Efficacy of pragmatic same-day ring prophylaxis for adult individuals exposed to SARS-CoV-2 in Switzerland (COPEP): protocol of an open-label cluster randomised trial

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ABSTRACT

Introduction Lopinavir/ritonavir (LPV/r) has been proposed as repurposed drugs for pre-exposure and postexposure prophylaxis as well as therapy of COVID-19. Coronavirus postexposure prophylaxis (COPEP) trial aims at assessing their efficacy as postexposure ring-prophylaxis among adults exposed to SARS-CoV-2.

Methods and analysis COPEP is a two-arm open-label cluster-randomised trial conducted in three cantons of Switzerland. Asymptomatic contacts (≥16 years) of individuals diagnosed with COVID-19 will be randomised (2:1) to either LPV/r (400 mg/100 mg two times per day) for 5 days, or a standard of care arm (no treatment). Asymptomatic individuals may be either SARS-CoV-2 positive or negative. Contacts living in the single household will form a cluster and will be randomised into the same arm. All participants will be followed-up for 21 days and undergo daily monitoring for COVID-19 symptoms. The primary endpoint is 21-day incidence of laboratory-confirmed COVID-19 with ≥1 compatible symptoms, analysed in an intention-to-treat (ITT) analysis. The secondary endpoints include the 21-day incidence of COVID-19 as well as SARS-CoV-2 infection in a modified ITT analysis, excluding participants who had a positive SARS-CoV-2 RT-PCR from oropharyngeal swab and/or a positive SARS-CoV-2 IgG serology at baseline. Assuming a 21-day incidence for COVID-19 of 20% among contacts without postexposure chemoprophylaxis, to detect a relative risk reduction of 60% (ie, translating in an absolute reduction from 20% to 8%), with a power of 80%, an alpha of 5%. Accounting for design effect of cluster design of circa 1.1, we plan to enrol 200 participants to the LPV/r arm and 100 to the standard of care arm, 300 participants in total.

Strengths and limitations of this study

Among the first clinical trials to study two prophylactic candidates for COVID-19 in the general population as well as healthcare workers.

The trial will test pragmatic ring prophylactic strategies that can be prescribed the same day with minimal clinical and laboratory assessment.

Recruitment may be slowed down by the evolution of the epidemic.

This is an unblinded trial.

INTRODUCTION

Since the beginning of the SARS-CoV-2 outbreak in December 2019, the medical and scientific community has scrambled to identify effective pharmacological candidates for its prevention and treatment. COVID-19 (which is the clinical manifestation of SARS-CoV-2 infection) has caused unprecedented pressure on modern health systems across the world. The sudden and uncontrollable influx of severe COVID-19 cases has caused critical shortage of hospital-based resources and forced many countries into lockdown in early 2020.
Containing the COVID-19 pandemic may necessitate a multipronged strategy, including immunisation, treatment and prophylaxis. The latter would facilitate the containment of future outbreaks of the disease in non-immunised populations. This current pandemic wave will likely be followed by subsequent outbreaks and authorities will need to consider less strict isolation measures when appraising multiple economic impacts (including on health-related outcome) over a prolonged period of time.

Several drugs have been identified as possible prophylactic candidates. These include the protease inhibitors lopinavir/ritonavir (LPV/r) used for the treatment of HIV. Protease inhibitors work by preventing viral replication and LPV/r has been shown to bind to active site of the SARS-CoV protease in vitro, with studies confirming that the spatial structure of the binding site was conserved between SARS-CoV and SARS-CoV-2. LPV/r is licensed in most countries, are part of the WHO essential list, and are available globally and at low cost.

However, to date, there are no published randomised studies assessing the efficacy of LPV/r as prophylaxis for COVID-19. Coronavirus postexposure prophylaxis (COPEP) is a Swiss two-arm open-label cluster randomised trial that will assess the efficacy, safety and acceptability of same-day LPV/r-based prophylaxis, compared with standard of care alone (no treatment) for asymptomatic individuals exposed to SARS-CoV-2.

METHODS

COPEP is an open-label two-arm (2:1) cluster randomised superiority trial to assess efficacy, safety and acceptability of same-day LPV/r prophylaxis vs standard of care (no treatment) for asymptomatic adult individuals exposed to SARS-CoV-2. Asymptomatic individuals may be either SARS-CoV-2 positive or negative. The study will be performed at three sites in Switzerland, Geneva, Basel and Lugano, and seeks to recruit 300 participants over 8-9 months. The main objective of COPEP is to assess prevention of laboratory-confirmed COVID-19 with ≥1 compatible symptom, with secondary objectives, including preventing of SARS-CoV-2 infection and attenuation of COVID-19 severity.

Participants

This trial enrols individuals (≥16 years) who have come into close contact with a confirmed case of SARS-CoV-2 infection. Individuals are eligible to be enrolled if the contact occurred within the last 7 days but no more than 72 hours after the index case was diagnosed. A close contact is defined, as per the Swiss Federal Office of Public Health, as a person who spent >15 min in <2 m distance or shared closed space with a confirmed case for a prolonged period (e.g., more than 2 hours) in the period extending from 48 hours before onset of symptoms (or before the date of testing in the absence of symptoms), or a person who had direct contact with the body fluids or laboratory specimens of a case without recommended personal protective equipment (PPE) or in case of failure of PPE.

Recruitment of participants to this study will be performed through two routes; (1) the Public Health Authorities in Geneva, Basel and Lugano, who, as the standard of care, contact all individuals tested positive for SARS-CoV-2 to provide them with instructions on isolating procedures and self-monitoring and (2) healthcare workers will be recruited via social media platforms and internal hospital-based platforms. All close contacts of confirmed SARS-CoV-2 positive index case will be screened for eligibility (table 1).

Intervention

Participants will be randomised in household clusters, 2:1, to receive either LPV/r, 400 mg/100 mg (2× 200 mg/50 mg pills) two times per day for 5 days; or standard of care (no treatment). The first dose of LPV/r will

### Table 1 Inclusion and exclusion criteria

| Inclusion criteria                                      | Exclusion criteria                                                                 |
|--------------------------------------------------------|------------------------------------------------------------------------------------|
| 1. Documented close contact with a PCR-confirmed SARS-CoV-2 positive individual | 1. Fever (temperature >38.0°C) and/or respiratory symptoms (cough, dyspnoea) and/or new anosmia/ageusia |
| 2. ≥16 years of age                                    | 2. Individuals with previous confirmed SARS-CoV-2 infection within the last 6 months |
| 3. Informed consent documented by signature (including parent’s or legal guardian’s signature if the participant is 16 and 18 years old) | 3. Known impairment of liver function |
|                                                        | 4. Known hypertensitivy to any of the study medications                           |
|                                                        | 5. Use of any medications that are contraindicated with LPV/r using the website www.hiv-druginteractions.org/checker |
|                                                        | 6. Individuals on boosted protease inhibitor as part of an antiretroviral therapy |
|                                                        | 7. Inability to be followed-up for the trial period                              |

LPV/r, lopinavir/ritonavir.
be taken during the baseline visit, as directly observed therapy.

Outcomes
Primary endpoint
The primary endpoint is the 21-day incidence of laboratory-confirmed COVID-19 with ≥1 compatible symptom in individuals exposed to SARS-CoV-2 who are asymptomatic (either SARS-CoV-2 positive or negative) at baseline. The primary analysis includes all individuals enrolled, irrespective of their baseline oropharyngeal swab results or the baseline SARS-CoV-2 serology (intention-to-treat (ITT) analysis).

Secondary endpoints
- 21-day incidence of COVID-19 in individuals exposed to SARS-CoV-2 who are asymptomatic, and have negative SARS-CoV-2 PCR and serology at baseline (modified ITT).
- 21-day incidence of SARS-CoV-2 infection in individuals exposed to SARS-CoV-2 who are asymptomatic and have negative SARS-CoV-2 PCR and serology at baseline (modified ITT).
- Severity of clinical COVID-19 on a 7-point ordinal scale.
- Incidence of serious adverse events.

Explorative endpoints
- Acceptability of a prophylaxis for COVID-19.
- Reported adherence to LPV/r for participants on the LPV/r arm and HCQ and LPV/r drug levels on day 5 among all individuals.

Measurements
The following clinical definitions will be used. COVID-19 is defined as ≥1 symptom compatible with COVID-19 (cough, dyspnoea, anosmia, ageusia, elevated temperature (>38.0°C)) and either
1. a positive PCR for SARS-CoV-2 in oropharyngeal swab and/or
2. a seroconversion of IgG only or IgG and IgA for SARS-CoV-2 at day 21 in individuals with negative serology at baseline. In case of seroconversion of IgA only, seroconversion of IgG using more sensitive spike-based recombination immunofluorescence assay (S-rIFA) will be assessed.

New SARS-CoV-2 infection is defined as
1. a positive PCR for SARS-CoV-2 (oropharyngeal swab) among those with a negative PCR at baseline and/or
2. a seroconversion of IgG only or IgG and IgA for SARS-CoV-2 at day 21 in individuals with negative serology at baseline. In case of seroconversion of IgA only, seroconversion of IgG using more sensitive S-rIFA will be used.

Seroconversion for SARS-CoV-2 is defined as negative results for IgG in ELISA at baseline and either
1. positive result for IgG in ELISA and confirmation by S-rIFA at day 21 or
2. doubtful result for IgG in ELISA at day 21 and confirmation by S-rIFA. In case of negative result for IgG in ELISA at day 21 seroconversion can alternatively be defined as follows: Negative result of IgA in ELISA at baseline and (a) positive or doubtful result for IgA in ELISA at day 21 and (b) positive result for IgG in S-rIFA at day 21.

Procedures
All participants will be followed-up for 21 days (figure 1 and table 2). Participants will undergo an oropharyngeal swab for SARS-CoV-2 PCR and a SARS-CoV-2 serology at baseline and on day 21. In addition, participants will be asked to provide on day 5, a capillary puncture on dried blood spot (DBS) to assess blood levels of LPV/r. Participants will be given the option of ‘self-test’, or of a home-visit for this DBS procedure.

During follow-up, individuals will be asked to complete a daily online COVID-19 symptoms questionnaire. The online questionnaire generates alerts when individuals report a symptom associated with COVID-19 and if participants do not complete the questionnaire for 2 consecutive days. The online questionnaire also serves as a reminder for those on the LPV/r arm, to take their daily medication up to day 5. Paper questionnaires will be made available for those without access to internet. All questionnaires are available in several languages.

The team will contact the participant in case an alert is triggered and do a first clinical assessment by phone. Participants who report COVID-related symptoms (eg, dyspnoea, cough, fever (>38.0°C), anosmia) will be asked to come onsite for an extra visit and undergo clinical assessment and an oropharyngeal swab to confirm/exclude SARS-CoV-2 infection. If found positive, participants will be provided with appropriate care, as per local protocol. A follow-up visit 14 days after a positive diagnosis will be conducted, in order to establish the level of clinical manifestation of disease.

On day 21, participants will further complete a questionnaire on adverse events and acceptability of the treatment. Participants on the LPV/r arm are asked to bring the phials which contained the medication to day 21 visit, so that the study team can record the number of returned pills, as per good clinical practice standard.

Sample size
The sample size for the primary endpoint assumes that without treatment 20% of close contacts will develop COVID-19, based on the clinical observations made by the team. To detect a relative risk reduction of 60% (ie, translating in an absolute reduction from 20% to 8%), with a power of 80%, an alpha of 5% and accounting for design effect of cluster design of circa 1.1 (based on DE=(n-1)r+1, where n=3 denotes the average cluster size, and r=0.05 is the assumed intraclass correlation coefficient). A sample size of 200 participants on LPV/r and 100 on standard of care arm, or 300 in total will thus be needed.
For the first of the secondary endpoint (occurrence of new SARS-CoV-2 infection) we assume 40% of close contact without prophylaxis will become infected and 16% with prophylaxis; but we also expect a baseline prevalence of positive PCR of 30% (these participants will be excluded from this analysis in a modified ITT). With an effective sample size of 140 individuals on the LPV/r arm (200×0.7) and 70 on the standard of care arm (100×0.7) the power will be 95% for this endpoint.

Randomisation and blinding
Randomisation will be done in variable-sized blocks (sizes 6 or 9) in random sequence, stratified by site (Geneva, Basel and Lugano). Randomisation will be done by cluster, that is where close contacts reside in the same household (eg, the family or friends of the index case that live together). The randomisation procedure will consist of sealed envelopes, one for each household cluster, prepared by individuals who are not associated with the recruitment for the trial. Treatment will be identical within households because of the risk of cross-contamination and to simplify treatment administration to individuals in the same household. This is an open-label trial.

Statistical analysis
For the analysis of the primary endpoint, 21-day incidence of COVID-19, we will perform an ITT, including all individuals who were randomised (including those who will retrospectively be found to be PCR-confirmed SARS-CoV-2 positive at baseline as well as individuals retrospectively found SARS-CoV-2 seropositive by serology).

The intervention arm, LPV/r, will be compared with the surveillance arm, using separate indicator variables for the active treatment arms. Because the hypotheses about treatment efficacy are unrelated and independent, and because we will focus on estimation and CIs rather than on statistical tests, no adjustment will be done for multiplicity. Since individual observations will be clustered within households (randomisation units), themselves nested within index cases, we will use mixed complementary log-log regression models for the main analysis (complementary log-log regression is similar to logistic regression, but yields relative hazards, rather than ORs; relative hazards are more readily interpretable in the context of disease incidence). The outcome variable will be the occurrence of COVID-19 by day 21.
A random intercept will be defined by each household, nested within the index case. The main fixed effect will be treatment. The main statistical model will be adjusted for potential confounding variables, guided by the most up to date evidence. We foresee that for the incidence of COVID-19, adjustment variables are: age, presence of comorbidities (specifically cardiac, liver or pulmonary disease), treatment of the index case, occupational versus non-occupational exposure and/or positive serology.

In a modified intent to treat analysis, we will evaluate the secondary endpoints; (1) 21-day incidence of COVID-19 in individuals exposed to SARS-CoV-2 who are asymptomatic, PCR-confirmed SARS-CoV-2 negative and have negative SARS-CoV-2 serology at baseline (modified ITT), (2) 21-day incidence of SARS-CoV-2 infection in individuals exposed to SARS-CoV-2 who are asymptomatic, PCR-confirmed SARS-CoV-2 negative and have negative SARS-CoV-2 serology at baseline (modified ITT). The output will be an adjusted relative hazard of the treatment effect versus surveillance, with a 95% CI.

For the secondary outcome ‘severity of disease’, rated on a 7-point scale, we will use a Mann-Whitney test to compare this ordinal outcome variable between intervention arm and surveillance. For the analysis of individual-reported adverse events, the population will include all randomised participants, and the comparisons will be by cross-tabulations and \( \chi^2 \) tests, since the probability of experiencing adverse events will likely not be affected by clustering. For the analysis of acceptability/compliance, only the intervention arm will be evaluated. We will use R software V.1.2.5019 and Stata software V.16 for the analysis.

**Data collection and management**

Any data collected during the course of this study will be protected under secure password, or kept under lock and key in the research office. No records bearing patient identification will be provided to anyone outside of the institution except regulatory agencies. Patients will not be identifiable as individuals in any publication or presentation of this study. Data will be recorded in RedCap and analysed as described above. The trial will follow all standard procedures, including reporting of adverse events. The protocol was written in accordance with Standard Protocol Items: Recommendations for Interventional Trials checklist (see online supplemental appendix 1).

**Patient and public involvement**

No patient or patient advisor was involved with study design, recruitment or conduct.
ETHICS AND DISSEMINATION

The study (protocol v3 09.06.2020) has been approved by the following boards: Commission Cantonale d’éthique de la Recherche, Geneva, Switzerland (2020-00864), Ethikkommission Nordwest- und Zentralschweiz, Comité Ethique Cantonale and Swissmedic (Swiss Agency for Therapeutic Products).

All participants will be asked for written informed consent with a dedicated member of the research team. This trial will be conducted in accordance with good clinical practice and the Helsinki Declaration. The WHO Trial Registration Data Set is in online supplemental appendix 2.

The COPEP-trial will establish whether LPV/r is effective as postexposure chemoprophylaxis against clinical COVID-19 and/or SARS-CoV-2 infection compared with surveillance among asymptomatic individuals with recent contact to a person infected with SARS-CoV-2. Furthermore, this trial will aim to investigate whether LPV/r reduces the severity of clinical COVID-19 over surveillance alone, and whether these prophylaxes are safe and acceptable for postexposure prophylaxis of SARS-CoV-2 and COVID-19.

If effective, such a pragmatic approach with prescription the same day and without waiting for laboratory results, will provide a feasible and relatively low-cost strategy to contain local COVID-19 outbreaks in future. All results will be disseminated in peer-reviewed journals and national and international conferences.

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Contributors
AC and NDL conceived the overall trial, including the overall design and therapeutic candidates and are the PI and Co-PI. AC, NDL, MSm, TP, MSI and AM designed the trial, including decision on intervention dosage and administration, study endpoints and procedures. MSm and TP developed all statistical aspects of the trial design. MSm will perform the trial data analysis, with the support of TP. DL and SF contributed to the formulation of recruitment via the cantonal health authorities, together with MB and CS. AM and DOA developed the patient and laboratory pathways. BM, LK and LD helped with all laboratory aspects of the trial design. MSm wrote the first draft of this manuscript. GJN, SR, AM, MSt, NDL and AC are the study physicians and together with FJ, TA, HS, JS, MB and FC who manage the respective patients with COVID-19 or infectious diseases unit contributed to the patient pathway. All authors contributed to the redrafting of the manuscript and in the process of approving the final draft.

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