/Randomization visit. Screening cannot be on the same day as the Baseline/Randomization visit. After the informed consent, the following assessments will be performed at screening to determine eligibility:

- Demographics
- Medical History; including SARS-CoV-2 test result date (if available), signs and symptoms, and treatment (including hospitalization, Intensive Care Unit (ICU) status, supplemental O₂ status), COVID-19 vaccination status and dates, and PASC history (symptoms and duration). At every clinic visit after screening visit, changes in medical history including SARS-CoV-2 infection since last visit will be collected.
- Collection of concomitant medications taken within 14 days of informed consent
- COMPASS-31

²Only applicable for those who are randomized to coordinated non-pharmacologic care. Not applicable to those receiving usual care. The coordinated non-pharmacologic care will be for a total duration of 3 months only. During this time, a care coordinator will reach out to the participants weekly.

³ The DSQ-PEM will be done twice for the baseline visit: once on the day of the visit with a look back period of "past 7 days" and once within one day after the visit with a look back period of "since last study visit (SLV)". DSQ-PEM SLV will be administered via phone or electronic survey. The DSQ-PEM administered at every timepoint after baseline will be within one day after strenuous in-person study visits and will ask about the period "SLV".

⁴Blood collection for safety parameters, biomarkers, and biospecimens for biorepository. Blood collection may also be collected as needed based on PI discretion and adverse events.

⁵Adherence to drug (pill count) will be documented. Participants receiving oral study drugs to be taken at home will receive weekly phone calls for the first 30 days. Tab

⁶It is required to have completed the Screening prior to the Baseline/Randomization visit. Screening cannot be on the same day as the Baseline/Randomization visit.

⁷Skin biopsy is only applicable to Appendix A: IVIG. If the baseline skin biopsy shows evidence of small fiber neuropathy, the biopsy will be repeated at EOI or ET.

⁸ It is required to wear the wearable 7 days before start of study drug intervention and 7 days before end of intervention visit. All other times, it is strongly encouraged.

⁹ At every clinic visit after screening visit, changes in medical history including SARS-CoV-2 infection since last visit will be collected.

- OHQ/OIQ
- Review appendix-level eligibility criteria, refer to Appendices

8.3 BASELINE/RANDOMIZATION (DAY -10 TO -7)

The following will occur at the Baseline/Randomization visit:

- Concomitant Medication review
- OHQ/OIQ
- MAPS
- VOSS
- Active Stand Test
- 6-minute walk test
- Modified Depaul Symptom Questionnaire-Post Exertional Malaise (DSQ-PEM)
 - At Baseline, this will be done twice: once on the day of the Baseline visit with a look back period of "past 7 days" and once the day after the Baseline study visit with a look back period of "since last study visit (SLV)". DSQ-PEM SLV will be administered via phone or electronic survey.
- Autonomic function tests, at specialized sites
- Skin biopsy will only be done in Appendix A: IVIG. If the baseline result shows evidence of small fiber neuropathy, the biopsy will be repeated at EOI or ET.
- PROMIS-29 + 2 Questionnaire
- PASC Symptom Questionnaire
- Nasal Swab for SARS-CoV-2 rapid antigen test
- Blood collection for the following evaluations:
 - Complete Blood Count (local, if not done within the past 3 months)
 - Comprehensive Metabolic Panel (local, if not done within the past 3 months)
 - Biomarkers that may include viral biomarkers, immune biomarkers, Inflammatory biomarkers, Endothelial function biomarkers, and biomarkers for autonomic and cognitive function). Refer to Section 8.9.12.
 - Biospecimens (serum, plasma, and stool) to be stored in biorepository (frozen for retrospective analysis). Refer to Section 8.10.
- Pregnancy test, urine in individuals of child-bearing potential
- Height and weight
- Distribution of wearable, it is required to wear the wearable 7 days before start of study drug
 intervention and 7 days before end of intervention visit. All other times, it is strongly
 encouraged.
- Randomization: 1:1:1:1 randomization to combination of study drug intervention/control with coordinated non-pharmacologic care/usual care

- Safety assessment of SAEs
- Review appendix-level eligibility criteria, refer to Appendices

8.4 STUDY INTERVENTION PERIOD (DAY 0 TO END OF INTERVENTION [EOI])

Refer to Appendices for study drug-specific intervention duration. The following will occur during the drug dosing period:

- Concomitant Medication review
- Study intervention administration, as per specific study Appendix
- Coordinated non-pharmacologic care or usual care, as applicable. Refer to Section 6.6.
- Clinic visits, as per specific study Appendix
- OHQ/OIQ
- COMPASS-31
- MAPS
- VOSS
- Active Stand Test
- 6-minute Walk Test
- DSQ-PEM
- PROMIS-29 + 2 Questionnaire
- PASC Symptom Questionnaire
- Safety assessment of AEs, including non-serious AEs, SAEs and ESIs
- · Height and weight, as applicable for dosing
- Vital signs monitoring
- Adherence to study intervention(s)
 - Care-coordinators will reach out weekly to the participants randomized to coordinated non-pharmacologic care
 - Pill count and weekly phone calls (for the first 30 days) for participants receiving oral study drugs to be taken at home

8.5 EARLY TERMINATION CLINIC VISIT

- Concomitant medication review
- OHQ/OIQ
- COMPASS-31
- MAPS
- Active Stand Test
- VOSS
- 6-minute Walk Test

- DSQ-PEM
- PROMIS-29 + 2 Questionnaire
- PASC Symptom Questionnaire
- Blood collection for safety parameters (complete blood count [CBC] with differential and complete metabolic panel [CMP]), biomarkers, and biospecimens for biorepository, including distribution of stool sample kit
- Skin biopsy, if applicable
- Autonomic function tests at specialized sites
- Safety assessment of AEs, including non-serious AEs, SAEs and ESIs
- Adherence to study interventions

8.6 END OF INTERVENTION CLINIC VISIT (DAY EOI + 1 WEEK)

The EOI Clinic Visit will occur after the final study intervention administration. The following will occur at the EOI Clinic Visit:

- Concomitant Medication review
- OHQ/OIQ
- COMPASS-31
- MAPS
- VOSS
- Active Stand Test
- 6-minute Walk Test
- DSQ-PEM
- PROMIS-29 + 2 Questionnaire
- PASC Symptom Questionnaire
- Blood collection for safety parameters (CBC with differential and CMP), biomarkers, and biospecimens for biorepository, including distribution of stool sample kit
- Autonomic function tests at specialized sites
- Skin biopsy, if applicable
- Safety assessment of AEs, including non-serious AEs, SAEs and ESIs
- Adherence to study interventions

8.7 FOLLOW-UP (EOI + 3 MONTHS)

Follow-up clinic visits will occur 3 months after EOI. The following assessments will occur as part of follow-up:

- OHQ/OIQ
- COMPASS-31
- MAPS
- VOSS
- Active Stand Test
- 6-Minute Walk Test
- DSQ-PEM
- PROMIS-29 + 2 Questionnaire
- PASC Symptom Questionnaire
- Blood Collection for biomarkers and biospecimens for biorepository
- Safety assessment of SAEs and ESIs

8.8 CLINICAL LABORATORY ASSESSMENTS

A CBC with differential and CMP are required as part of baseline study assessments. If a participant has these laboratory assessments available within 3 months of study enrollment, they do not need to be repeated as part of the study.

Blood will be collected from participants and sent to a central lab for biomarkers assessment (refer to Section 8.9.12)

8.9 STUDY ASSESSMENTS

8.9.1 ORTHOSTATIC HYPOTENSION QUESTIONNAIRE

The Orthostatic Hypotension Questionnaire (OHQ) is a PRO that measures orthostatic intolerance, which has been the primary presentation of patients with PASC-related autonomic dysfunction. This measure includes the 6-item symptom assessment (OHSA) and the 4-item Daily Activity Scale (OHDAS). Each item is scored from 0 (none/no interference) to 10 (worst possible/complete interference), describing the preceding week. Prior data in patients with PASC suggests that this measure well-discriminates those with symptoms compared to healthy controls. Measurements for the first 50 subjects will be used for test-retest reliability.

8.9.2 COMPASS-31

The COMPASS-31 is a PRO that measures autonomic symptoms across multiple domains commonly seen in patients with PASC. A COMPASS-31 score will be used to assess participant eligibility for specific study appendices. Refer to the Appendices for further details.

8.9.3 MALMO POTS SYMPTOM SCORE (MAPS)

The MAPS assesses symptom burden in POTS. It is a self-rating, 12-item score (0-10 per item, total range 0-120) based on patients' own perception of symptoms through visual analogue scale assessment.¹³

8.9.4 ACTIVE STAND TEST

The active stand test is a well-characterized performance measure of short-term neurologic and cardiovascular function that requires limited resources and aids in diagnosis of patient symptoms of orthostatic intolerance.

It is important to standardize and measure change from supine to standing and not from sitting with legs down to standing. Participants will remain supine for 10 minutes, and data will be acquired at 5 and 10 minutes. Standing test should be performed with HR and BP monitoring at 1, 3, 5 and 10 minutes. In

patients with orthostatic intolerance this test induces a BP drop within the first minute that correlates with HR and BP responses to tilt up, and can be used as a measure of orthostatic intolerance.¹⁴

8.9.5 VANDERBILT ORTHOSTATIC SYMPTOMS SCORE (VOSS)

The VOSS consists of 9 orthostatic symptoms rated on a scale of 0 (no symptom) to 10 (worst the participant has experienced) at the end of each HUT and after the active stand test. These symptoms are mental clouding, blurred vision, shortness of breath, rapid heartbeat, tremulousness, chest discomfort, headache, lightheadedness, and nausea.

8.9.6 6-MINUTE WALK TEST

Normal walking speed will be measured using a standard 6 minute walk, which is an established measure of overall health.¹⁵

8.9.7 MODIFIED DEPAUL SYMPTOM QUESTIONNAIRE-POST EXERTIONAL MALAISE

The Modified Depaul Symptom Questionnaire-Post Exertional Malaise (DSQ-PEM) PRO assessment. ¹⁶ Some patients have experienced debilitating PEM that may occur following exertion or on the day after exertion. To assess PEM, the Depaul Symptom Questionnaire PEM short form will be adapted. This scale assesses symptom frequency and severity over a 6-month look back period. However, for the purposes of this study, it will be modified to assess over a 7-day look back period at baseline, and at all other timepoints, the look back period will be since the last study visit.

It was previously validated in patients with myalgic encephalomyelitis/chronic fatigue syndrome. Frequency is rated on a 5-point Likert scale: 0 = none of the time, 1 = a little of the time, 2 = about half the time, 3 = most of the time, and 4 = all of the time. Severity is also rated on a 5-point Likert scale: 0 = symptom not present, 1 = mild, 2 = moderate, 3 = severe, 4 = very severe. Post-Exertional Malaise (PEM) will be deemed to be present if the participant reports having at least one moderate (rated severity ≥ 2) PEM symptom for a frequency rated ≥ 2 .

The DSQ-PEM questionnaire will be via phone or electronic survey within one day after strenuous inperson study visits. This evaluation will allow us to assess the effect of study procedures on exacerbation or development of PEM in a population who may be particularly vulnerable to this condition. For the purposes of assessing PEM related to the specific activity, it will be modified to assess PEM related specifically to the activity that occurred the previous day.

8.9.8 PROMIS-29+2

Patient-Reported Outcomes Measurement Information System (PROMIS-29) global health scale: The PROMIS was developed out of the "Roadmap for Medical Research" created by the NIH in 2002 as valid, generalizable items to standardize clinical research across NIH-funded research dealing with PROs. Multiple PROMIS scales have been validated across many clinical populations. The PROMIS-29 consists of 29 items that assess general domains of health and functioning, including overall physical health,

mental health, social health, pain, fatigue, and overall perceived quality of life. PROMIS-29+2 is the PROMIS-29 Profile v2.1 and two items from Cognitive Function – Abilities v2.0. The PROMIS global health scales has been correlated against the EuroQol EQ-5D. Additionally, PROMIS scales have been used with PASC patients. 9

8.9.9 SKIN BIOPSY

This will be required for all participants enrolled in specific Appendices only. This test evaluates intraepidermal small fiber density and sudomotor fiber density in distal, intermediate, and proximal legs. If the baseline result shows evidence of small fiber neuropathy, the biopsy will be repeated at EOI or ET.

8.9.10 AUTONOMIC FUNCTION TESTS AT SPECIALIZED SITES

Additional autonomic function testing will be completed at specific sites that have the capability to perform these tests:

- Head-up Tilt (HUT) Test This test stimulates the autonomic nervous system with orthostatic stress by passive movement from a supine position to an upright tilt in a controlled laboratory setting. Tilt test provides both sympathetic and parasympathetic assessments. The variables measured will be continuous BP, HR, end tidal CO₂, and cerebral blood flow velocity (CBFv) using transcranial doppler monitoring (CBFv and end tidal CO₂ only at sites that have the appropriate device). The technique used for this assessment will be 10 minutes of baseline and 10 minutes 70 degree head-up tilt (HUT). The outcome will be all variables at baseline and at every 1 minute of the tilt (total 10 measurements during the tilt).
 - Transcranial doppler (TCD), if available The test measures the velocity of blood flow through the brain's blood vessels by measuring the echoes of ultrasound waves.
- Valsalva maneuver This test evaluates sympathetic adrenergic functions using BP responses and cardiovagal (parasympathetic) functions using the HR responses. Valsalva maneuver consists of forced expiration against the resistance with the expiratory pressure during strain at 40 mm Hg for 15 seconds. The measured variables will be continuous BP and HR.
- Deep breathing test The test evaluates changes in the instant HR that is provoked by "deep" breathing. The test is performed in supine position. The measured outcomes will be the ratio of the HR during inspiration and expiration as well as the calculated respiratory sinus arrhythmia (RSA). RSA amplitude is defined as the difference between the end of expiration and end of inspiration in HR. Typically, 6 respiratory cycles are identified and the respective RSA amplitudes are averaged.

The final outcome of these measures is the quantitative autonomic symptoms score (QASAT). The total and sub scores (cardiovagal, adrenergic, respiratory, cerebral blood flow, small fibers, and autonomic failure) will be obtained.

8.9.11 PASC SYMPTOM QUESTIONNAIRE

Participants will be asked to complete a questionnaire that asks about the presence of PASC symptoms. This questionnaire includes additional PASC symptoms that are not directly related to autonomic dysfunction.

8.9.12 BIOMARKERS

Blood will be collected from participants and sent to a central lab for biomarkers assessment. The biomarkers assessed will include inflammatory cytokines, skin biopsies, and other viral biomarkers, immune biomarkers, Endothelial function biomarkers, and biomarkers for autonomic and cognitive function.

Refer to the MOP for further details regarding specimen collection, preparation, and shipment.

8.10 BIOREPOSITORY FOR FUTURE RESEARCH

The RECOVER Biorepository is designed to collect and store blood plasma, serum samples, and stool samples for future research related to the various studies of the RECOVER Program. Such research might include developing diagnostic and/or prognostic tests, improving our understanding of the underlying pathophysiology of PASC, and developing new therapeutic targets. Samples from biorepositories have proven to be enormously important in the last 20 years, as information on the components of blood has expanded rapidly. Important insights have been gained from biorepository samples from clinical trials and the stored samples from the RECOVER Program will prove equally productive and important. This biorepository will be conducted under the coordination of the DCRI which serves as the CT - DCC for all RECOVER clinical trials.

The existence of the biorepository will provide the opportunity to devise new hypotheses, since blood collection techniques are standardized across all approved protocols thereby allowing cross-protocol sample comparisons if scientifically justified.

Within this framework, the design of the biorepository is to collect additional blood and prepare EDTA plasma and serum, to be stored in the biorepository for future, as yet unspecified, analyses and studies. A serologic test of COVID-19 infection will be done as part of the biorepository for all participants. Approximately eighty (80) mL of blood will be collected to prepare the aliquots for storage at -80°C. Specifically, two 1.0 ml aliquots of EDTA plasma and two 1.0 ml aliquots of serum will be prepared. These samples will be stored at the Biorepository in a lab for up to 7 years.

SAFETY ASSESSMENTS AND REPORTING

9.1 ADVERSE EVENTS AND SERIOUS ADVERSE EVENTS

9.1.1 DEFINITION OF ADVERSE EVENTS (AE) AND SERIOUS ADVERSE EVENTS (SAE)

An AE is any untoward medical occurrence in humans, whether or not considered drug-related, which occurs during the conduct of a clinical trial. An AE can therefore be any change in clinical status, routine labs, x-rays, physical examinations, etc., that is considered clinically significant by the study investigator.

An SAE or serious suspected adverse reaction (SAR) or serious adverse reaction, as determined by the investigator or the sponsor, is an AE that results in any of the following serious outcomes:

- Death
- Life-threatening AE ("life-threatening" means that the study participant was, in the opinion of the investigator or sponsor, at immediate risk of death from the reaction as it occurred and required immediate intervention)
- Persistent or significant incapacity or substantial disruption of the ability to conduct normal life functions
- Inpatient hospitalization or prolongation of existing hospitalization
- Congenital abnormality or birth defect
- Important medical event that may not result in one of the above outcomes, but may jeopardize the health of the study participant or require medical or surgical intervention to prevent one of the outcomes listed in the above definition of serious event

Hospitalization for elective treatment of a pre-existing condition that did not worsen from baseline does not meet the definition of an SAE.

An unanticipated problem involving risk to non-participants, including an environmental exposure and exposure to a breast-feeding infant, will be reported to the sponsor and the site IRB and central IRB, as appropriate. An event occurring in a non-participant will not be entered into the electronic data capture (EDC) system. Refer to the MOP for details.

Medication errors with the study drug(s) resulting in an SAE are reportable. The medication error will be captured as a protocol deviation and the SAE captured on the SAE electronic CRF (eCRF).

9.1.2 COLLECTION PERIOD FOR AE AND SAE INFORMATION

Safety event collection will occur at the pre-specified study visits but all participants will be instructed to self-report concerns by calling the site.

Serious adverse events (SAEs) will be collected from the first invasive study procedure (blood collection or nasal swab) through the end of the Follow-up period (EOI = 3 months). Serious adverse events (SAEs) or ESIs (per Appendix) will be extracted by site personnel from the participant's medical record if the participant seeks medical care or if hospitalization occurs, each of which notifies the site to conduct follow-up.

Medical occurrences that begin before the first invasive study procedure (blood collection or nasal swab), but after obtaining informed consent, will not be considered an AE. The medical occurrence or condition will be captured on the Medical History eCRF.

Each study drug appendix will identify whether if all non-serious AEs will be collected to the study database from Start of study drug administration through the end of study drug dosing. Any non-serious AEs that result in study drug discontinuation will be identified as the reason for study drug discontinuation in the study database and reported as an AE. Any non-serious AEs that are ESIs will be collected from the start of study drug administration through the end of follow-up period.

9.1.2.1 SEVERITY OF EVENT

For reportable events, the following guidelines will be used to describe severity grades, described in the Common Terminology Criteria for Adverse Events (CTCAE) v5.0, published November 27, 2017.

Grade refers to the severity of the AE. The CTCAE displays Grades 1 through 5 with unique clinical descriptions of severity for each AE based on this general guideline:

- Grade 1 Mild; asymptomatic or mild symptoms; clinical or diagnostic observations only; intervention not indicated.
- Grade 2 Moderate; minimal, local or noninvasive intervention indicated; limiting ageappropriate instrumental activities of daily living (ADL)*.
- Grade 3 Severe or medically significant but not immediately life-threatening; hospitalization or prolongation of hospitalization indicated; disabling; limiting self-care activities of daily living
- Grade 4 Life-threatening consequences; urgent intervention indicated.
- Grade 5 Death related to AE.

9.1.2.2 RELATIONSHIP TO STUDY INTERVENTION

All reportable events must have their relationship to the study intervention assessed by the by site investigator based on temporal relationship and his or her clinical judgment. The degree of certainty about causality will be graded using the categories below. In a clinical trial, the study product must always be suspect.

- Related The AE is known to occur with the study intervention, there is a reasonable possibility
 that the study intervention caused the AE, or there is a temporal relationship between the study
 intervention and event. Reasonable possibility means that there is evidence to suggest a causal
 relationship between the study intervention and the AE
- Not Related There is not a reasonable possibility that the administration of the study intervention caused the event, there is no temporal relationship between the study intervention and event onset, or an alternate etiology has been established

9.1.3 TIME PERIOD AND FREQUENCY FOR EVENT ASSESSMENT AND FOLLOW-UP

The occurrence of an AE, ESI, or SAE may come to the attention of study personnel during study visits and interviews of a study participant presenting for medical care, or upon review by a study monitor.

All reportable events will be captured on the appropriate CRF. Information to be collected includes event description, date/time of onset, clinician's assessment of severity, relationship to study product (assessed only by those with the training and authority to make a diagnosis), action taken with study drug (e.g. study drug discontinuation), and date/time of resolution/stabilization of the event. All events occurring within the pre-specified reportable time windows must be documented appropriately regardless of relationship.

Any medical condition that is present at the time the participant is screened will be considered as baseline and not reported as an AE. However, if the study participant's condition deteriorates at any time during the study, it will be recorded as an AE.

Changes in the severity of an AE will be documented to allow an assessment of the duration of the event at each level of severity to be performed. Adverse events (AEs) characterized as intermittent require documentation of onset and duration of each episode.

The investigator will follow all SAEs until resolution, stabilization, or the event is otherwise explained. The DCRI Safety Surveillance Team will follow all SAEs until resolution, stabilization, or until otherwise explained.

9.1.4 REPORTING AND MONITORING OF SAES

All of the study drugs used in this platform protocol will be under an IND and subject to IND regulations in 21 CFR 312 since their investigational use for treatment of PASC is not an approved indication. The IND sponsor or DSMB will review aggregate safety data. The IND sponsor will be responsible for determining if the safety reporting criteria are met per 21 CFR 312.32(c)(1)(i)(C) and 21 CFR 312.32(c)(1)(iv) and will notify the CT-DCC to prepare an aggregate report for submission to the US FDA. An aggregate safety report will be submitted to FDA as soon as possible, but in no case later than 15 calendar days after the IND sponsor determination. If the IND sponsor determines that an unexpected fatal or life-threatening SAR occurs markedly more frequently in a study drug arm than in the control arm, an aggregate safety report will be submitted to the FDA as soon as possible, but in no case later than 7 calendar days after the IND sponsor determination. Information on individual SAEs will be available upon request from the FDA following the submission of any aggregate reports.

Individual SAEs must be entered into the data system within 24 hours of site awareness. The DCRI Safety Surveillance team will notify pharmaceutical partners of SAEs, if applicable within 1 to 2 business days of their receipt that occur involving the specific appendix of the supplied study drug/control, as required. Serious Adverse Events that are related and confirmed unlisted by the DCRI Safety Medical Monitor and IND sponsor will be reported to the FDA as suspected unexpected serious adverse reaction (SUSARs); as 7-day reports for unexpected fatal or life-threatening adverse reactions and 15-day reports for serious

and unexpected adverse reactions. The SUSARs will be shared with the pharmaceutical partner of the supplied study drug according to the same timelines. If the IND sponsor, DSMB, or FDA note a clinically important increase in the rate of a SUSAR, the IND sponsor or designee will notify investigators no later than 15 calendar days after determining that the information qualifies for reporting. The investigators will notify their local IRB according to local guidelines if applicable. The CT-DCC will notify the central IRB. On a monthly basis, the DSMB chair will receive a report of the listing and overall pattern of SAEs, including hospitalizations. Refer to the Safety Management Plan for further details regarding specific reporting timelines. In addition, see other sections (7.4 and 13.10) for guidelines for pausing or stopping drug in individual patients and the guidelines for pausing the trial while the DSMB reviews the safety data to make their recommendation.

Investigators are not obligated to actively seek information on AEs or SAEs after the participant has concluded study participation. However, if the investigator learns of any SAE, including a death, at any time after a participant has completed the study (but before the study itself has ended), and they consider the event to be reasonably related to the study intervention, the investigator must promptly report the SAE to the sponsor via EDC entry.

9.1.5 EVENTS OF SPECIAL INTEREST

Each study drug may have a unique list of ESIs. Refer to the relevant Appendix.

Events of special interest (ESIs) meeting serious criteria will be reported to the DSMB chair within 1-2 business days, and all events (non-serious AEs/SAEs) will be reported to the DSMB chair via a monthly listing.

9.1.6 REPORTING OF PREGNANCY

Pregnancies that occur following the first study drug administration while on-study will be collected. If the study drug appendix excludes pregnancy, the participant will be advised to discontinue study drug(s). Pregnancies will be followed until pregnancy outcome unless the participant refuses to provide consent. Pregnant participants will be asked to complete a pregnancy-specific consent form if the final outcome of the pregnancy is not reached while the participant is on-study. Any pregnancy-associated ESI or SAE should be reported if information can be collected and entered into the CRF. The DCRI Safety Surveillance team will notify pharmaceutical partners if applicable of a pregnancy within 1-2 business day of receipt that occur involving the specific appendix of the supplied study drug/control, as required.

10 STATISTICAL ANALYSES

All statistical analyses will be performed using SAS (SAS Institute, Inc. Cary, NC, USA) version 9.4 or higher. Baseline demographic and clinical data will be summarized by intervention arm. Descriptive statistics will include mean, standard deviation, median, 25th and 75th percentiles for continuous variables and frequency and percentage for categorical variables. Statistical comparisons will be

performed using two-sided tests at 0.05 significance level. Additional details regarding statistical analyses will be provided in the SAP that is available prior first patient enrollment and that will be finalized prior to the database lock.

10.1 STATISTICAL HYPOTHESES

Primary Endpoint:

1. Improvement in OHQ/OIQ composite score at the EOI compared to baseline in intervention(s) vs. control.

Secondary Endpoints:

- 1. Improvement in symptom-specific outcomes (COMPASS-31 and MAPS) at the EOI compared to baseline in intervention(s) vs. control
- 2. Improvement in performance-based and vital sign outcome measures (Active Stand Test, BP, HR, 6-minute Walk Test) at the EOI compared to baseline in intervention(s) vs. control
- 3. Improvement in quality of life (PROMIS-29+2 summary score) at the EOI compared to baseline in intervention(s) vs. control
- 4. Improvement in specific tracked accelerometry measurements (step count, HR) via wearable device at the EOI compared to baseline in intervention(s) vs. control
- 5. Study drug(s) are safe in the PASC population

10.2 SAMPLE SIZE DETERMINATION

This study uses an adaptive platform trial design that will allow study drugs to be added or dropped from consideration based on accruing evidence. Initial sample size estimates are based on 2 trials (Appendix A and Appendix B), each with a 2 x 2 factorial design, for each study drug in combination with non-pharmacologic intervention in a 2 x 2 factorial design. With 4 different combinations in the trial, this results in 1:1:1:1 random allocation. If additional study drugs with the same route are added later in the study drug factor in any of the 2 x 2 factorial studies that can utilize a shared control, the design may be (m+1) x 2 factorial design with m active study drugs and the sample size will be adjusted accordingly.

The primary interest of studies in this platform protocol is to evaluate the effect of the study interventions vs control. Studies in this platform design will also explore the potential interaction of study drug and non-pharmacologic intervention. However, we do not expect to have interactions among effects of the study drug with effects of non-pharmacologic care. The primary comparisons are the main effects of the interventions, i.e., study drug vs. placebo and coordinated non-pharmacologic care vs. control (usual care) within each study appendix. Power calculation is provided for each study appendix based on sample size without adjusting for multiplicity because the study is a phase II where we try to identify potential efficacy. Furthermore, the main effect comparisons are considered in a factorial design where the degree of need for multiplicity adjustment is low due to the comparisons are nearly statistically independent due to orthogonal contrasts of the cells if the variances in the different cells are equal.²⁰

For the primary comparisons in each appendix, the power is calculated to detect a clinically significant difference of one unit (equivalent to effect size of 0.5 based on SD of 2) in changes of OHQ/OIQ composite score between any of the 2 groups, assuming a missing date rate with 0.05 type I error with a 2-sided test. Further details on sample size, power and missing data rate are found in the specific appendices.

Our justifications for the above assumptions are based on the literature. The clinically significant difference is chosen to be one unit in OHQ/OIQ composite score because the estimated minimally important difference (MID) ranges from 0.8 to 1.0 in a validation study of OHQ/OIQ.²¹ The assumption of standard deviation (SD) of 2 in change of OHQ/OIQ composite score is based on the observed SDs of 2.07 (active arm) and 1.69 (control arm) in change of OHQ/OIQ composite score within 7 days in a phase 3 randomized, placebo-controlled clinical trial on droxidopa for neurogenic orthostatic hypotension.²² The observed SD of OHQ/OIQ composite score in 48 POTS subjects is 1.68 and we would have SD of 2 or 2.1 in change of OHQ/OIQ composite score if the correlation between OHQ/OIQ composite score at baseline and the EOI is 0.3 or 0.2.

10.3 POPULATIONS FOR ANALYSES

Population for the primary efficacy analysis will be based on an intention-to-treat (ITT) population, consisting of all randomized participants who receive the study intervention or control. All randomized participants will be included and will be analyzed according to their randomly assigned treatment group. Population for safety analyses will be performed among participants in the ITT population analyzed according to their randomly assigned treatment groups.

10.4 STATISTICAL ANALYSES

This section describes analysis methods for the primary, secondary, safety and exploratory outcomes. Full details will be provided in the SAP. If additional study drugs are added later, the statistical analysis section will be revised appropriately.

10.4.1 ANALYSIS OF THE PRIMARY ENDPOINT

For primary comparison, the analysis population includes all randomized participants to study drug intervention or its control, and the primary endpoint is the change of OHQ/OIQ composite score from baseline to the EOI.

Same analysis approach will be used in all these primary comparisons. A linear regression approach will be used for the primary analyses where the change of OHQ/OIQ composite score is the dependent variable and group indicator (for coordinated care or for IVIG depending on the primary analysis), and baseline OHQ/OIQ composite score and other pre-specified baseline variables are covariates (independent variables). The null hypothesis of no group difference corresponds to the coefficient for the group indicator is zero. Rejection of this null hypothesis will provide evidence for presence of the intervention effect.

In addition to these primary analyses, we will also perform 2 exploratory analyses as a supplement. Specifically, we will include the testing of interaction between pharmacologic and non-pharmacologic treatments within each trial. Furthermore, we will assess the equality of non-pharmacologic treatment effect by testing the interaction of non-pharmacologic treatment and the trial indicator. Details will be provided in the SAP.

10.4.2 ANALYSIS OF THE SECONDARY ENDPOINT(S)

Symptom-specific outcomes (COMPASS-31 and MAPS). Each of the symptom-specific outcomes is a continuous measure. Changes from baseline to EOI for each outcome will be analyzed using similar methods as those outlined for the primary endpoint. Our pilot preliminary data showed that COMPASS-31 has mean (SD) of 33.41 (18.92) in subjects with long COVID-19 and POTS. A 30% reduction in COMPASS-31 is considered to be clinically significant. This implies that 10-unit difference is clinically significant if the mean COMPASS-31 is 34 at the baseline. The power for this secondary outcome is provided in the appendix.

Performance-based and vital sign outcome measures (Active Stand Test, BP, HR, 6-minute Walk Test). Each of the performance and vital sign -specific outcomes is a continuous measure. Changes from baseline to EOI for each outcome will be analyzed using similar methods as those outlined for the primary endpoint.

Quality of life (PROMIS-29+2). Each of the symptom-specific outcomes is a continuous measure. Changes from baseline to EOI for each outcome will be analyzed using similar methods as those outlined for the primary endpoint.

Specific tracked accelerometry measurements (step count, HR). Each of the symptom-specific outcomes is a continuous measure. Changes from baseline to EOI for each outcome will be analyzed using similar methods as those outlined for the primary endpoint.

Safety endpoint. Safety endpoints include the proportion of participants who experience individual SAEs and the proportion who experience any one or more SAEs. These will be analyzed in the safety population. Events of Special Interest (ESIs) will be summarized by study drug appendix and duration. Incidence of AEs/SAEs leading to discontinuation will also be summarized.

10.4.3 ANALYSIS OF THE EXPLORATORY ENDPOINT(S)

The quantitative autonomic symptoms score (QASAT) from the autonomic function tests, VOSS score after the active test or HUT, PASC symptom questionnaire scores; biomarkers (viral, autonomic function, immune, inflammatory and endothelial function biomarkers), and cognitive function (CCL11) are the exploratory endpoints. Each of the outcome is a continuous measure. Changes from baseline to EOI for each outcome will be analyzed using similar methods as those outlined for the primary endpoint.

10.4.4 MISSING DATA

The primary and secondary analyses will be performed without missing data imputation. Additional sensitivity and supplementary analyses with adjustments for missing data will be described in the SAP.

10.4.5 PLANNED INTERIM ANALYSES

Interim examination of clinical endpoints and key safety events will be performed at regular intervals during the course of the trial. An independent, NIH-appointed, DSMB will monitor participant safety and review performance of the trial. The primary objective of these interim analyses will be to ensure the safety of the participants enrolled in the trial and evaluate the accumulating endpoint data by treatment group in order to make interim assessments of futility. In addition, interim monitoring will involve a review of participant recruitment, compliance with the study protocol, status of data collection, and other factors which reflect the overall progress and integrity of the study.

This protocol does not have planned early stopping rules for efficacy. Because PASC presentations and outcomes are highly varied; an important study objective is to estimate the effect of treatment on a wide range of participant-relevant outcomes. If the study were to be stopped early with less than the full sample size, it would decrease precision and reduce the study's ability to characterize treatment risks and benefits based on important secondary effectiveness and safety outcomes. Stopping early would also limit the collection of data that are critical for planning future trials in similar patient populations.

10.4.6 PLANNED SUBGROUP ANALYSES

The following 2 subgroups are pre-specified with the same analysis approaches as those for **the primary and secondary outcomes.**

Confirmed antigenemia/immune biomarker subgroup is a subset of the ITT group and consists of participants who have detectable antigen or immune biomarkers. This subgroup is of particular interest because the efficacy of antiviral/immune-targeting agents may be greater in participants with documented evidence of viral persistence/immune response. Theoretically, if the hypothesized benefit of study drug(s) is concentrated in participants with viral/immune response persistence, then the ITT treatment effect may be attenuated by inclusion of participants with uncertain viral/immune response persistence status.

No confirmed antigenemia/immune biomarker population subgroup is a subset of the ITT population and consists of participants who tested negative for antigenemia/immune biomarkers Analysis of this subgroup is of particular relevance because viral persistence testing is not widely available in practice. If efficacy is demonstrated for the overall study cohort, it may not be automatically assumed that the benefit extends equally to the subset of participants without confirmation of antigenemia/immune biomarkers.

10.4.7 PLANNED TEST-RETEST RELIABILITY ANALYSES

Subjects will be asked to fill out the OHQ/OIQ questionnaire twice, once at screening and once at baseline. The data from the first 50 subjects will be used for the test-retest reliability analyses. Details will be provided in the SAP.

10.4.8 PLANNED SAMPLE SIZE AND POWER RE-ESTIMATION

In order to assist the DSMB in monitoring potential risk of increased drop out and subsequent decision, we plan to carry out blinded sample size and power re-estimation when approximately half of the targeted number of subjects have enrolled in each trial. The observed overall dropout rate and the observed overall standard deviation in the change of OHQ will be used for these blinded calculations. No efficacy analysis will be performed.

11 SUPPORTING DOCUMENTATION AND OPERATIONAL CONSIDERATIONS

11.1 REGULATORY, ETHICAL, AND STUDY OVERSIGHT CONSIDERATIONS

11.1.1 INFORMED CONSENT PROCESS

11.1.1.1 INSTITUTIONAL REVIEW BOARD (IRB)

The protocol, informed consent form(s) [ICF(s)], recruitment materials, and all participant materials will be submitted to the Institutional Review Board(s) [IRB(s)] of record for review and approval. This approval must be obtained before any participant is enrolled. Any amendment to the protocol will require review and approval by the IRB(s) before being implemented in the study. All changes to the consent form will also be IRB-approved and a determination will be made regarding whether previously consented participants need to be re-consented.

11.1.1.2 CONSENT PROCEDURES AND DOCUMENTATION

All consenting will occur either via an electronic consent process or a paper process. Consent forms describing in detail the study drug/control, study procedures, and risks will be given to the participant and documentation of informed consent is required prior to starting study procedures. Informed consent is a process that is initiated prior to the individual's agreement to participate in the study and continues throughout the individual's study participation. A description of risks and possible benefits of participation will be provided to the participants. Consent forms will be IRB-approved and the participant will be asked to read and review the document. The participant will be provided contact information in the event they have questions about study participation. This will allow them to communicate with the investigators (or their delegate), for further explanation of the research study and to answer any questions that may arise, as necessary. Participants will have the opportunity to carefully review the consent form and ask questions prior to signing.

The participants should have the opportunity to discuss the study and think about it prior to agreeing to participate. The participant will sign the informed consent document prior to any procedures being done specifically for the study. The participants may withdraw consent at any time throughout the course of the study. A copy of the informed consent document will be provided to the participants for their records. The rights and welfare of the participants will be protected by emphasizing to them that the quality of their medical care will not be adversely affected if they decline to participate in this study.

The study team will distinguish between the desire to discontinue study drug and the desire to withdraw consent for study follow-up.

11.1.2 CONFIDENTIALITY AND PRIVACY

Participant confidentiality is strictly held in trust by the participating investigators, their staff, and the sponsor(s) and their agents. This confidentiality is extended to cover testing of biological samples in addition to the clinical and private information relating to participants. Therefore, the study protocol, documentation, data, and all other information generated will be held in strict confidence. No information concerning the study or the data will be released to any unauthorized third party without prior written approval of the sponsor. The study participant's contact information will be securely stored in the clinical study database.

Study participant research data, which is for purposes of statistical analysis and scientific reporting, will be transmitted to and stored at the CT-DCC. The study data entry and study management systems used by clinical sites and by research staff will be secured and password protected. At the end of the study, all study-related data storage systems will be archived according to local processes.

11.1.3 KEY ROLES AND STUDY GOVERNANCE

The RECOVER program is overseen by the RECOVER Executive Committee. The RECOVER program also includes a Clinical Trial Steering Committee, which is a multi-stakeholder committee that oversees the study and includes patients, the CT-DCC, the NIH, the FDA, and academic and subject matter experts.

The CT-DCC is overseen by a Principal Investigator. The CT-DCC is responsible for study coordination, site management, communication, financial administration, treatment allocations, receipt and processing of data, quality control programs, and statistical analysis and reporting.

The DSMB will oversee the safety and welfare of trial participants as well as provide recommendations for continuation, discontinuation, or revision of the trial.

11.1.4 DATA AND SAFETY MONITORING BOARD

Safety oversight will be under the direction of the RECOVER DSMB composed of individuals with the appropriate expertise. Members of the DSMB should be independent from the study conduct and free of conflict of interest, or measures should be in place to minimize perceived conflict of interest. The DSMB will meet at least semi-annually to approve protocols, assess safety and efficacy data, and at

appropriate intervals to meet requirements for the Interim Analyses on each arm of the study. The DMSB will operate under the rules of an approved charter that will be written and reviewed at the organizational meeting of the DSMB. At this time, each data element that the DSMB needs to assess will be clearly defined. The DSMB will provide its input to the NIH.

11.1.5 CLINICAL MONITORING

This study will employ a centralized risk-based approach to monitoring with routine and periodic review of site-submitted data to review the informed consent process, select eligibility criteria, medical history, identify and follow-up on missing data, inconsistent data, data outliers, etc. and ensure completion of administrative and regulatory processes. The study team will facilitate regular communication through training sessions, teleconferences, videoconferencing, email, etc. Using quality-by-design principles, steps will be taken at the study design stage to foresee and limit significant problems that might occur during the study conduct. Follow-up from the sites is expected to keep participants engaged.

11.1.6 QUALITY ASSURANCE AND QUALITY CONTROL

The study team will work in tandem to ensure that the data collected in this study are as complete and correct as possible. A four-step, multi-functional approach to quality control will be implemented:

- Training: Prior to the start of enrollment, the clinician investigators and key study personnel at
 each site will be trained with the clinical protocol and data collection procedures, including how
 to use the EDC system. Follow-up training and training for new study personnel or new versions
 of the protocol will be conducted as needed.
- Monitoring: The RECOVER CT-DCC will ensure that data collection is handled properly, will
 provide in-service training, and will address questions from site investigators and coordinators.
 Electronic review of data quality and completeness will occur on a regular and ongoing basis.
 Any issues will be addressed.
- Managing data: After the data have been transferred for statistical summarization, data description, and data analysis, further crosschecking of the data will be performed with discrepant observations being flagged and appropriately resolved through a data query system.
- Reviewing data: Data regarding events of interest will be reviewed to ensure appropriate documents are collected for DSMB review. The CT-DCC will monitor study data and contact site study teams when events comprising the primary endpoint are not complete.

11.1.7 DATA HANDLING AND RECORD KEEPING

11.1.7.1 DATA COLLECTION AND MANAGEMENT RESPONSIBILITIES

Streamlining research activities and conducting the trial in a pragmatic manner will increase the ability to complete the trial in the face of strained clinical and research resources. Data may be collected by

electronic methods, supplemented by telephone or videophone follow-up, and from the electronic health record.

Data will be collected directly from participants using REDCap through text messaging or email with a survey link, or phone call as back up. The process for using text messaging and email is Health Insurance Portability and Accountability Act (HIPAA) compliant.

Site personnel or participants will enter study data into a secure online database. Data will be maintained in a secure online database until the time of study publication. At the time of publication, the CT-DCC will generate a de-identified version of the database for archiving (see Section 11.1.9). All source documents at the sites should be completed in a neat, legible manner to ensure accurate interpretation of data.

11.1.7.2 STUDY RECORDS RETENTION

Study documents should be retained for a minimum of 6 years after the study has ended. However, if required by local regulations or the US FDA, these documents should be retained for a longer period. No records will be destroyed without the written consent of the sponsor. It is the responsibility of the sponsor to inform the investigator when these documents no longer need to be retained.

11.1.8 PROTOCOL DEVIATIONS

A protocol deviation is defined as non-compliance with the clinical study protocol or GCP requirements. The non-compliance may be on the part of the participant, site investigator, or the site staff.

A major protocol deviation is a significant divergence from the protocol that may have significant effect on the participant's safety, rights, or welfare and/or on the integrity of the study data. Major protocol deviations must be sent to the study IRB and local IRB per their guidelines, recorded in source documents, and reported to the coordinating center. All protocol deviations will be documented. For this study, any missed or delayed survey completion will not be considered a major protocol deviation, unless it is a study procedure that is required for the primary endpoint.

11.1.9 PUBLICATION AND DATA SHARING POLICY

This study will comply with the NIH Public Access Policy, which ensures that the public has access to the published results of NIH-funded research. It requires scientists to submit final peer-reviewed journal manuscripts that arise from NIH funds to the digital archive PubMed Central upon acceptance for publication.

11.1.10 CONFLICT OF INTEREST POLICY

The independence of this study from any actual or perceived influence, such as by the pharmaceutical industry, is critical. Therefore, any actual conflict of interest of persons who have a role in the design, conduct, analysis, publication, or any aspect of this trial will be disclosed and managed. Furthermore, persons who have a perceived conflict of interest will be required to have such conflicts managed in a

way that is appropriate to their participation in the design and conduct of this trial. The study leadership in conjunction with the NIH has established policies and procedures for all study group members to disclose all conflicts of interest and will establish a mechanism for the management of all reported dualities of interest.

11.2 ABBREVIATIONS

AE	Adverse Event	
BP	Blood Pressure	
bpm	Beats per minute	
CFR	Code of Federal Regulations	
CONSORT	Consolidated Standards of Reporting Trials	
COVID-19	Coronavirus Disease of 2019	
CRF	Case Report Form	
CT-DCC	Clinical Trial – Data Coordinating Center	
DCRI	Duke Clinical Research Institute	
DSMB	Data Safety Monitoring Board	
EOI	End of Intervention	
ESI	Events of Special Interest	
FDA	Food and Drug Administration	
GCP	Good Clinical Practice	
HIPAA	Health Insurance Portability and Accountability Act	
HR	Heart Rate	
ICF	Informed Consent Form	
ICH	International Conference on Harmonization	
ICMJE	International Committee of Medical Journal Editors	
ICU	Intensive Care Unit	
IND	Investigational New Drug Application	
IRB	Institutional Review Board	
MAPS	Malmo POTS Symptom Score	
mITT	Modified Intention-To-Treat	
MOP	Manual of Procedures	
NCT	National Clinical Trial	
NIH	National Institutes of Health	
OHQ	Orthostatic Hypotension Questionnaire	
OIQ	Orthostatic Intolerance Questionnaire	
PASC	Post-acute Sequelae of SARS-CoV-2 Infection	
PHI	Personal Health Information	
PRO	Patient-Reported Outcome	
QASAT	Quantitative Autonomic Symptoms Score	
RECOVER	Researching COVID to Enhance Recovery	
SAE	Serious Adverse Event	
SAP	Statistical Analysis Plan	
SAR	Suspected Adverse Reaction	

SARS-CoV-2	Severe Acute Respiratory Syndrome Coronavirus 2		
SD	Standard Deviation		
SUSAR	Suspected Unexpected Serious Adverse Reaction		
ULN	Upper Limit of Normal		
US	United States		
VOSS	Vanderbilt Orthostatic Symptoms Score		

11.3 PROTOCOL AMENDMENT HISTORY

Version	Date	Description of Change
1.0	22SEPT2023	None, original protocol
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		IVIG. If the baseline skin biopsy shows evidence of small fiber
		neuropathy, the biopsy will be repeated at EOI or ET
		 The description for DSQ-PEM was updated throughout the protocol to remove the phrase "remote follow-up"

 The description of QHQ/OIQ was updated to reflect the appropriate look back periods: 7 day look back period at baseline and at all other timepoints, the look back period will be since last visit

- Section 9.1.5 was updated with the following statement: Events of special interest (ESIs) meeting serious criteria will be reported to the DSMB chair within 1-2 business days and all events (non-serious AEs/SAEs) will be reported to the DSMB chair via a monthly listing.
- Following statement was added to Section 8.2 to describe the Screening and Baseline visit: It is required to have completed the Screening prior to the Baseline/Randomization visit.
 Screening cannot be on the same day as the Baseline/Randomization visit.
- Updated description of active stand test: Participants will remain supine for 10 minutes, and data will be acquired at 5 and 10 minutes. Standing test should be performed with HR and BP monitoring at 1, 3, 5 and 10 minutes.
- Updated QASAT score description for autonomic function testing.
- Added clarity to the section on VOSS to indicate that it will be done after at the end of each HUT and after the active stand test.
- Added indication for blood collection to the End of
 Intervention study visit in the Schedule of Procedures tables in
 Appendix A and B, along with the following footnote: Blood
 may be collected at any time as needed based on PI discretion
 and adverse events.
- Modified the wording of the dosing information for IVIG in Appendix A as follows: The dosing frequency of 0.5 g/kg once a week is strongly suggested; however, the frequency can be modified based on individual site preference
- Updated description of placebo capsules in Appendix B for consistency with manufacturing details
- Added additional exclusion criteria to Appendix B based on FDA recommendation and product label
- Removed adherence to wearable as a study procedure throughout the protocol.

3.0	10JAN2023	 The coordinated non-pharmacologic care patient log was changed from a daily log to a weekly log; similarly, the daily monitoring of BP was changed to weekly for those receiving coordinated non-pharmacologic care Clarified throughout the protocol that the DSQ-PEM will be done twice at baseline: once on the day of the visit with a look back period of "past 7 days" and once the day after the visit with a look back period of "since last study visit". Removed the following statement from Section 8.9.7 as it is not relevant: "At least one moderate symptom (severity ≥ 2, frequency ≥ 2) on the DSQ-PEM will make a participant
		 included in this cluster and will trigger other baseline assessments listed below." Reworded the statement in Section 8.9.10 to indicate that autonomic function tests will be completed at specific site (not select sites): "Additional autonomic function testing will be completed at specific sites that have the capability to perform these tests."
		 Updated Section 14.7 to present the information on Ivabradine 2.5 mg since the 2.5 mg strength (from Europe) will be the investigational product that is used in this study. The 2.5 mg tablet will be encapsulated to form a capsule (tablet + filler). Stool collection was inadvertently missing from the End of Intervention visit in the Appendix B Schedule of Procedures. It has now been added.
		 Added text to Section 8.4 Study Intervention Visit and footnotes to the appendix specific Schedule of Procedures to clarify procedures that will be completed to determine dosing. Added specification regarding medical history: At every clinic visit after screening visit, changes in medical history including SARS-CoV-2 infection since last visit will be collected. Added clarification that the CBC labs will be "with differential". Added concomitant medications review to each study visit and the early termination visit.

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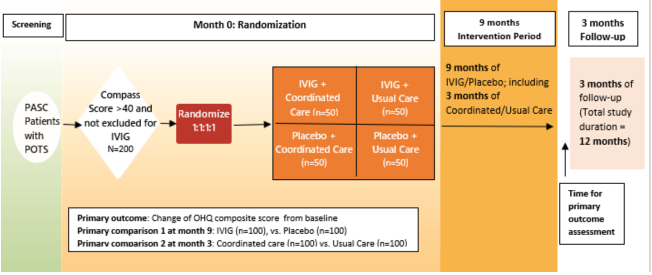
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13 APPENDIX A (IVIG)

13.1 STUDY DESIGN

Figure 2. Study Schema for Appendix A (IVIG)



*In this factorial design, participants will be randomized 1:1:1:1 to the combination of study drug /placebo and coordinated/usual care.

As depicted in **Figure 2**, approximately 200 participants will be enrolled in Appendix A (IVIG), based on eligibility criteria.

In the IVIG appendix, randomization will be implemented with 1:1:1:1 allocation among the combination of IVIG/placebo and coordinated/ usual non-pharmacologic care. All participants will receive IVIG or placebo for 9 months (36 weeks) with a follow-up period for an additional 3 months (total study duration for 12 months).

All participants will receive coordinated non-pharmacologic care or usual non-pharmacologic care (control) for a duration of 3 months. This will be concurrent with scheduled IVIG (or placebo) administration. coordinated non-pharmacologic care involves volume expansion through high salt diet (6-10 grams of salt), water intake (at least 2 liters per day), abdominal binder, exercise/rehabilitation, feedback, weekly monitoring of BP, motivation, education, and assisted care through a care coordinator.

The care coordinator will call each patient weekly during the 3 month non-pharmacologic intervention period to provide education and support on lifestyle modifications. Patients will be asked to maintain a weekly log of non-pharmacological care (average dietary intake of salt and water, physical rehabilitation) and provide a summary to the care-coordinators during the weekly calls. Further details of the non-pharmacologic intervention are provided in Section 6.6 and contained in the MOP.

Skin biopsies will be completed for all participants. In addition, participants enrolled in sites that are selected for autonomic function tests will be required to complete the other autonomic function tests as per the platform protocol schedule of procedures.

13.2 INTERVENTION RATIONALE

IVIG is approved by the FDA for the treatment of various immune deficiency syndrome and autoimmune diseases. IVIG at high dose is thought to neutralize pathogenic autoantibodies and suppresses autoimmune response, and the use of high titers of anti-SARS-CoV-2 antibodies for the treatment of COVID-19 is authorized in the US. From the past experience of using IVIG on severe cases of POTS, IVIG can be associated with improved autonomic symptoms, as well as with overall symptom burden. ²³⁻²⁷ IVIG may also improve other, non-cardiovascular dysregulation and dysautonomia and a potential to provide disease-modifying effects. Therefore, IVIG is a promising option for patients with severe PASC presenting as post-COVID-19-POTS.

13.3 RISK ASSESSMENT

Serious and unexpected AEs may occur that have not been previously reported with IVIG use. The most common adverse reactions of IVIG reported in greater than 5% of subjects during previous clinical trials are headache, cough, injection site reaction, nausea, pharyngitis, urticaria, vomiting, fever, back pain, rash, chills, hypertension, and asthenia. Previously reported AEs were identified in the OCTA-088 study, which was a multi-dose, open-label, multi-center study in patients with primary immunodeficiency. Common AEs included abnormal hematologic and clinical chemistry findings. Following approval of IVIG, other AEs have been reported including thromboembolic events, such as myocardial infarction (MI), stroke, pulmonary embolism (PE), and deep vein thromboses (DVT). These may be serious depending on the site and type of thrombosis. Cases of aseptic meningitis have been reported; however, no fatal cases have been observed. Acute renal failure has been observed, but in most cases, it was mild, but may be serious in elderly patients, patients with diabetes, and patients with pre-existing renal disease. Hemolytic anemia /hemolysis has also been observed, but in most cases, it was mild and self-limited.²⁸

The safety of IVIG for use in pregnancy and during lactation has not been established in controlled clinical trials. Clinical experience with immunoglobulins suggests that no harmful effects on the course of pregnancy, or on the fetus and the neonate are to be expected. Immunoglobulins are excreted into the milk and may contribute to the transfer of protective antibodies to the neonate.

Though side effects related to the pre- and post-medications (loratadine and acetaminophen) are rare, they can occur. Common adverse reactions of loratadine are headache, somnolence, fatigue, and dry mouth.²⁹ Common adverse reactions of acetaminophen are nausea, stomach pain, headache, hoarseness, loss of appetite, itching, rash, dark urine, clay-colored stools and swelling of the face, throat, tongue or limbs.

In a few cases, participants in this study may require steroids as pre-medication. General risks associated with steroids are increased appetite, acne, rapid mood swings, muscle weakness, and delayed wound healing.

13.4 ADDITIONAL APPENDIX-LEVEL INCLUSION CRITERIA

Abnormal active standing test defined as presence of orthostatic tachycardia (an increase of 30 beats per minute (bpm) or more in HR within 10 minutes upon standing without orthostatic hypotension) and experiencing orthostatic symptoms

2. COMPASS-31 Score > 40

13.5 ADDITIONAL APPENDIX-LEVEL EXCLUSION CRITERIA

- 1. Current or previous IVIG treatment
- 2. Contraindication to intravenous immunoglobulin.
- 3. Known allergic reactions to blood products including IVIG and/or subcutaneous immunoglobulin (SCIG), such as history of clinically relevant hemolysis after IVIG infusion, aseptic meningitis, recurrent severe headache, hypersensitivity, severe generalized or severe local skin reactions
- 4. Selective IgA deficiency
- 5. Current and recent (within 5 half-lives) use of high-dose corticosteroids (for example for prior solid organ transplant), omalizumab, anti-TNF-alpha inhibitors
- 6. Use of immunosuppressants such as Plaquenil, or low-dose steroid (prednisone, no more than 10mg a day) will be excluded unless the participant is on stable (>4 weeks) dose
- 7. Significant thrombotic events after the acute phase of COVID-19 and/or within 6 months of enrollment
- 8. Veins that are not viable for infusions
- 9. Not willing to adhere to dosing schedule for IVIG infusions for 9 months

13.5.1 PRECAUTIONS

The healthcare provider should consult other appropriate resources such as the prescribing information for the interacting drug for comprehensive information on dosing or monitoring.

There is clinical evidence of an association between the administration of immunoglobulins and thromboembolic events such as MI, PE, and DVT. Therefore, caution should be exercised when administering IVIG.

Most IVIG adverse effects are mild and are related to the rate of infusion. Therefore, IVIG should be started at a slow rate of infusion and gradually increased according to patient tolerance. Patients predisposed to renal failure include patients with any degree of pre-existing renal insufficiency, diabetes mellitus, age greater than 65, volume depletion, sepsis, paraproteinemia, or patients receiving known nephrotoxic drugs. Especially in such patients, IVIG products should be administered at the minimum concentration available and the minimum rate of infusion practicable.

As with other IVIG formulations, this product is prepared from large pools of human plasma, which may contain the causative agents of hepatitis and other viral diseases. The risk that such products will transmit an infectious agent has been reduced by screening plasma donors for prior exposure to certain viruses, by testing for the presence of certain current virus infections, and by inactivating and/or removing certain viruses.

IgA deficient patients with antibodies against IgA are at greater risk of developing severe hypersensitivity and anaphylactic reactions.

As recommended in good clinical practice, epinephrine should be administered for management of anaphylaxis. Instructions will be given to participants to understand signs of allergic reactions and when to seek medical attention for possible allergic reactions.

Hyperproteinemia, with resultant changes in serum viscosity and electrolyte imbalances may occur in patients receiving IVIG therapy.

Aseptic Meningitis Syndrome (AMS) has been reported with IVIG treatments, especially with high doses or rapid infusion.

Hemolysis, either intravascular or due to enhanced red blood cell sequestration, can develop subsequent to IVIG treatments. Risk factors include high doses and non-O blood group. Therefore, it is advised to monitor for hemolysis and hemolytic anemia.

It is advised to monitor for pulmonary adverse reactions (transfusion related acute lung injury) and volume overload.³⁰

The pre- and post-medications (loratadine and acetaminophen) should be used with caution in participants with decreased renal or liver function. These products are not to be used if the participant has had any allergic reactions to these ingredients in the past.²⁹ These medications will also be considered for participants who experience infusion related reactions.

13.6 CONCOMITANT MEDICATIONS

No medications or standard treatments for POTS should be withheld from study participants.

Participants who are already on stable (>4 weeks) doses of immunosuppressants as Plaquenil, methotrexate, and low-dose steroid (prednisone, no more than 10mg a day) will be allowed to continue the medication throughout the study.

During the study, initiation of steroids will not be allowed for more than 4 weeks, unless it is being used a pre-medication for IVIG treatment.

13.7 IVIG INFORMATION

Several formulations of IVIG have been approved by the FDA for use in various conditions including immune thrombocytopenic purpura, primary immunodeficiency, secondary immunodeficiency, Kawasaki disease, and prevention of graft versus host disease.

The FDA has not approved for the treatment of POTS or COVID-19. However, a number of case reports show improvement with immune modulating therapy, particularly IVIG, in patients with various post-COVID conditions

13.7.1 FORMULATION, APPEARANCE, PACKAGING, AND LABELING

IVIG is a sterile liquid preparation of highly purified immunoglobulin G (IgG) derived from large pools of human plasma. Each mL contains 100 mg of protein, of which ≥96% is IgG. a sterile liquid preparation of highly purified IgG derived from large pools of human plasma.

IVIG will be supplied as a sterile solution for injection supplied in 20 g (200 mL) single-use, tamper evident vials.

13.7.2 DRUG DISPENSING, STORAGE, AND STABILITY

IVIG may be stored for 36 months at 2-8°C (36-46°F) from the date of manufacture, and it may be stored at temperatures not to exceed 25°C (77°F) for up to 6 months anytime during the 36 month shelf life, after which the product must be immediately used or discarded.³⁰

13.7.3 DOSING AND ADMINISTRATION

IVIG (or placebo) will be administered intravenously by medical professional at the site. Double-blind will be maintained. The dose will be a **total of 2 g/kg every month**. The dosing frequency of 0.5 g/kg once a week is strongly suggested; however, the frequency can be modified based on individual site preference. There will be a dose cap at 80 kg (i.e. if weight > 80 kg, participant will be dosed at the 80 kg level).

The proposed dose (2 g/kg monthly) is evidence-based medicine. The dose is recommended for idiopathic thrombocytopenic purpura and chronic inflammatory demyelinating neuropathy.³¹ There is no plan to vary the IVIG dose since the proposed dose is evidence-based medicine, and there is no evidence that the higher or lower dose is superior over the proposed dose. The same dose was also used in anecdotal studies for long COVID patients with dysautonomia and small fiber neuropathy.^{32,33}

To reduce the occurrence of side effects and to enhance tolerability, fluids and pre- and post-meds are strongly encouraged. Therefore, 500 mL fluids (0.9% NaCl 500 mL IV), acetaminophen 650 mg, and loratadine 10 mg may be given as per PI discretion 30 minutes before IVIG (or placebo) infusion and after infusion. Some patients may require stronger pre- and post-medication (steroids, ketorolac) or additional fluids.

To ensure participant safety and monitor for adverse events, HR, BP, temperature, respiratory rate and oxygen saturation will be measured pre-, during, and post-infusion. Participants will be monitored for 30 minutes after receiving the infusion.

Decisions regarding dose interruptions or modifications will be made by the site investigator based on the clinical evaluation of the participant.

13.7.4 RATIONALE FOR SELECTION OF DOSE

Pre-clinical studies:

IVIG supplies a broad spectrum of opsonic and neutralizing IgG antibodies against bacterial, viral, parasitic, and mycoplasma agents, and their toxins. Immunoglobulins are fractionated blood products made from pooled human plasma. Immunoglobulins are endogenous proteins produced by B lymphocyte cells.

Pharmacokinetics

Following intravenous infusion, IVIG products show a biphasic decay curve. The initial (α) phase is characterized by an immediate post-infusion peak in serum IgG and is followed by rapid decay due to equilibration between the plasma and extravascular fluid compartments. The second (β) phase is characterized by a slower and constant rate of decay. The commonly cited "normal" half-life of 18 to 25 days is based on studies in which tiny quantities of radiolabeled IgG are injected into healthy individuals. When radiolabeled IgG was injected into patients with hypogammaglobulinemia or agammaglobulinemia, highly variable half-lives ranging from 12 to 40 days were observed. In other radiolabeled studies, high serum concentrations of IgG and hypermetabolism associated with fever and infection have been seen to coincide with a shortened IgG half-life.

In contrast, pharmacokinetic studies in immunodeficient patients are based on the decline of IgG concentrations following infusion of large quantities of immune globulin. In such studies, investigators have reported uniformly prolonged half-lives of 26 to 35 days.³⁴

Clinical studies:

Human IVIG preparations for intravenous or subcutaneous administration were approved for number of indications.³⁵ FDA approved IVIG for diseases such as primary humoral immunodeficiencies, prevention of infections in lymphocyte depleted populations (B-cell lymphoma, bone marrow transplants, pediatric HIV), demyelinating, and autoimmune diseases. The recommended dose of IVIG, 1 to 2 gm/kg given over one to 5 days, is administered when treating idiopathic thrombocytopenic purpura or other autoimmune related conditions. For chronic inflammatory demyelinating neuropathy (CIDP), the standard dose is 0.4 g/kg/day for 5 days (total dose 2 gram/kg), but in relapsing patients this dose may need to be repeated every 2-8 weeks to maintain improvement.

IVIG is a complex therapy and can lead to a number of adverse events. Most adverse reactions to IVIG are infusion rate related and mild. The most common are headaches (16%), fever (7%), hypertension (5%), chills (3%) and nausea (3%).³¹

The Immune Globulin Intravenous (Human), 10% Caprylate/Chromatography Purified CIDP Efficacy, also known as the ICE study, was a multi-center, randomized, double-blind, placebo-controlled trial conducted with Gammaked IVIG.³⁶ This study included 2 separately randomized periods to assess whether Gammaked IVIG was more effective than placebo for the treatment of CIDP (assessed in the Efficacy Period for up to 24 weeks) and whether long-term administration of Gammaked IVIG could maintain long-term benefit (assessed in the 24 week randomized withdrawal period). The Inflammatory Neuropathy Cause and Treatment (INCAT) was used to assess functional disability of both upper and lower extremities in demyelinating polyneuropathy. More subjects with CIDP responded to Gammaked: 28 of 59 subjects (47.5%) responded to Gammaked compared with 13 of 58 subjects (22.4%) administered placebo (25% difference; 95% CI 7%-43%; p=0.006).

A randomized, double-blind, placebo-controlled, cross-over withdrawal study was conducted to evaluate the efficacy and safety/tolerability of Gammagard Liquid in adult subjects (N=44) with multifocal motor neuropathy.³⁷ The study examined grip strength in the more affected hand (measured with dynamometer), and Guy's Neurological Disability Scale (GNDS) [upper limb part 6 subsection]. The Overall Disability Sum Score (ODSS) changed by -7.14% during placebo (indicating worsening of disability) and by -1.11% (indicating minimal change in disability) during treatment with Gammagard liquid. For this specific analysis of ODSS, lower scores represented more disability. With the dominant hand, subjects required 17% longer to complete the 9-hole peg test (a measure of dexterity) at the end of the placebo period, compared with baseline. By contrast, at the end of the Gammagard liquid treatment period, participants required 1.2% longer to complete the 9-hole peg test for the dominant hand compared with baseline. With the non-dominant hand, subjects required 33% longer to complete the 9-hole peg test at the end of the placebo period and 6.7% longer at the end of the Gammagard liquid treatment period, compared with baseline.

A randomized, double-blind, multi-center, investigator-initiated study compared IVIG (Aventis Behring LLC, King of Prussia, PA) with placebo 5% albumin.³⁸ On Days 1, 2, and 21, IVIG (1 g/kg) or placebo was given. The primary outcome measure was the change in muscle strength from baseline to Day 42, using the average muscle score (AMS). Secondary outcome measures included change from baseline AMS at Days 10 and 21, the Hughes' functional disability scale, forced vital capacity (FVC), and nerve conduction studies (NCS) of 4 motor nerves (median, ulnar, peroneal, and tibial). Total 33 patients were randomized. IVIG improved strength in patients with untreated CIDP by Day 10 with continued benefit through Day 42.

No randomized clinical trial has been performed to study IVIG in patients with PASC. Case studies of patients with PASC taking IVIG show improvements in symptom resolution.²⁸

13.8 CONTROL INFORMATION

Normal saline given intravenously will be the control (placebo) product.

13.8.1 FORMULATION, APPEARANCE, PACKAGING, AND LABELING

The normal saline will be of similar formulation and appearance to IVIG.

13.8.2 CONTROL DISPENSING, STORAGE, AND STABILITY

Store at controlled room temperature 20°C to 25°C (68°F to 77°F); excursions permitted between 15°C to 30°C (59°F to 86°F).

13.8.3 DOSING AND ADMINISTRATION

Control dosing and administration will occur according to Section 13.7.3 in order to maintain blinding.

13.9 EVENTS OF SPECIAL INTEREST

The following ESIs will be reported for IVIG:

- Aseptic meningitis
- Renal dysfunction (acute renal failure, osmotic nephrosis)
- Severe allergic reaction
- Thromboembolic events (DVT, PE, MI, ischemic stroke or other arterial thromboembolism)
- Acute hypertension greater than 180/110 (involving emergency visit or hospitalization)
- Episodes of anaphylaxis
- Episodes of other allergic reactions (rash, urticaria, shortness of breath, wheezing, tongue or lip swelling, injection site swelling and redness)
- Episodes of transfusion reactions (fever, chills, rigors)
- Episodes of acute renal failure unrelated to co-morbidities
- Episodes of aseptic meningitis
- Episodes of hemolytic anemia
- Episodes of acute lung injury with transfusion
- Episodes of thrombosis
- Any death
- Any other adverse reaction that resulted in the investigator or participant pausing or stopping study drug

13.10 ADDITIONAL APPENDIX-LEVEL HALTING AND STOPPING CRITERIA

- 1) Study drug infusion rate should be decreased by 50% if:
 - a. The participant has a concerning increase in symptoms of fatigue, aching, shortness of breath, nausea, headache, or substantial change in HR or BP or oxygen saturation during the infusion that is determined by the investigator to be due to study drug but not

representing an allergic reaction (but not representing an allergic reaction, in which case study drug should be interrupted or discontinued).

2) Study drug infusion should be interrupted if:

- a. The participant requests that the study drug be paused or stopped
- b. The investigator believes that for safety or tolerability reasons it is in the best interest of the participant to pause or stop study drug, for example for worsened shortness of breath and/or oxygen desaturation
- c. The participant develops a suspected allergic reaction; including new rash, new or worsened shortness of breath, or transfusion reaction during the infusion that is suspected to be related to the study medication

3) Redosing of study drug (subsequent weekly study drug dosing) should be interrupted (paused) if:

- a. The participant requests to interrupt study drug
- b. The investigator believes that, for safety reasons or tolerability reasons, it is in the best interest of the participant to interrupt study drug
- c. The participant develops any SUSAR
- d. The participant develops a suspected allergic or transfusion reaction that may be related to the study medication

13.11 STATISTICAL CONSIDERATIONS (APPENDIX-SPECIFIC)

Power calculation is provided for this IVIG appendix is based on a total of 200 enrolled participants. The power is 86% to detect 1-unit difference between 2 groups in change of OHQ/OIQ composite score if the missing data rate is 25% with a total sample size of 200. Table 3 provides variation of this assumption for a total sample size of 200 with corresponding powers for the primary endpoint.

We assumed that the missing data rate to range from 20% to 25% at month 9 based on the observed missing date of 30% at one year in the longitudinal study in POTS subjects.³⁹ The missing data rate of 25% is more likely in the trial with IVIG because of 9-month duration of intervention and the nature of intervention via injection

Table 3. Sample size and power calculation Power Calculation for the Primary Outcome, change of OHQ/OIQ composite score, from baseline to EOI with 0.05 type I error

Sample size	Missing data	Sample size	Clinically	Equal Group	Effect Size	Power
	rate	without	Significant	Standard		
		missing data	Difference	Deviation		
200	20%	160	1.0	2.0	0.5	88.2%
200	25%	150	1.0	2.0	0.5	86.0%
200	20%	160	1.0	2.1	0.48	84.9%

200	25%	150	1 / 1	1 71	0.48	82.6%
200	23/0	150	1.0	Z.1	0.40	02.070

For the secondary outcome, COMPASS-31, the power is 86% with a total sample size of 200 to detect 10-unit difference in change of COMPASS-31 from baseline to 9-month follow-up between the 2 groups assuming SD of 20 (equivalent to effect size of 0.5) and 25% missing data rate.

13.12 SCHEDULE OF PROCEDURES

Study Intervention Period Schedule of Activities for Appendix A; IVIG

	Study Intervention Period					
	Start of Study Intervention	3 Month Clinic Visit	6 Month Clinic Visit	End of Intervention (9 Month Clinic Visit)		
PROCEDURE	Day 0	Week 12 ± 1 week	Week 24 ± 1 week	Week 36 ± 1 week		
Concomitant Medications		X	X	X		
Pre and Post-Dose Administration ¹	х	х	х			
Study Drug Administration ^{1,5}	Х	Х	х			
Coordinated or Usual Non-pharmacologic Care ²	Х	Х				
OHQ/OIQ		х	Х	Х		
COMPASS-31		Х	Х	Х		
MAPS		Х	х	х		
voss		х	Х	Х		

	Study Intervention Period					
	Start of Study Intervention	3 Month Clinic Visit	6 Month Clinic Visit	End of Intervention (9 Month Clinic Visit)		
PROCEDURE	Day 0	Week 12 ± 1 week	Week 24 ± 1 week	Week 36 ± 1 week		
Active Stand Test		х	Х	Х		
6-minute Walk Test		X	X	X		
DSQ-PEM ³ (look back period "since last visit")		х	Х	Х		
PROMIS-29 + 2 Questionnaire		Х	Х	Х		
Skin Biopsy		pleted at baseline for a hy, the biopsy will be re		kin biopsy shows evidence		
Blood Collection ⁶				Х		
PASC Symptom Questionnaire		X	Х	X		
Vital Signs monitoring ⁷	Х					
Stool collection				Х		
Autonomic function tests at specialized sites				х		
Height and weight ⁸	х					

	Study Intervention Period					
				End of Intervention (9 Month Clinic Visit)		
PROCEDURE	Day 0	Week 12 ± 1 week	Week 24 ± 1 week	Week 36 ± 1 week		
Safety Assessment of AEs, including SAEs and ESIs ⁴	X	х	х	Х		

Abbreviations: EOI – End of Intervention, QOL – Quality of Life; PASC-Post-acute sequelae of SARS-CoV-2 infection, OHQ– Orthostatic hypotension questionnaire; OIQ- Orthostatic intolerance questionnaire; SAE – Serious Adverse Events; ESI – Events of Special Interest

¹ Pre-and post-dose administration and study drug administration will be done throughout the study intervention period

² Only applicable for those who are randomized to coordinated non-pharmacologic care. Not applicable to those receiving usual care. The coordinated non-pharmacologic care will be for a total duration of 3 months only.

³DSQ-PEM questionnaire will be a via phone or electronic survey within one day after strenuous in-person study visits.

⁴ Non-serious AEs will be reportable from start of study drug dosing to end of study drug administration.

⁵ The dosing frequency of 0.5 g/kg once a week is strongly suggested; however, the frequency can be modified based on individual site preference. The dose will be a **total of 2 g/kg every month** for 9 months.

⁶ Blood may be collected at any time as needed based on PI discretion and adverse events.

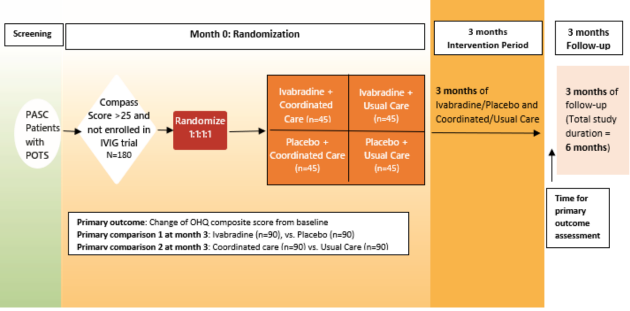
⁷ Vital signs monitoring will be done pre-, during, and post-infusion to ensure participant safety and monitor for adverse events.

⁸ Weight will be captured at each infusion visit to determine dosing.

14 APPENDIX B (IVABRADINE)

14.1 STUDY DESIGN

Figure 3. Study Schema for Appendix B (Ivabradine)



^{*}In this factorial design, participants will be randomized 1:1:1:1 to the combination of study drug/placebo and coordinated/usual care.

As depicted in **Figure 3**, approximately 180 participants will be enrolled in Appendix B (Ivabradine), based on eligibility criteria.

In the ivabradine study, randomization will be implemented with 1:1:1:1 allocation among the combinations of ivabradine/placebo and coordinated/usual care. All participants receive either ivabradine or placebo for 3 months (12 weeks) with a follow-up period for an additional 3 months (total study duration of 6 months).

All participants will receive coordinated non-pharmacologic care or usual non-pharmacologic care (control) for a duration of 3 months. This will be concurrent while taking ivabradine (or placebo). coordinated non-pharmacologic care involves volume expansion through high salt diet (6-10 grams of salt), water intake (at least 2 liters per day), abdominal binder, exercise/rehabilitation, weekly monitoring of BP, motivation, education, and assisted care through care coordinator.

The care coordinator will call each patient weekly during the 3 month non-pharmacologic intervention period to provide education and support on lifestyle modifications. Patients will be asked to maintain a weekly log of non-pharmacological care (average dietary intake of salt and water, exercise regimen) and provide a summary to the care-coordinators during the weekly calls. Further details of the non-pharmacologic intervention are provided in Section 6.6 and contained in the MOP.

Autonomic function tests are optional for this appendix but recommended as per the platform protocol schedule of procedures.

14.2 INTERVENTION RATIONALE

Ivabradine is a medication used to treat chronic heart failure with elevated HR. It reduces the HR without reducing contractility or BP. It has the greatest activity and most potent effect when the HR is elevated. It increases coronary flow reserve and collateral perfusion, promoting the development of coronary collaterals and improving endothelial function. In addition, it decreases norepinephrine levels in patients with POTS. Ivabradine has been successfully used to treat symptoms such as tachycardia and POTS in a small randomized trial, and its use has been associated with improvement in symptoms in patients with PASC and POTS.³⁴

14.3 RISK ASSESSMENT

Serious and unexpected AEs may occur that have not been previously reported with ivabradine use. Previously reported AEs were identified in the SHIFT study, which was a randomized, placebo-controlled trial in patients with systolic heart failure. Common AEs included (incidence ≥ 1.0% higher on ivabradine than placebo and occurring in > 1% on ivabradine) included bradycardia, hypertension, atrial fibrillation, and phosphenes (visual brightness). Following approval of ivabradine, the following AEs have been reported: syncope, hypotension, torsade de pointes, ventricular fibrillation, ventricular tachycardia, angioedema, erythema, rash, pruritus, urticaria, vertigo, and diplopia, and visual impairment.

No ivabradine human data are currently available to evaluate for a drug-associated risk of major birth defects, miscarriage, adverse maternal or fetal outcomes, or lactation effects. The risk/benefit of treatment with ivabradine for PASC has not been established in pregnant women.

14.4 ADDITIONAL APPENDIX-LEVEL INCLUSION CRITERIA

- Abnormal active standing test defined as presence of orthostatic tachycardia (an increase of 30 beats per minute (bpm) or more in HR within 10 minutes upon standing without orthostatic hypotension) and experiencing orthostatic symptoms
- 2. COMPASS-31 Score > 25 and not enrolled in the IVIG appendix

14.5 ADDITIONAL APPENDIX-LEVEL EXCLUSION CRITERIA

- 1. A person of child-bearing potential who is not taking effective contraception
- 2. Use of the following medications: clonidine, tizanidine, amphetamines, and serotonin and norepinephrine reuptake inhibitors (SNRIs) with the exception of modafinil
- 3. Use of beta-blockers (any formulation), calcium channel blockers, midodrine, pyridostigmine, fludrocortisone, and guanfacine will be excluded unless participant is on a stable dose (>4

weeks). Participants on stable doses will be allowed to continue the medication throughout the study.

- 4. Combination with verapamil or diltiazem which are moderate CYP3A4 inhibitors with heart rate reducing properties
- 5. Lactating and breast-feeding women
- 6. Severe hepatic impairment
- 7. Use of drugs known to prolong the QT-interval (e.g., quinidine, disopyramide, bepridil, sotalol, amiodarone, pimozide, ziprasidone, sertindole, mefloquine, halofantrine, pentamidine
- 8. Concomitant use of digoxin
- 9. Participants who are pacemaker dependent
- 10. Patients with hypokalemia (serum K+<3.5 mEq/L)
- 11. Patients taking potassium-depleting diuretics
- 12. A history of congenital or acquired long QT syndrome, with or without torsade de pointes
- 13. Patients with high degree AV block such as Mobitz II

14.5.1 PRECAUTIONS

The healthcare provider should consult other appropriate resources such as the prescribing information for comprehensive information on dosing and monitoring.

Initiation of medications that inhibit or induce CYP3A may increase or decrease concentrations of ivabradine, respectively. These interactions may lead to:

- Clinically significant adverse reactions, potentially leading to severe, life-threatening, or fatal events from greater exposures of concomitant medications
- Clinically significant adverse reactions from greater exposures of ivabradine
- Loss of therapeutic effect of ivabradine

Concomitant administration of ivabradine with negative chronotropes (such as calcium channel blockers and beta-blockers) increases the risk of bradycardia. Therefore, close monitoring of heart rate is required.

14.6 CONCOMITANT MEDICATIONS

Participants who are already on stable (>4 weeks) doses of beta-blockers (any formulation), calcium channel blockers, midodrine, pyridostigmine, fludrocortisone, and guanfacine will be allowed to continue the medication throughout the study. During the study, initiation of the following medications will not be allowed for more than 2 weeks:

- Beta-blockers (any formulation)
- Midodrine
- Calcium channel blockers

Pyridostigmine

14.7 IVABRADINE INFORMATION

Ivabradine is a hyperpolarization-activated cyclic nucleotide-gated channel blocker. In April 2015, the US FDA approved ivabradine for use in adult and pediatric patients to treat stable symptomatic heart failure with an ejection fraction of ≤35%. Currently, ivabradine is not FDA approved for the treatment of POTS or PASC.

14.7.1 FORMULATION, APPEARANCE, PACKAGING, AND LABELING

Ivabradine 2.5 mg capsules for oral administration, equivalent to 2.5 mg of ivabradine hydrochloride will be used. The capsule will consist of the 2.5 tablet plus fillers (microcrystalline cellulose).

The tablet contains the following inactive ingredients: betadex, microcrystalline cellulose, croscarmellose sodium, and magnesium stearate. The film-coating contains hypromellose, lactose monohydrate, titanium dioxide, macrogol 4000, iron oxide red, iron oxide yellow, and iron oxide black.

14.7.2 DRUG DISPENSING, STORAGE, AND STABILITY

Store at controlled room temperature 20°C to 25°C (68°F to 77°F); excursions permitted between 15°C to 30°C (59°F to 86°F).

14.7.3 DOSING AND ADMINISTRATION

The starting dose will be 5 mg twice a day, and the dose will be modified if needed at the 1 month clinic visit. At this visit, HR will be measured and the dose will be modified as applicable. The maximum dose will be 7.5 mg twice daily if HR is > 90 bpm.

The table below provides the dosing of ivabradine based on HR.

Supine Resting HR 60-80	2.5 mg BID
Supine Resting HR >80	5 mg BID
Supine Resting HR >90	7.5 mg BID

^{*}Resting HR should be measured 5 minutes after lying down

During the study intervention period, the study sites will contact participants once a week to monitor compliance with study drug administration.

Ivabradine tablets should generally be taken with food and the tablets should be swallowed whole, not chewed, broken, or crushed.

If the participant misses a dose of ivabradine within 8 hours of the time it is usually taken, the participant should take it as soon as possible and resume the normal dosing schedule. If the participant misses a dose by more than 8 hours, the participant should not take the missed dose but instead take the next dose at the regularly scheduled time. The participant should not double the dose to make up for a missed dose.

Overdose (defined as an excess of 7.5 mg twice a day) may lead to severe and prolonged bradycardia. In the event of bradycardia with poor hemodynamic tolerance, temporary cardiac pacing may be required. Supportive treatment, including intravenous (IV) fluids, atropine, and intravenous beta-stimulating agents such as isoproterenol, may be considered.

In the event of an overdose, the investigator should:

- Closely monitor the participant for any AEs/SAEs and laboratory abnormalities for at least 5 half-lives or 28 calendar days after the overdose of study intervention (whichever is longer).
- 2. Document the quantity of the excess dose as well as the duration of the overdose in the CRF.
- 3. If there is an associated SAE, the SAE is reported within the CRF within 24 hours of site awareness.

Decisions regarding dose interruptions or modifications will be made by the site investigator based on the clinical evaluation of the participant.

14.7.4 RATIONALE FOR SELECTION OF DOSE

Pre-clinical studies:

There was no evidence of carcinogenicity when mice and rats received ivabradine up to 104 weeks by dietary administration. High doses in these studies were associated with mean ivabradine exposures of at least 37 times higher than the human exposure (AUC_{0-24hr}) at the maximum recommended human dose.

Clinical studies:

In the SHIFT trial, all subjects were initiated on ivabradine 5 mg (or matching placebo) twice daily and the dose was increased to 7.5 mg twice daily or decreased to 2.5 mg twice daily to maintain the resting HR between 50 and 60 bpm.

No pharmacokinetic differences have been observed in elderly (≥ 65 years) or very elderly (≥ 75 years) patients compared to the overall population. However, ivabradine has only been studied in a limited

number of patients ≥ 75 years of age.

No dose adjustment is required in patients with mild or moderate hepatic impairment.

No dosage adjustment is required for patients with creatinine clearance 15 to 60 mL/min.

No randomized clinical trial has been performed to study ivabradine in patients with PASC. Case studies of patients with PASC taking ivabradine show improvements in symptom resolution.⁴⁰

14.8 CONTROL INFORMATION

The control (placebo) oral tablets will be similar to the study drug, ivabradine.

14.8.1 FORMULATION, APPEARANCE, PACKAGING, AND LABELING

The placebo will be salmon-colored, film-coated tablets, that match the appearance of ivabradine.

The control packaging matches the packaging described in Section 14.7.1.

14.8.2 CONTROL DISPENSING, STORAGE, AND STABILITY

Store at controlled room temperature 20°C to 25°C (68°F to 77°F); excursions permitted between 15°C to 30°C (59°F to 86°F).

14.8.3 DOSING AND ADMINISTRATION

Control dosing and administration will occur according to Section 14.7.3 in order to maintain blinding.

14.9 EVENTS OF SPECIAL INTEREST

The following ESIs will be reported for ivabradine:

- Bradycardia
- Hypertension
- Atrial fibrillation
- Luminous phenomena (phosphenes)

14.10 STATISTICAL CONSIDERATIONS (APPENDIX-SPECIFIC)

Power calculation is provided for this Ivabradine appendix is based on a total of 180 enrolled participants. The power is 86.5% to detect 1-unit difference between two groups in change of OHQ/OIQ composite score if the missing data rate is 15% with a total sample size of 180. Table 4 provides variation of this assumption for a total sample size of 180 with corresponding powers for the primary endpoint.

We assumed that the missing data rate to be 15% at month 9 based on the observed missing date of 30% at one year in the longitudinal study in POTS subjects.³⁹

Table 4. Sample size and power calculation Power Calculation for the Primary Outcome, change of OHQ/OIQ composite score, from baseline to EOI with 0.05 type I error

Sample size	Equal or Unequal Group Missing data rate(s)	Total or group Sample size(s) without missing data	Clinically Significant Difference	Equal or Unequal Group Standard Deviation(s)	Power
180	15%	153	1.0	2.0	86.5%
180	20%	144	1.0	2.0	84.6%
180	15%	153	1.0	2.1	83.1%
180	20%	144	1.0	2.1	81.0%

For the secondary outcome, COMPASS-31, the power is 86.9% to detect 10-unit difference in change of COMPASS-31 from baseline to 3-month follow-up between the two groups assuming SD of 20 (equivalent to effect size of 0.5) and 15% missing data rate.

14.11 SCHEDULE OF PROCEDURES

Study Intervention Period Schedule of Activities for Appendix B: Ivabradine

	Study Intervention Period				
	Start of Study Intervention	1 Month Clinic Visit	End of Intervention (3 Month Clinic Visit)		
PROCEDURE	Day 0	Week 4 ± 1 week	Week 12 - 1 week		
Concomitant Medications		X	х		
Study Drug Administration ¹	Х	Х			
Coordinated Non- pharmacologic Care ²	х	X	Х		
Weekly site calls ³	Х	Х			
OHQ/OIQ Assessment		Х	Х		
COMPASS-31		X	Х		
MAPS		Х	х		
Stool Collection			Х		
Active Stand Test and Vanderbilt Orthostatic Symptoms Score		Х	х		
6-minute Walk Test		Х	Х		

	Study Intervention Period		
	Start of Study Intervention	1 Month Clinic Visit	End of Intervention (3 Month Clinic Visit)
PROCEDURE	Day 0	Week 4 ± 1 week	Week 12 - 1 week
DSQ-PEM ⁴			
(look back period "since last visit")		X	X
PROMIS-29 + 2 Questionnaire		Х	х
Blood Collection ⁶			х
PASC Symptom Questionnaire		Х	х
Autonomic function tests at specific sites			Х
Safety Assessment of AEs, including SAEs and ESIs	Х	х	х
Vital signs monitoring ⁷		Х	
Adherence ⁵	Х	X	Х

Abbreviations: EOI – End of Intervention, QOL – Quality of Life; PASC-Post-acute sequelae of SARS-CoV-2 infection, OHQ– Orthostatic hypotension questionnaire; OIQ- Orthostatic intolerance questionnaire; AE – Adverse Events; SAE – Serious Adverse Events; ESI – Events of Special Interest

¹Study drug administration will occur every day during the intervention period

² Only applicable for those who are randomized to coordinated non-pharmacologic care. Not applicable to those receiving usual care. The coordinated non-pharmacologic care will be for a total duration of 3 months only.

³Participants will receive weekly phone calls for the first 30 days of the study intervention period

⁴ DSQ-PEM questionnaire will be via phone or electronic survey within one day after strenuous in-person study visits.

⁵ Adherence to drug (pill count) will be documented.

⁶ Blood collection may be collected at any time as needed based on PI discretion and adverse events.

⁷ HR will be captured at the 1 month clinic visit to determine dosing.