Antivirals for adult patients hospitalized with SARS-CoV-2 infection: A randomized, Phase II/III, multicenter, placebo-controlled, adaptive study, with multiple arms and stages. COALITION COVID-19 BRAZIL IX – REVOLUTION: protocol and statistical analysis plan

ABSTRACT

Repurposed drugs are important in resource-limited settings because the interventions are more rapidly available, have already been tested safely in other populations and are inexpensive. Repurposed drugs are an effective solution, especially for emerging diseases such as COVID-19. The REVOLUTION trial has the objective of evaluating three repurposed antiviral drugs, atazanavir, daclatasvir and sofosbuvir, to provide evidence on whether these medications are capable of decreasing the SARS-CoV-2 load in patients with COVID-19. The REVOLUTION trial is designed to first identify whether any of these drugs alone or in combination reduce the viral load. If they do, a Phase III trial will be tested simultaneously in a Phase II trial to determine if these medications are capable of increasing the number of days free of respiratory support. Participants must be hospitalized adults aged ≥18 years with initiation of symptoms ≤9 days and SpO2 ≤94% in room air or a need for supplemental oxygen to maintain an SpO2 > 94%. The expected total sample size ranges from 252 to 1,005 participants, depending on the number of stages that will be completed in the study. Hence, the protocol is described here in detail together with the statistical analysis plan. In conclusion, the REVOLUTION trial is designed to provide evidence on whether atazanavir, daclatasvir or sofosbuvir decrease the SARS-CoV-2 load in patients with COVID-19 and increase the number of days patients are free of respiratory support. In this protocol paper, we describe the rationale, design, and status of the trial.

Keywords: COVID-19; Coronavirus infections; Antiviral agents; Protocol; Respiratory insufficiency; Daclatasvir; Sofosbuvir

ClinicalTrials.gov identifier: NCT04468087

INTRODUCTION

Background and rationale

Coronavirus disease 2019 (COVID-19) reached a pandemic status, and several approaches have been suggested to control severe acute respiratory
syndrome coronavirus 2 (SARS-CoV-2) replication in patients with moderate to severe cases. One of these strategies includes the repurposing of antiviral drugs used to treat severe acute respiratory syndrome coronavirus 1 (SARS-CoV-1) and Middle East respiratory syndrome coronavirus (MERS-CoV), including antiretroviral agents.\(^1\) Drugs such as remdesivir, lopinavir and ritonavir, darunavir and cobicistat, umifenovir and oseltamivir potentially exert therapeutic effects against COVID-19.\(^2\) However, among these repurposed drugs, only remdesivir showed clinical benefits for COVID-19 treatment.\(^3\)

Other specific and nonspecific antivirals have been proposed to treat the disease. Some are based on the results from in vitro and clinical studies conducted with a small sample size in specific populations. Atazanavir (ATV) blocked the major protease activity of SARS-CoV-2 in a protease-free cell assay;\(^4\) sofosbuvir (SOF) showed EC\(_{50}\) values against SARS-CoV-2 replication of 6.2 and 9.5\(\mu\)M in HuH7 (hepatoma) and Calu-3 (type II pneumocytes) cells, respectively;\(^5\) and daclatasvir (DCV) consistently inhibited the production of SARS-CoV-2 infectious particles in Vero cells, the HuH-7-cell line and Calu-3 cells, preventing the induction of interleukin (IL) 6 and tumor necrosis factor alpha (TNF-\(\alpha\)) production.\(^5\) Available clinical studies are synthesized in recent meta-analysis, SOF/DCV may reduce the mortality rate and need for intensive care unit (ICU)/invasive mechanical ventilation (IMV) in patients with COVID-19 while increasing the chance for clinical recovery with a low to moderate quality of evidence.\(^6\) However, no studies investigating DCV or ATV alone have been published.

Implementing studies that allow more than one new treatment to be tested simultaneously may be advantageous over classic parallel group studies. The main objectives of this type of clinical trial are to quickly reject any new therapies that do not seem to be better than controls and to identify those that are significantly better in terms of clinical outcomes.\(^7\) Thus, we propose a randomized, placebo (PbO)-controlled, adaptive, multiarm, multistage Phase II trial to evaluate ATV, DCV and DCV plus SOF simultaneously to first identify whether any of these drugs alone or in combination reduce the viral load. If they do, a Phase III trial will be initiated to investigate clinical outcomes.

**Objectives**

The main objective of the REVOLUTION trial is to evaluate whether repurposed antiviral drugs alone or in combination are effective at decreasing the viral load and increasing the number of days free of respiratory support. The primary and secondary objectives are described in table 1.

**Study design**

A randomized, adaptive, PbO-controlled, multiarm, multistage trial conducted in 3 continuous stages (ClinicalTrials.gov Identifier: NCT04468087); version 4.0 of Protocol 10/07/2020. The first two stages are Phase II studies, and the third is a Phase III study, as shown in figure 1.

**Methods**

**Study settings**

The study will be conducted in approximately 60 Brazilian hospitals.
### Table 1 - Objectives and outcomes

| Primary objective                                                                 | Outcome/primary variables                                                                                                                                                                                                                                                                                                                                 |
|----------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Phase II first stage (II/1): compare the effect of treatment with single antivirals against placebo in reducing the load of SARS-CoV-2 in nasopharyngeal swab samples | Decay rate (slope) of the SARS-CoV-2 viral load logarithm in nasopharyngeal and swab samples evaluated at D0, D3, D6 and D10 after randomization                                                                                                                                                                      |
| Phase II second stage (II/2): compare the effect of treatment with combinations of antivirals compared to isolated ones in reducing the SARS-CoV-2 viral load in nasopharyngeal swab samples | Decay rate (slope) of the SARS-CoV-2 viral load logarithm in nasopharyngeal and swab samples evaluated at D0, D3, D6 and D10 after randomization                                                                                                                                                                      |
| Phase III: compare the efficacy of antivirals alone or in combination to the placebo in increasing the number of days free of respiratory support | Days free of respiratory support, defined as the number of days without oxygen, noninvasive ventilation/high-flow nasal cannula or the need for mechanical ventilation within 15 days from randomization  
1. This parameter is counted as follows:  
D = zero (if the patient dies within 15 days (either in the hospital or at home or remains on respiratory support with oxygen through a nasal catheter, noninvasive ventilation, high-flow nasal catheter, or mechanical ventilation ≥ 15 days)  
D = 15 - x (if the patient is released from the hospital in < 15 days, where x represents the number of days with respiratory support during hospitalization) |

| Secondary objectives | Outcomes/secondary variables |
|----------------------|-----------------------------|
| Evaluate the status using the 7-stage ordinal scale for clinical outcomes on D15 | Percentage of patients in various stages:  
1. Not hospitalized with resumption of normal activities  
2. Not hospitalized, but unable to resume normal activities  
3. Hospitalized, with no need for supplemental oxygen  
4. Hospitalized, requiring supplemental oxygen  
5. Hospitalized, requiring high-flow nasal oxygen therapy, noninvasive mechanical ventilation, or both  
6. Hospitalized, requiring blood oxygenation through a membrane system, invasive mechanical ventilation or both  
7. Death |
| Evaluate the status using the 6-stage ordinal scale for clinical outcomes on D7 | Percentage of patients in various stages:  
1. Nonhospitalized  
2. Hospitalized, with no need for supplemental oxygen  
3. Hospitalized, requiring supplemental oxygen  
4. Hospitalized, requiring high-flow nasal oxygen therapy, noninvasive mechanical ventilation, or both  
5. Hospitalized, requiring blood oxygenation through a membrane system, invasive mechanical ventilation or both  
6. Death |
| Evaluate 28-day mortality | Percentage of deaths in 28 days |
| Evaluate the number of days free from mechanical ventilation within 28 days | D = 28 - number of days requiring mechanical ventilation  
D = zero if death occurs or the patient continues to require mechanical ventilation after 28 days |
| Evaluate the number of days out of the hospital within 28 days | D = 28 - number of days after admission to the hospital  
D = zero if death occurs or the patient remains hospitalized after 28 days |
| Evaluate the time to discharge | Number of days from randomization to discharge, within 28 days  
D = 28 - number of days from randomization to hospital discharge  
D = zero if death occurs or the patient remains hospitalized after 28 days |
| Evaluate the number of days free of respiratory support within 15 days for Phases II/1 and II/2 | D = 15 - number of days with respiratory support on hospitalization  
D = zero if death occurs or the patient remains hospitalized with a need for respiratory support defined as the use of low-flow, high-flow oxygen, IMV, or MV in 15 days |

**Safety objective**

| Outcomes/safety variables |
|---------------------------|
| Evaluate Grade 2, 3 or 4 adverse events, which were not present at the patient’s entrance, defined by the Division of AIDS table for Grading the Severity of Adult and Pediatric Adverse Events | Percentage of Grade 2, 3, or 4 adverse events in the Division of AIDS table |
| Evaluate serious adverse events | Percentage of serious adverse events |
| Evaluate discontinuation of study drug-related treatment | Percentage of patients who needed to discontinue the intervention (study drug) |

D - day; IMV - invasive mechanical ventilation; MV - mechanical ventilation.
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Interventions

Intervention drugs administered in each stage

**Phase II, first stage** - six arms with six different interventions will be designed:
1. ATV.
2. DCV.
3. SOF/DCV.
4. PbO ATV.
5. PbO DCV.
6. PbO SOF/DCV.

**Phase II, second stage** - four arms with four different interventions will be designed:
1. Best of the first stage.
2. Best of the first stage + the second best.
3. PbO best first stage.
4. PbO best first stage + PbO second best.

**Phase III, third stage** - two arms with two different interventions will be designed:
1. Best antiviral alone or in combination (defined in Stage 1 or 2).
2. PbO best antiviral alone or in combination.

Eligibility criteria

The inclusion and exclusion criteria are described in table 2. Procedures for the initial diagnosis of COVID-19, collection of biological material, viral identification, genetic sequencing and viral isolation in culture are explained in Supplementary material 1.

### Table 2 - Inclusion and exclusion criteria

| Inclusion criteria                                                                 | Exclusion criteria                                                                 |
|-----------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|
| 1. Adults (≥ 18 years) hospitalized with COVID-19 with one of the following conditions: | Presence of any of the following conditions:                                        |
| - Positive RT-PCR test for SARS-CoV-2 OR                                           | - Patients requiring invasive mechanical ventilation                                |
| - Typical clinical history AND chest CT with typical findings, pending the results of the RT-PCR test for SARS-CoV-2 | - ALT or AST level > 5 times the upper limit of normal                               |
| 2. Time between symptom onset and inclusion ≤ 9 days                               | - Total bilirubin level > 2mg/dL                                                   |
| 3. SpO₂ ≤ 94% in room air or need for supplemental oxygen to maintain an SpO₂ > 94% | - Platelet count < 50,000 cells/L                                                  |
| 4. The patient consents to participate in the study and is willing to comply with all study procedures, including the collection of virology samples | - Total neutrophil count < 750 cells/L                                              |
|                                                                                  | - Renal failure (eGFR < 30mL/min/1.73m², using the MDRD or CKD-EPI method);         |
|                                                                                  |   and predefined renal failure Stage 3 according to the AKIN™ classification with a serum creatinine level > 4mg/dL or patient already on renal replacement therapy |
|                                                                                  | - History of liver disease with moderate to severe impairment (liver cirrhosis with Child Pugh B and C classification) previously known* |
|                                                                                  | - Decompensated congestive heart failure*                                          |
|                                                                                  | - Pregnant or breastfeeding patients                                                |
|                                                                                  | - Known allergy or hypersensitivity to any study drug                              |
|                                                                                  | - Carrier of hepatitis C [positive HCV RNA], active hepatitis B (positive surface antigen in the past), or HIV (ELISA and confirmatory Western blot in the past) |
|                                                                                  | - Patients currently using nucleoside or nucleotide analog drugs for any indication |
|                                                                                  | - Corrected QT interval > 480 on the electrocardiogram                              |
|                                                                                  | - Heart rate < 55 bpm                                                              |
|                                                                                  | - Patients using or who recently used (< 90 days) amiodarone                        |
|                                                                                  | - Women of childbearing potential* or men with a partner of childbearing potential who do NOT agree to use two contraceptive methods (including barrier method) for 100 days |

RT-PCR - real time polymerase chain reaction; CT - computed tomography; SpO₂ - oxygen saturation; ALT - alanine aminotransferase; AST - aspartate aminotransferase; eGFR - estimated glomerular filtration rate; MDRD - Modification of Diet in Renal Disease; CKD-EPI - Chronic Kidney Disease Epidemiology Collaboration; AKIN - Acute Kidney Injury Network; HCV - hepatitis C virus; ELISA - enzyme-linked immunosorbent assay. * Defined in Supplementary material 1.

Figure 1 - Study flowchart.

ATV - atazanavir; SOF - sofosbuvir; DCV - daclatasvir.

**Eligibility criteria**

The inclusion and exclusion criteria are described in table 2. Procedures for the initial diagnosis of COVID-19, collection of biological material, viral identification, genetic sequencing and viral isolation in culture are explained in Supplementary material 1.
The doses and administration routes of the study drugs are described in Table 3. If swallowing is impossible for any reason, the protocol provides for the use of the study drugs via a nasogastric or nasoenteral tube. The study drugs will be donated by pharmaceutical companies, with ATV (and PbO) donated by Fundação Oswaldo Cruz (Fiocruz) and DCV, SOF and respective PbOs donated by Blanver Farmoquímica e Farmacêutica.

**Discontinuation of the drug treatment and safety criteria**

The set of primary analyses for safety and efficacy will include individuals who received at least one dose of the study drug. Emerging treatment data will be analyzed and defined as data collected from the administration of the first study drug dose to the date of administration of the last study drug dose plus 30 days.

At medical discretion, the study drug(s) will be discontinued if a research participant meets one of the following criteria:

1. The researcher considers that the discontinuation of the study drug is in the best interest of the research participant.
2. Elevated levels of alanine aminotransferase (ALT) or aspartate aminotransferase (AST) greater than ten times the value of the upper limit of normal (ULN). Elevations in ALT or AST levels often follow the clinical course of COVID-19, whether they are due to shock, sepsis or even direct SARS-CoV-2 infection in liver cells. These changes may occur in the two study groups: active drug or PbO.
3. Elevated levels of ALT or AST greater than 3x the ULN WITHOUT an increase in alkaline phosphatase levels confirmed in a new test performed within 48 hours AND the presence of one of the following conditions:
   a. Total bilirubin level > 2 times the ULN.
   b. International Normalized Ratio (INR) > 2.
4. Elevated ALT or AST levels greater than 3x the ULN accompanied by the presence of the following two conditions:
   a. Onset or worsening of fatigue, nausea, vomiting, discomfort in the upper abdomen, or fever.
   b. Rash and/or peripheral eosinophilia (> 5%).
   c. Any Grade 3 or greater rash accompanied by symptoms.
   d. Any Grade 4 AE or laboratory abnormality considered related to the study drug.

**Discontinuation of one arm of the study for safety**

The criteria for discontinuing one arm of the study for safety are the occurrence of one of the following 3 conditions:

1. Presence of the components of Hy’s Law\(^{11}\) which consists of the presence of all of the following conditions confirmed by an adjudicating committee of 2 independent evaluators:
   a. The study drug causes a higher incidence of hepatocellular injury confirmed by a > three times ULN increase in ALT and AST levels than the PbO group.
   b. Increase in ALT or AST levels > three times ULN and elevation of total bilirubin levels > two times ULN without signs of cholestasis (elevated alkaline phosphatase levels).
   c. No other explanation for the combination of the changes described above, such as the presence of hepatitis A, B, C, use of vasopressors or IMV; acute or preexisting hepatic disease or the presence of another drug capable of causing the observed injury.
2. Elevated ALT and AST levels > five, ten or twenty times ULN more frequently in the treated group than in the control group after 48-hour reassessments.
3. Presence of more serious adverse events (SAEs) defined in Section S.5 (Supplementary material 1) in the treatment arm than in the PbO arm.

Patients with COVID-19 may progress to sepsis-associated liver injury, which makes characterizing all the Hy Law criteria in these patients very challenging. For this study, we will consider a modified definition of Hy’s Law that will be present when the 3 criteria described above are met in the absence of shock or severe acute respiratory failure with the need for mechanical ventilation.

### Table 3 - Study drug dosage

| Drug                  | Pharmaceutical form | Route of administration/instructions for use | Frequency of administration | Conservation            |
|-----------------------|---------------------|---------------------------------------------|----------------------------|-------------------------|
| Daclatasvir/placebo DCV | Coated tablets/28 tablets bottles* | Orally with or without food | 2 tablets 1 once daily on D1, 1 tablet once daily from D2 to D10 | Room temperature 15 to 30°C |
| Sofosbuvir/placebo SOF | Coated tablets/28 tablets bottles* | Orally with or without food | 1 tablet twice daily on D1, 1 tablet once daily from D2 to D10 | Room temperature 15 to 30°C |
| Atazanavir/placebo ATV | Capsules with packs of 30 capsules* | Orally with food | 2 capsules twice daily on D1, 1 capsule twice daily from D2 to D10 | Room temperature 15 to 30°C |

DCV - daclatasvir; SOF - sofosbuvir; ATV - atazanavir; D - day. *Coated tablets for daclatasvir and sofosbuvir treatments and placebos and capsules for atazanavir treatment and placebo are identical in physical appearance.
Rules for changing the study stage (Figure 2)

1. If ATV and a presentation of DCV alone are effective and safe in the first stage, a second stage will be performed with the first- and second-best drugs from the previous stage combined according to the rate of decay of the viral load logarithm (slope). The drug with the highest decay rate will be administered first, followed by the drug with the second highest decay rate. The third phase will investigate the drug with the highest decay rate in Phase II/1 or Phase II/2.

2. If only DCV presentations are effective and safe, a Phase II/2 trial will not be conducted, and Phase III will start with the formulation of DCV alone or combined with SOF which produces the higher rate of decay and it is safe according to the interpretation of the DSMB.

3. If only ATV is effective and safe, a Phase II/2 will not be conducted, and Phase III starts with ATV alone.

4. If none of the study drugs are effective and safe in Phase II/1, the study ends without conducting Phase II/2 and/or Phase III.

7. Transfer to advanced units (ward and ICU) according to the clinical judgment and patient’s need.

8. The addition of other therapies, such as antibiotics, corticosteroids, IL-6 receptor inhibitors and other antivirals, as indicated by the attending physician.

Study intervention compliance

- Hospitalized patients: Each dose of the study drug will be administered and supervised by a member of the clinical research team who is appropriately trained. The administration date and time will be entered into the case report form (CRF).

- Out-of-hospital patients: Patients who are discharged before the end of treatment will have their drug dispensed for home use and be monitored by telephone interviews, a medication diary and physical reevaluation in return visits on the missing day from the 4 prespecified schedule times (baseline and Days - D - 3, 6 and 10). Assessments of study medication use will be performed at each study visit. The subject should be instructed to bring all unused study drugs to each visit and any empty bottles. The dates and number of tablets dispensed and returned must be recorded on the drug accountability form.

Drug accountability

The principal investigator at each site may delegate responsibility for study product accountability to the research pharmacist at the participating site. He/she will be responsible for maintaining complete records and documentation of study product receipt, accountability, dispensation, storage conditions, and final disposition of the study product(s). The time of study drug administration to the subject will be recorded on the appropriate data collection form (CRF). All study product(s), whether administered or not, must be documented on the appropriate study product accountability record or dispensing log available at the pharmacy of each center. The Coordinating Center team will verify the study product accountability records and dispensing logs of each participating site.

Concomitant therapy and drug interaction

Therapy administered before enrollment with any other experimental treatment or off-label use of marketed medications must be discontinued upon enrollment. Concomitant therapy will be recorded daily until D10. A subject cannot participate in another clinical trial or receive any experimental treatment other than the study drugs for the treatment of COVID-19. Some concomitant medications must be stopped or adjusted according to table 4 due to drug interactions.

Figure 2 - Rules for changing stages.

SOF - sofosbuvir; ATV - atazanavir; DCV - daclatasvir.

All participants’ daily routines will include

1. A clinical evaluation by the attending physician.
2. Routine laboratory tests at the discretion of the attending physicians.
3. Respiratory and motor physiotherapy.
4. Surveillance of vital signs.
5. Addition of ventilatory support measures, such as increased oxygen flow, use of noninvasive ventilation or high-flow nasal cannula, as recommended by the attending physicians.
6. Prophylaxis of stress ulcers and venous thromboembolism according to the protocol of each institution.
| Drug                  | SOF | DCV  | ATV       | Effects                                      | Action                                           |
|----------------------|-----|------|----------|----------------------------------------------|-------------------------------------------------|
| Amiodarone           | X   | X    | X        | Bradycardia when administered with another antiviral | Do not administer. Switch to propafenone         |
| Digoxin              |     | X    |          | ↑Digoxin                                      | Avoid administration. If necessary, decrease the dose by 30% |
| Quinidine            | X   | X    | ↑Quinidine| A dose adjustment is not needed (when not using ritonavir). Avoid use with DCV |                                                   |
| Lidocaine            |     |      | X        | A dose adjustment is not needed (when not using ritonavir) |                                                 |
| Carbazamazepine      | X   | X    | X        | ↓Efficacy SOF                                 | Switch to 500mg of levetiracetam twice a day IV or VO |
| Phenytoin            |     |      |          |                                                                 |                                                 |
| Phenobarbital        |     |      |          |                                                                 |                                                 |
| Lamotrigine          | X   |      |          | ↓Lamotrigine if administered with ritonavir      | A dose adjustment is not needed (when not using ritonavir) |
| Rifampicin           | X   | X    | X        | ↓Efficacy SOF and DCV                         | Do not administer or suspend RMP for 10 days     |
| Itraconazole         |     | X    | ↑ATV     |                                                                 |                                                 |
| Ketoconazole         | X   | X    | ↑itraconazole      |                                                                 |                                                 |
| Voriconazole         | X   | X    | ↑Ketoconazole   |                                                                 |                                                 |
| Dexamethasone        |     |      |          | ↓Efficacy DCV                                 | Do not adjust dose. All patients will use this drug |
| Bosentan             |     |      |          |                                                                 |                                                 |
| Dabigatran           | X   | X    | ↑Dabigatran  |                                                                 | Substitute for enoxaparin                       |
| Rivaroxaban          |     |      | ↑Rivaroxaban   |                                                                 |                                                 |
| Apixaban             |     |      | ↑Apixaban   |                                                                 |                                                 |
| Warfarin             | X   | X    | ↑Warfarin   |                                                                 | Monitor INR for a warfarin dose adjustment or switch to enoxaparin/heparin |
| Atorvastatin         | X   | X    | ↑Atorvastatin | Temporary suspension: risk of myopathy or statin dose reduction by 50% |                                                   |
| Pravastatin          |     |      | ↑Pravastatin   |                                                                 |                                                 |
| Simvastatin          |     |      | ↑Simvastatin   |                                                                 |                                                 |
| Quetiapine           |     |      | ↑Quetiapine   | Decrease quetiapine dose by 50%                |                                                 |
| Midazolam            | X   | X    | ↑Midazolam   | Decrease midazolam dose by 50%                 |                                                 |
| Ergot derivatives    | X   | X    | ↑Ergot derivatives | Suspend for 10 days                          |                                                 |
| Omeprazole           |     |      |          | ↓Efficacy ATV                                 | Switch to H2 antagonists. In case of gastrointestinal bleeding, maintain with the administration of the omeprazole dose with a difference of 12 hours from the ATV dose |
| Salmeterol           |     |      |          | ↑Salmeterol                                   | Switch to formoterol or salbutamol              |
| Sildenafil           |     |      |          | ↑Sildenafil                                    | Reduce the dose of sildenafil to 25mg every 2 days |
| Tadalafil            |     |      | ↑Tadalafil   | Reduce the tadalafil dose to 10mg every 72 hours |                                                   |
| Antacids             |     |      |          | ↑Efficacy ATV                                 | Administer 2 hours before or 1 hour after ATV    |
| Tricyclic antidepressants |     |      |          | ↑Antidepressants                              | Decrease antidepressant doses by 50%            |
| Trazodone            |     |      | ↑Trazodone   | Decrease the trazodone dose by 50%             |                                                 |
| Colchicine           | X   | X    | ↑Colchicine  | Use a 0.6mg (1 capsule) dose followed by 0.3mg (1/2 tablet) 1 hour later. Do not readminister in less than 3 days |                                                   |
| Diltiazem            |     |      |          | ↑Diltiazem and DCV                            | Reduce the diltiazem dose by 50%                |
| Nifedipine           | X   | X    | ↑Nifedipine   | Reduce the nifedipine dose by 50%              |                                                  |
| Verapamil            |     |      | ↑Verapamil and DCV                             | Reduce the verapamil dose by 50%                |
| Famotidine           | X   | X    | ↑Famotidine  | Administer ATV 10 hours after and at least 2 hours before the famotidine dose | Administer ATV 10 hours after and at least 2 hours before ranitidine dose |
| Ranitidine           |     |      |          | Administer ATV 10 hours after and at least 2 hours before ranitidine dose | Administer ATV 10 hours after and at least 2 hours before the cimetidine dose |
| Cimetidine           |     |      |          |                                                                 |                                                 |
| Cyclosporine         | X   |      | ↑Cyclosporine | Monitor serum level                           |                                                 |
| Tacrolimus           |     |      | ↑Tacrolimus  |                                                                 |                                                 |
| Sirolimus            |     |      | ↑Sirolimus  |                                                                 |                                                 |
| Clarithromycin       |     |      | ↑DCV      | Reduce the dose of clarithromycin by 50% (QTc prolongation) or use azithromycin |                                                 |
| Oral contraceptives* | X   |      | ↑Ethinyl estradiol |                                                                 | Exchange for desogestrel                          |
| Atazanavir           |     |      | ↑DCV only if using ritonavir                   | Do not change the dose. Ritonavir will not be used |
| Fluticasone          | X   |      | ↑Fluticasone |                                                                 | Switch to budesonide                             |

DCV - daclatasvir; SOF - sofosbuvir; ATV - atazanavir; IV - intravenous; VO - oral; RMP - rifampicine; INR - International Normalized Ratio. *Contraceptives: for patients in the sofosbuvir and daclatasvir groups, contraceptives with ethinyl estradiol will be provided. Patients participating in the atazanavir group will be provided contraceptives with desogestrel or gestodene.
Outcomes

Primary, secondary and safety outcomes are described and defined in table 1.

Sample size

We intend to include 252 patients in Phase II/Stage 1 (189 in the active groups and 63 in the PbO group) and 189 patients in Phase II/Stage 2 (126 in the active groups and 63 in the PbO group). For Phase III, 564 additional participants will be randomized (376 in the active group and 188 in the PbO group). Thus, at the end of Phase III, 314 patients will have been evaluated in the PbO group and between 439 and 502 patients will have been evaluated in the treatment group if the study persists through the three phases. More details about the sample size calculation are described in Supplementary material 2 (Statistical Analysis Plan).

Recruitment and patient retention

Recruitment will be granted for every patient hospitalized with COVID-19, who will be screened for eligibility criteria, sign the Informed Consent Form and be followed by a local study team who have been properly trained until discharge. Loss to follow-up is not expected in this period. If hospital discharge occurs before 15 days after the randomization date, these patients will be evaluated by telephone call, which will be performed by the center 15 days after randomization and 28 days later. Those who return home before D10 will be assessed daily until completion of the study drug treatment period (10 days) by telephone call or in person on D3, 6 or 10 to collect nasal swabs and to assess treatment compliance.

Sequence allocation

The randomization list will be generated electronically using appropriate software. Randomization will be performed in blocks and stratified by center. In Phase II Stage 1, the blocks will have 12 codes (positions), with each treatment group represented by three different codes and each PbO by a single code. In Phase II, Stage 2, the blocks will have six codes, the treatment groups will be represented by two different codes, and each PbO will be represented by a single code. Block sizes will be adapted if any arm is discontinued. In the third stage of the study, the blocks will have six codes, four codes representing the treatment group and 2 codes representing the PbO group.

Allocation concealment

The concealment of the randomization list will be maintained through a centralized, automated, internet-based randomization system, available 24 hours a day, developed by a team of programmers and researchers from the Research Institute of HCor-Hospital do Coração (HCor).

Blinding

This study is not a global double-blind study in Stages 1 and 2, as we have 3 drugs with different physical characteristics, rendering global blinding impossible. Both the participant and investigator can know, after randomization, to which drug group the patient was allocated. However, none will know whether the capsule or coated tablet to be administered contained the active drug or PbO, ensuring blinding of participants and investigators within that specific group, as well as the outcome assessors.

In Phase 3, the global blinding of the stage is possible, since we will have only one active group and a PbO.

A partnership will be established with a handling pharmacy duly licensed to carry out the fractioning, repacking, labeling and blinding process of the drugs. The pharmacy will deliver all the drugs to a logistic contractor, who will deliver them to the centers.

Data collection, management and analysis

Collection and management methods

The data collection system to be adopted in the development of this study is widely used in research projects and easily accessible via the web with Redcap® software. Only users qualified by the system administrator receive access, each registered user accesses the system only with their login and password, and the sharing of this information between project collaborators is prohibited.

The data collected and the participant timeline are described in table 5.

Retention

Data will be collected from patients admitted to the ICU/hospital, which reduces the risk of data loss due to loss of follow-up. Telephone contact will be made centrally by professionals who have not participated in other stages of the study.

- Postdischarge follow-up: Patients discharged less than 15 days after randomization will be contacted by telephone daily until D10 and on D15 and 28 and questioned using a structured form by an interviewer who is blinded to the research participant’s allocation group. In this contact, the participant or his or her family member will be asked about the observation of adverse events, and ordinal scales of 6 and 7 points will be applied for the secondary outcomes.
Statistical methods

The main analysis of the data from this study will be performed on patients of the intention-to-treat (ITT) population. The definition of the analysis populations and more details on statistical methods are available in the Statistical Analysis Plan. The analyses will be performed with R software (R Core Team, 2020).

All the different PbOs for ATV, DCV and SOF will always be analyzed together as a single PbO group.

Analysis of primary outcomes

Phase II/1

In Phase II/1, patients will be allocated in ratio of 3:3:3:1:1:1 (3 for each treatment group and 1 for PbO). For Phase II/1, the parameter of interest for the decision will be the comparison of the decay rates of the viral load logarithm from real time polymerase chain reaction (RT-PCR) to COVID-19 in 9 days between the treatment groups compared with the control using a mixed linear model.

The interim analysis, which is planned to begin when 126 patients are randomized and followed for at least 10 days, will be exclusively conducted for a patient safety assessment (based on safety outcomes and adverse events). At the end of Phase II/1, each group will be compared with the PbO group considering a significance level for each comparison of 0.067 (Bonferroni correction for multiple comparisons), such that the global type I error of Phase II/1 will be 0.20. If no treatment is significantly different from the PbO group at the stipulated level of significance, the study will be terminated at this stage.

Among the treatments significantly different from the PbO group in relation to the linear rate of viral load decay, the one with the highest rate of decay and considered safe by the independent study safety committee will be a candidate treatment for inclusion in Phase II/2 of the study. It will be combined with the drug responsible for the second highest linear decay rate to comprise the second Phase II/2 treatment group.

Phase II/2

In the 2nd stage, the patients will be allocated in the ratio of 2:2:1:1:2 for each active arm and 1 for each PbO, and the same statistic will be used as in Phase II/1. Details are described in the Statistical Analysis Plan - SAP (Supplementary material 2).

As in Phase II/1, the interim analysis planned after inclusion and data are collected from half of the planned patients will only assess safety data.
Phase III

The third stage of the study will proceed in a 2:1 allocation (two active treatments for each PbO), with a minimum inclusion of 189 patients and a maximum of 564. The study will perform interim analyses that may interrupt the study due to safety, futility, or efficacy, using all participants already randomized in Phases II/1 and II/2, since the primary efficacy outcome of Phase III (time of use of ventilatory support within 15 days) will be analyzed for all patients from these earlier phases as well.

The hypothesis test for the treatment effect will be performed using a generalized additive model of location and scale considering the distribution of a beta-binomial mixture with inflated zeros for the data adjusted by age and considering the random effect of the center for each intercept (model for the beta-binomial part and for the probability of zeros).

Phase III has three interim analyses planned, m = 3, for each third of the collected sample. The interim analyses consider stop limits according to the O’Brien-Fleming criterion, with significance levels of 0.06%, 1.51% and 4.71%, respectively, to maintain the global significance level at 5%. In Phase III, discontinuations due to futility will be allowed and use the same stopping criteria defined for efficacy.

Although the PbO group is composed of individuals receiving PbOs for three separate drugs (sometimes in combination), it will always be evaluated as a single group after gathering information from all “PbO arms”.

Secondary/exploratory outcomes and additional analyses are described in Supplementary material 1 (Section S4.1).

Interim analyses

One interim analysis is planned in Phase II (first stage), 1 in Phase II (second stage) and 3 in Phase III, and 3 interim analyses will be conducted, as described in detail in the statistical methods section.

The database will be blocked after obtaining a 15-day follow-up for all patients, and all necessary actions to obtain follow-up data will be performed. All interim analyses would be made available to Agência Nacional de Vigilância Sanitária (Anvisa).

Ethical aspects and good clinical practices

Ethical considerations and dissemination

The trial was designed according to the guidelines for good clinical practice, follows the principles of the Declaration of Helsinki, and was approved by the Comissão Nacional de Ética em Pesquisa (Conep; n° 4.799.171, June 23, 2021), Anvisa (n° 107/2020) and the Ethics Committee (EC) guidelines at each center.

Patients are included after the signed Informed Consent Form is obtained by the investigators participating in the study. In addition, all eventual amendments to the protocol must be approved by the REC/Conep system before its implementation by the participating centers.

Summary of versions

| Version | Date         |
|---------|--------------|
| v4.0    | Feb 26th / 2021 |
| v3.0    | Dec 28th / 2020 |
| v2.0    | Oct 30th / 2020 |
| v2.0    | Oct 15th / 2020 |
| v1.0    | Sep 09th / 2020 |
| v1.0 August 2020 | Aug 07th / 2020 |
| Original project v1.0 July 2020 | Jul 27th / 2020 |

The study will be submitted for publication after completion, regardless of its findings. Manuscript preparation will be an inalienable responsibility of the Steering Committee. The main paper will be authored by the Steering Committee members plus the principal investigators of the ten top recruiting sites, who will contribute intellectually to the manuscript.

Adverse events

All adverse events will be recorded by the investigators on the clinical records (CRF), regardless of severity, and graded according to the Division of AIDS (DAIDS) Severity table. Table 1S (Supplementary material 1) shows the known adverse events resulting from the use of the products under investigation and explains the main actions to implement for SAEs.

The principal investigator at the research center is responsible for informing research ethics committees of any SAEs within 24 hours, as required by local regulations, except those that are classified as study outcomes.

The HCor Research Institute is responsible for receiving and monitoring all adverse events that occur during the clinical study. All adverse events classified as serious according to the definition of RDC Anvisa 09/15 must be informed to the sponsors within a maximum period of 24 hours of knowledge through a specific email. After evaluation, Fiocruz and Blanver Farmoquímica e Farmacêutica must notify Anvisa of serious, unexpected reports whose causality is possible, probable or defined in relation to the product under investigation within the regulatory deadlines.

Study organization

Protocol violations and deviations

Major deviations related to the inclusion or exclusion criteria, study conduction, and patient management must be reported to the coordinating site.
Trial organization and oversight

The Steering Committee comprises the study investigators and is responsible for the development of the study protocol, manuscript drafts and study submission for publication. A team from the HCor Research Institute coordinates the study in association with the Brazilian Research in Intensive Care Network (BRICNet) and Coalition Brazil. The HCor Research Institute has been responsible for conducting the study since its inception and managing and controlling data quality. The Data Safety and Monitoring Board (DSMB) is composed of an external statistician and 2 other researchers who are experts in critical care medicine. The DSMB is responsible for the interim analysis and for providing guidance to the Steering Committee regarding the continuation and safety of the trial after the interim analyses.

Current status

We randomized 256 patients from April until July 2021. The first DSMB meeting occurred in May and June 2021. After analyzing data according to DSMB Charter, the DSMB recommended continuing the trial as planned.

Dissemination policy

The results of the study will be disseminated in presentations at congresses and submitted for publication in high impact scientific journals.

DISCUSSION

This manuscript describes the protocol of the REVOLUTION trial, which is a randomized, PbO-controlled, adaptive, multiarm, multistage study to evaluate the efficacy of ATV, DCV and SOF simultaneously compared with their PbOs. The main objective is to identify whether any of these drugs alone or in combination are capable of reducing viral load and, if so, investigate the clinical outcome of increasing the number of days free of respiratory support in a larger population.

Expeditious and methodologically rigid clinical trials are demanding to identify safe and effective treatments. Nevertheless, rapid identification and abandonment of harmful or ineffective approaches are equally important, especially during a pandemic. The principal aim of the REVOLUTION trial is to aid in the recognition of any of these 3 repurposed antivirals as treatments for COVID-19.

Repurposed drugs are important in resource-limited settings because the interventions are more rapidly available, have already been tested safely in other populations and are inexpensive.

Limitations of the protocol are described below. First, decisions related to changing stages rely on a decreasing viral load in the initial phases despite extensive debate on the association of viral load with worse clinical outcomes. Second, late initiation in recruiting participants in the pandemic timeline slowed our recruitment rate, as pandemic cases steadily decreased after vaccination outset and, third, late protocol publication, although the content of this protocol paper is absolutely consistent with the protocol submitted and approved by the national/local Ethics Research Committee, our national regulatory agency (Anvisa) and ClinicalTrials.gov.

CONCLUSION

In conclusion, the REVOLUTION study protocol explains the design and purpose of the study, which has the objective of answering whether repurposed antiviral drugs are effective and safe treatments for COVID-19 in an adaptive, multiarm, multistage design starting with a Phase II clinical trial and ending with a Phase III trial.

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Roles of protocol contributors:

I.S. Maia is the principal investigator; T. M. L. Souza and A. B. Cavalcanti conceived the study, led the proposal and protocol development. A. Marcadenti, F. G. Zampieri, L. P. Damiani and A. B. Cavalcanti contributed to study design and to development of the proposal. I. S. Maia and A. B. Cavalcanti were the trial methodologists. All authors read and approved the final manuscript.

Committees, Leadership, and Investigators Coalition COVID-19 Brazil:

Coalition COVID-19 Brazil is a research alliance coordinated by the following institutions: Hospital Israelita Albert Einstein, HCor-Hospital do Coração, Hospital Sírio-Libanês, Hospital Moinhos de Vento, Hospital Alemão
Antivirals for adult patients hospitalized with SARS-CoV-2 infection: A randomized, Phase II/III, multicenter, placebo-controlled, adaptive study, with multiple arms and stages.

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