Conflicts of interest. All authors report no conflicts of interest relevant to this article.

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To the Editor—In March 2020, a call to action was issued for antimicrobial stewardship programs (ASP) to assist in the SARS-CoV-2/COVID-19 response.1 Specific attention was focused on the common ASP infrastructures of prospective audit, existing partnerships with microbiology laboratories, and experience in stewarding medication resources as justification for ASP involvement. We leveraged our existing Enterprise ASP prospective audit platform to contribute to the response. Here we describe the logic and development of COVID-19 ASP flags, which were rapidly operationalized in the enterprise electronic medical record (EMR).

Our prospective audit system utilizes a longstanding, home-grown flagging system that was converted to function within the EMR (Epic Systems, Verona, WI).2-4 It generates a patient list based on a series of “rules” with complex logic incorporating medications and order elements, laboratory values, and microbiology, etc. The system also allows for the documentation of actions taken and provider response. Interventions deemed complete can be dismissed (ie, removed), and those requiring follow-up can be deferred for later review.

As the burden of COVID-19 patients began to increase, and amid concerns regarding medication shortages, our ASP needed a mechanism to identify patients with a SARS-CoV-2 polymerase chain reaction (PCR) testing performed and/or patients receiving medications in need of careful stewardship. We elected to incorporate both the PCR result and potential COVID-19 therapies into the rule logic because flagging the medications alone would not filter out non-COVID-19 indications, leading to the addition of low value flags or “noise” into the system. We considered flagging only PCR-positive patients, but this approach would fail to identify pending testing or PCR-negative patients who remained on potentially inappropriate medications. Conversely, incorporation of all ordered PCRs would also have contributed a great deal of noise.

We developed a hybrid approach by designing 2 flags that identify COVID-19 indications, leading to the addition of low value flags or “noise” into the system. The system also allows for the documentation of actions taken and provider response. Interventions deemed complete can be dismissed (ie, removed), and those requiring follow-up can be deferred for later review.

As the burden of COVID-19 patients began to increase, and amid concerns regarding medication shortages, our ASP needed a mechanism to identify patients with a SARS-CoV-2 polymerase chain reaction (PCR) testing performed and/or patients receiving medications in need of careful stewardship. We elected to incorporate both the PCR result and potential COVID-19 therapies into the rule logic because flagging the medications alone would not filter out non-COVID-19 indications, leading to the addition of low value flags or “noise” into the system. We considered flagging only PCR-positive patients, but this approach would fail to identify pending testing or PCR-negative patients who remained on potentially inappropriate medications. Conversely, incorporation of all ordered PCRs would also have contributed a great deal of noise.

We developed a hybrid approach by designing 2 flags that identify opportunities for stewardship of medications and confirm infectious diseases (ID) consultation.

The first rule (ASP COVID-19 rule 1) uses logic that identifies inpatients with a negative SARS-CoV-2 PCR test, collected within the previous 7 days, who also have a medication order that may represent “active therapy” (Table 1). Despite the 2 studies by Gautret et al5,6 claiming the benefit of the combination of

Practical implementation of COVID-19 patient flags into an antimicrobial stewardship program’s prospective review

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Fig. 1. A transparent shield at the slit lamp.

Acknowledgments.

Financial support. No financial support was provided relevant to this article.
hydroxychloroquine and azithromycin, we deliberately omitted azithromycin from the “active therapy” list.5,6 We recognized that despite significant weaknesses in this literature, providers may still order the combination; however, including azithromycin would have introduced flags for appropriately prescribed azithromycin for non–COVID-19 indications. Additionally, patients prescribed the combination of hydroxychloroquine and azithromycin would be flagged by the hydroxychloroquine order, thus making azithromycin inclusion unnecessary.

The second rule (ASP COVID-19 rule 2) is triggered by 2 criteria. The first is an inpatient with a positive PCR result irrespective of active medication orders. Our Enterprise team felt that review of all PCR-positive patients would be important to ensure appropriate involvement of the ID team, consideration for enrollment in clinical trials, and candidacy for off-label therapy. Had we stopped here, we would have realized a gap in the ability to review patients for which tests were ordered and pending. This feature was particularly important for facilities within the enterprise with a slower PCR turnaround time. The second criterion is an inpatient with a pending test, but to limit noise, the rule is only triggered when there is also an “active therapy” order. This criterion prompts the reviewer to use the flag “defer” logic for follow-up of test results so that, if positive, it may be re-reviewed by the ASP team. If the result is negative and therapy order remains active, ASP COVID-19 rule 1 is triggered.

Once either of the rules triggers a review, it is accompanied by text that displays the rule name, the active medication order contributing to the logic, and the date, time, and result of the PCR test. Developing rules that satisfy the needs of the Enterprise ASP as a whole required consideration of each facility’s typical flag burden, testing availability and turnaround time, availability of ID consultation, and onsite clinical trials. Notably, we are not currently an epicenter of the outbreak, and we recognize that, for facilities experiencing a high volume of COVID-19 hospitalizations, the tools described may not be applicable or may need modification prior to implementation. The landscape of COVID-19 management is rapidly evolving. Therefore, we remain nimble in our ability to add or subtract medications from the “targeted therapy” list, and we understand that as SARS-CoV-2 community prevalence or testing recommendations change, the rules should be modified to produce the highest benefit within limited ASP resources. Another factor that contributed to the success of our design is the availability of an internally developed SARS-CoV-2 PCR test. Facilities desiring to use test results as an element of the logic in their ASP triggers should assure that, regardless of testing location, the result is discreetly documented in the EHR.

We recognize that this functionality has limitations, and we anticipate that further challenges may arise. However, our goal is to carefully consider how to leverage existing infrastructure to effectively steward critical medication resources without overburdening the ASP team. We hope that describing our ASP’s efforts empowers others to identify optimal design, critical tasks, and high-value interventions contributing to the identification, triage, and management of COVID-19 patients. For healthcare teams of all kinds, it truly is time for “all hands on deck.”

### Table 1. Antimicrobial Stewardship COVID-19 Rule Logic

| Name | Criteria | Display |
|------|----------|---------|
| **ASP COVID-19 rule 1: negative SARS-COV-2 PCR w/ active drug order (inpatients only)** | IF negative COVID-19 PCR in last 7 days AND IF active order for 1 of the following: Chloroquine, Darunavir/ritonavir, Hydroxychloroquine, Lopinavir/ritonavir, Nazaoxanide, Remdesivir, Ribavirin, Sarilumab, Tocilizumab, Lenzilumab, IVIg | THEN fire alert |
| **ASP COVID-19 rule 2: positive SARS-COV-2 PCR or pending lab w/ active drug order (inpatients only)** | IF positive COVID-19 PCR in last 7 days OR IF pending COVID-19 PCR in last 7 days AND IF active order for 1 of the following: Chloroquine, Darunavir/ritonavir, Hydroxychloroquine, Lopinavir/ritonavir, Nazaoxanide, Remdesivir, Ribavirin, Sarilumab, Tocilizumab, Lenzilumab, IVIg | THEN fire alert |

Note. SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; PCR, polymerase chain reaction; COVID-, coronavirus disease 2019; IVIg, intravenous immune globulin.
Positive RT-PCR tests among discharged COVID-19 patients in Shenzhen, China

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To the Editor—According to the current guideline of the National Health Commission of China, discharge of inpatients with the coronavirus 2019 (COVID-19) infection in China have to fulfill 2 recovery criteria: (1) symptoms disappear and computed tomography (CT) images become normal and (2) test negative for 2 consecutive times in reverse transcriptase-polymerase chain reaction (RT-PCR) tests for SARS-CoV-2.1 However, Lan et al.2 recently reported 4 cases who were tested positive for SARS-CoV-2 at 5 days after discharge, suggesting positive status among discharged patients.3 To date, the prevalence and associated risk factors remain unclear.

We investigated all 209 patients with laboratory-confirmed SARS-CoV-2 infection who were discharged from the designated hospital in Shenzhen, China, between January 23 and February 21, 2020. Demographic data, laboratory profile, clinical data, and CT images were collected from these patients’ electronic medical records. Throat swabs and anal swabs were collected from all patients for RT-PCR tests according to the following scenarios: (1) on February 18, 2020, for those discharged before February 12, 2019; (2) on February 19, 2020 for those discharged between February 13 and 19, 2019; (3) on days 7 and 14 after discharge thereafter. This study was approved by the Shenzhen Center for Disease Control and Prevention review board and the need for informed consent was waived. All data used in this work are available upon request and approval of Shenzhen Center for Disease Control and Prevention.

We compared the settings in the study by Lan et al.2 with those in this study (Appendix Table S1 online). Logistic regression models were adopted to explore the factors associated with the RT-PCR test results. Odds ratios (ORs) were calculated for the probability of positive test in throat swabs, or anal swabs, or either, and the rest were considered negative in each of the 3 scenarios. The results are as follows:

- Scenario 1: 9 positive RT-PCR test results from throat swabs
- Scenario 2: 13 positive RT-PCR test results from anal swabs
- Scenario 3: 22 positive RT-PCR for test results from either throat or anal swabs

Normally, only scenario 3 should be considered, but we included scenario 1 to be consistent with Lan et al.2

Among all 209 discharged patients, 9 (4.3%) tested positive in throat swabs only, 13 patients (6.2%) tested positive in anal swabs only, and 22 (10.5%) tested positive in either. Together, 10.5% of discharged patients showed virus shedding around an average of 4.7 days after discharge (range, 2–13 days). Under scenario 3, the logistic regression models revealed that a high risk of positive test

Acknowledgments. We thank Brian Fung, PharmD, Evan Draper, PharmD, Aaron Tande, MD, and the Mayo Clinic Enterprise Antimicrobial Stewardship Team. We dedicate this article to the memory of Dr James Steckelberg, a pioneer of antimicrobial stewardship.

Financial support. No financial support was provided relevant to this article.

Conflicts of interest. All authors report no conflicts of interest relevant to this article.

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