Opaganib in Coronavirus Disease 2019 Pneumonia: Results of a Randomized, Placebo-Controlled Phase 2a Trial

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Background. Opaganib, an oral sphingosine kinase-2 inhibitor with antiviral and anti-inflammatory properties, was shown to inhibit severe acute respiratory syndrome coronavirus 2 replication in vitro. We thus considered that opaganib could be beneficial for moderate to severe coronavirus disease 2019 (COVID-19) pneumonia. The objective of the study was to evaluate the safety of opaganib and its effect on supplemental oxygen requirements and time to hospital discharge in COVID-19 pneumonia hospitalized patients requiring supplemental oxygen.

Methods. This Phase 2a, randomized, double-blind, placebo-controlled study was conducted between July and December 2020 in 8 sites in the United States. Forty-two enrolled patients received opaganib (n = 23) or placebo (n = 19) added to standard of care for up to 14 days and were followed up for 28 days after their last dose of opaganib/placebo.

Results. There were no safety concerns arising in this study. The incidence of ≥ Grade 3 treatment-emergent adverse events was 17.4% and 33.3% in the opaganib and placebo groups, respectively. Three deaths occurred in each group. A numerical advantage for opaganib over placebo was observed in this nonpowered study reflected by total supplemental oxygen requirement from baseline to Day 14, the requirement for supplemental oxygen for at least 24 hours by Day 14, and hospital discharge.

Conclusions. In this proof-of-concept study, hypoxic, hospitalized patients receiving oral opaganib had a similar safety profile to placebo-treated patients, with preliminary evidence of benefit for opaganib as measured by supplementary oxygen requirement and earlier hospital discharge. These findings support further evaluation of opaganib in this population.

Keywords. hospitalization; SARS-CoV-2; sphingosine-kinase-2; supplemental oxygen.

Acute pneumonia due to severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) infection leading to coronavirus disease 2019 (COVID-19) can be life-threatening. Typical symptoms of SARS-CoV-2 infection include fever, fatigue, dry cough, and shortness of breath, but it can evolve to acute respiratory distress syndrome. Among patients with COVID-19, 80% present with symptoms of mild to moderate severity, with or without pneumonia, whereas 5%–14% develop serious or critical respiratory disease, requiring intensive care unit support [1–3]. The mortality rate of COVID-19 ranges from 0.1% to 19.7%, depending on country of origin, socioeconomic background, comorbidities such as diabetes and cardiovascular disease, and disease severity [4–8]. These comorbidities and advanced age (especially ≥65 years) increase the risk of mortality [9, 10].

Patients who are hospitalized with COVID-19 often require supplemental oxygen to maintain adequate oxygenation [3, 9]. Some pharmacological agents (corticosteroids and remdesivir) have been widely used as standard of care (SoC), with remdesivir first approved in the United States for emergency use in hospitalized COVID-19 patients in May 2020 [11].

Opaganib (RedHill Biopharma) is a first-in-class orally available, small molecule, selective sphingosine kinase-2 (SK-2) inhibitor with antiviral and anti-inflammatory properties [12]. Multiple ongoing clinical trials are evaluating opaganib as therapy for cancer. The antiviral effect of opaganib was demonstrated in various preclinical studies. Reid et al [13] showed that decreased expression or pharmacologic inhibition of SK-2 significantly inhibited Chikungunya virus replication. In a work by Xia et al [14], opaganib was shown to reduce fatality rates from influenza A infection in mice. Moreover, in a preliminary study using an in vitro model of human bronchial tissue, we recently demonstrated that opaganib has potent antiviral activity against SARS-CoV-2 [15]. We hypothesized that the antiviral and anti-inflammatory effects of opaganib would be beneficial for the treatment of SARS-CoV-2 infection, particularly for patients with moderate to severe COVID-19 pneumonia.
requiring supplemental oxygen. In a small cohort of patients with severe COVID-19 pneumonia, in which opaganib was offered through compassionate use, it appeared to improve clinical and laboratory parameters, and it was safe and well tolerated [16].

This Phase 2a proof-of-concept study was designed to provide a preliminary evaluation of the safety and efficacy of opaganib when added to SoC in hospitalized patients with COVID-19 pneumonia who required supplemental oxygen. Aside from safety and tolerability assessments, efficacy endpoints measured included change in oxygen requirements and clinical improvement parameters.

METHODS

Design
This was a Phase 2a, proof-of-concept, multicenter, randomized, double-blind, parallel arm, placebo-controlled study conducted in 8 sites in the United States. The study was not powered for efficacy. Eligible patients received opaganib or matching placebo twice daily added to SoC (per regional, institutional/physician practices). Eligible participants were adults (18–80 years inclusively), with SARS-CoV-2 infection determined by polymerase chain reaction analysis in a nasopharyngeal sample and pneumonia determined by chest x-ray. The full list of eligibility criteria is provided in the Supplementary Data, Section-1.

Intervention
Participants were randomized 1:1 to receive 2 × 250 mg of opaganib oral capsules or matching placebo every 12 hours for up to 14 days (Days 1–14). This dose had been determined in a Phase 1 clinical study in oncology patients as the maximum safe and tolerable dose [17]. Patients discharged before or at Day 10 completed only 10 days of treatment. After their last study drug dose, participants were followed for up to 28 days (safety follow-up). Treatment assignments remained blinded to the patients, investigators, hospital staff, and sponsor (RedHill Biopharma). The clinical trial (ClinicalTrials.gov Identifier NCT 04414618) was performed in accordance with the Declaration of Helsinki (1964), as revised most recently in Seoul (2008), US Food and Drug Administration (FDA) regulations, and the ICH Guideline for Good Clinical Practice, E6(R1) and local rules and regulations.

Outcomes
Given the exploratory nature of this proof-of-concept study, we sought to evaluate several clinically meaningful outcomes. We designed a primary outcome measure comparing total supplemental oxygen requirements from baseline to Day 14 (area under the curve) using daily oxygen flow measurements. Primary endpoint details are provided in the Supplementary Data, Section-2, together with a complete list of secondary, post hoc, safety, and exploratory endpoints. Presented herein are the safety outcomes, that is, incidence of treatment-emergent adverse events (TEAEs) and serious TEAEs (TESAEs) over the safety follow-up period, as well as the following clinically relevant secondary outcomes: (1) time to 50% reduction from baseline in supplemental oxygen requirement by Day 14, (2) percentage of patients no longer receiving supplemental oxygen for at least 24 hours by Day 14, (3) time to intubation and mechanical ventilation by Day 14, and (4) mortality by Day 30. The post hoc outcomes were as follows: (1) percentage of patients no longer receiving supplemental oxygen for at least 24 hours by Day 14 per SoC regimen, (2) time to intubation and mechanical ventilation by safety follow-up, (3) mortality by safety follow-up, (4) incidence of hospital discharge by Day 14 and by safety follow-up, and (5) time to ≥ 2 points improvement in the World Health Organization (WHO) Ordinal Scale [18] by Day 14.

Statistics
Because the sample size (~40 patients) was not chosen for statistical consideration, efficacy interpretation is based on numerical comparisons of descriptive statistics. Statistical methodology details are provided in the Supplementary Data, Section-3. Baseline characteristics are presented for the intent-to-treat (ITT) population (all randomized patients). Safety evaluations were performed for the safety population (patients who took at least 1 dose of study drug). Efficacy endpoints were analyzed for the modified ITT (mITT) population (patients who took at least 1 dose of study drug).

Patient Consent Statement
Written consent was obtained from all patients who participated in this study. The design of the study has been approved by local ethical committees and by the national regulatory agencies in all the countries where the protocol was activated including the FDA, the Israeli Ministry of Health, and the Argentine Administration of Medicines, Food and Medical Technology.

RESULTS
A diagram of the study flow is presented in Figure 1. The ITT population comprised 42 eligible patients, enrolled in July–November 2020. Of these, 23 patients were randomized to receive opaganib and 19 patients were randomized to receive placebo (randomization details are provided in the Supplementary Data, Section-4. One patient in each group did not receive study drug and was excluded from the safety and mITT populations. One patient randomized to placebo was mistakenly given opaganib on Days 10 and 11.

Demographic and Other Baseline Characteristics
The main demographic and baseline characteristics of patients in the ITT population are presented in Table 1. Most patients...
were male (64.3%) with an overall median age of 58.0 years. Randomization resulted in a difference between the opaganib and placebo arms in median age (years; 52.0 and 61.0, respectively) and median baseline oxygen requirement (L/minute; 6.0 and 10.5, respectively). The median time (days) from onset of symptoms to randomization was 10.0 for the opaganib group and 9.0 for placebo.

Remdesivir and/or high-dose corticosteroids (dexamethasone, prednisone, and methylprednisolone) comprised known effective SoC for COVID-19 and were coadministered with opaganib in 22 (95.7%) patients, and with placebo in all 18 (100%) patients. This included remdesivir in 10 (43.5%) patients on opaganib and 9 (50.0%) on placebo, and high-dose corticosteroids in 21 (91.3%) and all 18 (100%) patients, respectively.

**Treatment Compliance**

Compliance was assessed in a subset of the safety population comprising 21 patients on opaganib and 18 patients on placebo (further details are provided in the Supplementary Data, Section-5). For 2 patients on opaganib, compliance was not defined: for the patient randomized to placebo who took opaganib on one day, the expected number of opaganib doses was zero; the second patient was lost to follow-up, and thus compliance could not be assessed. The mean (± standard deviation) compliance was 90.05% (±21.523) in the opaganib and 98.41% (±3.917) in the placebo groups, respectively.

**Treatment-Emergent Adverse Events**

The incidence of TEAEs was similar between groups (52.2%, in the opaganib group versus 50.0% in the placebo group) (Table 2). In general, each unique TEAE term was reported singly by 1 patient (either in the opaganib or placebo group). The only TEAEs reported by 2 patients (8.7% in the opaganib group), and with a higher incidence in the opaganib group than in placebo, were hypokalemia (reported in 1 patient on placebo) and diarrhea (not reported in patients on placebo).
The TESAEs were reported in 13.0% of patients on opaganib and 27.8% patients on placebo. The TESAEs by Medical Dictionary for Regulatory Activities System Organ Class included the following: infections and infestations (8.7% in the opaganib and 11.1% in the placebo group); respiratory, thoracic, and mediastinal disorders (8.7% and 16.7%, respectively); and renal and urinary disorders (4.3% and 11.1%, respectively). None of the TESAEs were deemed as treatment related by the investigator.

The TEAEs of Grade 3 and above were reported in 4 patients (17.4%) on opaganib and 6 patients (33.3%) on placebo, with no TEAEs of Grade 3 and above reported at a higher frequency in the opaganib group compared with placebo (Table 2). Treatment-related TEAEs were recorded in 2 patients in the opaganib group, including 1 patient with Grade 1 diarrhea and 1 patient with Grade 3 rash. Both events were nonserious, and in both cases, the drug was withdrawn, and the patients recovered.

There were no reported TEAEs resulting in dose reductions. The TEAEs leading to treatment withdrawal were reported in 4 patients (17.4%) in the opaganib group (including a case of Grade 1 heart palpitations [not related to study treatment], a
be regarded as such.

In addition to the safety analysis, several efficacy analyses (presented below) were performed with the aim of providing initial data on potential beneficial effects of opaganib in the study population. Because the study was not powered for efficacy, the findings from these analyses provide a preliminary evaluation of opaganib's potential benefits, and they should be regarded as such.

### Total Supplemental Oxygen Requirement Using the Maximal Daily Oxygen Flow Over 14 Days

A relative benefit derived for each group was calculated (as described in the Supplementary Data, Section-2). The relative benefit calculated was 61.6% for opaganib (n = 21) and 46.7% for placebo (n = 18) (Supplementary Figure 1). This was a post hoc analysis to the predefined primary endpoint; as more information on the course of COVID-19 pneumonia became available, this endpoint, namely, quantifying total supplemental oxygen required over time (across various devices), proved to be of less clinical value than initially anticipated.

### Requirement of Supplemental Oxygen by Day 14

The cumulative incidence for no longer requiring supplemental oxygen for at least 24 hours by Day 14 was estimated using the Kaplan-Meier analysis. The analysis showed a higher estimated cumulative incidence in patients on opaganib (50%) compared with those on placebo (22.2%) (Supplementary Figure 2, Table 1).

### Intubation and Mechanical Ventilation, and Mortality During the Course of the Study and at the End of Follow-up

The Kaplan-Meier estimated time to 50% reduction from baseline in supplemental oxygen requirement based on oxygen flow (L/minute) Day 1–Day 14 was 5 days in the opaganib versus 8 days in the placebo group. The Kaplan-Meier estimated cumulative incidence of this event by Day 14 was 81% in the opaganib group compared to 66.7% in the placebo group.

### Hospital Discharge

The Kaplan-Meier estimated cumulative incidence of hospital discharge was 68.2% in the opaganib and 50.0% in the placebo group.
groups by Day 7 and 86.4% and 55.6%, respectively, by Day 14 (Supplementary Figure 3). A related post hoc analysis, whereby the 2-point improvement in the WHO Ordinal Scale was derived for patients, is presented in Supplementary Table 1.

DISCUSSION

This proof-of-concept study explored the safety and effectiveness of oral opaganib in hospitalized patients with COVID-19 pneumonia requiring supplemental oxygen. Opaganib’s safety profile in this study was not materially different than that of placebo. Results suggest that clinical improvement, as measured by reduced need for supplemental oxygen, improved WHO level in the scale for clinical improvement and in time to discharge (Supplementary Table 2).

Several lines of preliminary evidence suggested that opaganib may be particularly relevant to COVID-19 therapy. Before the onset of the COVID-19 pandemic, opaganib had demonstrated antiviral and anti-inflammatory effects in several preclinical models, including the following: amelioration of Pseudomonas aeruginosa-induced lung injury and reduction of fatality rates from influenza infection [19, 20]; inhibition of host inflammatory responses in several disease models [21, 22]; and exertion of various effects on the immune system [23]. Opaganib has since been shown to have potent antiviral activity against SARS-CoV-2 in a preclinical study using an in vitro model of human bronchial tissue [15] and was associated with favorable outcomes when offered through compassionate use to patients with moderate to severe COVID-19 pneumonia requiring supplemental oxygen via high-flow nasal cannula [16]. Because opaganib targets a host cell factor rather than the virus directly, development of resistance is less likely. This is of particular interest in SARS-CoV-2 infection, given its rapidly emerging mutations and viral strains.

The antiviral agent remdesivir has been approved for treatment in certain stages of COVID-19; however, it is administered intravenously and lacks anti-inflammatory properties [5], whereas opaganib is an oral agent with both antiviral and anti-inflammatory properties. Moreover, opaganib is stable at room temperature for 5 years (RedHill Biopharma Ltd., unpublished data, 2022). These factors support opaganib as a much-needed, practical, and convenient potential treatment of COVID-19 pneumonia.

Opaganib was shown to be well tolerated in this study, with no new safety signals emerging. The only TEAE of Grade 3 and above reported at a higher frequency in the opaganib group than placebo was a Grade 3 rash in a single patient, which resolved upon stopping study drug. Rash is a known uncommon toxicity of opaganib. Three patients on opaganib and 5 patients on placebo experienced TESAEs. The TEAEs leading to treatment withdrawal were reported in 17.4% of patients on opaganib and 5.6% of patients on placebo. By the safety follow-up timepoint, there were 3 deaths recorded in each group, and none were deemed treatment related. The overall incidence of TEAEs and TESAEs were similar in both treatment groups.

The current study was not powered for efficacy, and, as such, its efficacy findings provide a preliminary evaluation of opaganib’s beneficial effects in the study population. A numerical superiority was found for opaganib over placebo across several clinical outcomes. The prespecified primary efficacy analysis of change in total supplemental oxygen requirements proved difficult to analyze and required a post hoc analysis utilizing percentage change. This post hoc analysis demonstrated an apparent difference, with less total supplemental oxygen requirement for patients on opaganib compared with those on placebo. As the pandemic progressed, improvement in supplemental oxygen requirement had become a commonly used criterion for hospital discharge in COVID-19 patients [24]. Hence, the predefined secondary analysis of no longer requiring supplemental oxygen for at least 24 hours was the most clinically relevant endpoint. A number of other prespecified and post hoc analyses measuring clinical benefit supported the improvement in oxygen requirements. Of note, however, there was an imbalance in median age between the 2 arms of the study, with a younger population in the opaganib group (52.0 years) compared with placebo (61.0 years). In addition, the mean baseline oxygen requirement (L/minute) was lower in the opaganib group (6.0) than in placebo (10.5). Because younger individuals are expected to have generally better disease outcomes than their older counterparts, we cannot rule out that these imbalances may have impacted the preliminary efficacy results of the study, contributing to opaganib’s observed benefit in the studied population. However, such potential impact remains hypothetical, because most studies looking at the effect of age on disease severity report the age of >65 years as a risk factor.

Limitations

The main limitation of this study is its small sample size, which allows for descriptive analyses rather than meaningful statistical interpretation. A further limitation is the imbalance in age and supplemental oxygen requirement at baseline, which is a result of the small sample size despite randomization. The clinical interpretation of the primary endpoint was limited, as described under Discussion, leading to the addition of post hoc analyses to help with the overall interpretation of the data. An additional limitation of the study is that certain predefined secondary endpoints that relied on inpatient data collection could not be assessed due to changing standard of practice leading to earlier discharge of patients than anticipated at the time of protocol development. Furthermore, discharge criteria were somewhat subjective, depending on hospital capacity and regulations. It is notable that a greater proportion of patients on opaganib versus placebo were discharged before Day 14.
CONCLUSIONS

Taken together, the results of this Phase 2a study suggest that opaganib is safe and well tolerated and suggest a potential benefit of opaganib across various clinically meaningful outcome measures in patients with COVID-19 pneumonia. Based on these findings, further evaluation of opaganib is currently ongoing in larger randomized studies as a potential treatment for moderate to severe COVID-19 pneumonia in hospitalized patients requiring supplemental oxygen (a global Phase 2/3 [ClinicalTrials.gov Identifier NCT04467840] already completed enrollment [15]).

Supplementary Data

Supplementary materials are available at Open Forum Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

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Potential conflicts of interest. A. B. and G. R. are employees of RedHill Biopharma Ltd. P. A., R. F., V. K. B.-Y., and M. L. L. are full-time consultants for RedHill Biopharma Ltd. G. E. and K. L. W. are consultants to RedHill Biopharma Ltd. A. W. S.’s spouse purchased RedHill Biopharma Ltd. stocks. A. M. R. and H. S. M. have received research support from other companies. M. S. G. has worked as an oncology consultant to RedHill Biopharma Ltd. and has participated in research studies conducted by other companies.

All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

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