Managing nirmatrelvir/ritonavir during COVID-19: pharmacists’ experiences from the Perak state of Malaysia

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Abstract

Novel therapeutic agents for SARS-CoV-2 have emerged over time, serving to reduce the severity of the disease, admission and mortality, especially among high-risk populations. Oral nirmatrelvir/ritonavir (Paxlovid®) was found to reduce the risk of disease progression. Pharmacists played multiple roles in handling the COVID-19 pandemic. This article highlights the roles of pharmacists in managing nirmatrelvir/ritonavir within the Malaysian context. Pharmacists were actively involved in Paxlovid® inventory management. To ensure the balance between supply and demand of new therapeutic drugs, pharmacists in health facilities constantly monitor the inventory levels of the medications. As Paxlovid® was initially reserved for a certain population who met the clinical eligibility criteria based on a scoring system, pharmacists were required to screen and exclude patients with non-indications or contraindications to the medication. During dispensing, pharmacists convey clear instructions on how to take the medications to ensure adherence and medication safety. The novel nature of the medications necessitates pharmacists to counsel patients regarding its indication, the mode of action, actions to take when missing a dose or overdose happens, side effects, storage and disposal methods, as well as mechanism of reporting adverse drug reactions. Pharmacists were required to follow-up all patients via phone call on Day 3 and Day 5 post-initiation, examining both adherence and adverse drug reactions associated with Paxlovid®. Pharmacists experienced multiple challenges in managing Paxlovid®, particularly due to increased workload, suboptimal follow-up response, stringent medication storage requirements, and adherence issues. Universal research and innovation initiatives were proposed to improve the delivery of novel therapeutic agents in the future health system.

Introduction

The SARS-CoV-2 Omicron subvariants BA.2.12.1 and BA.4/5 have shown a sharp rise. The effectiveness of COVID-19 vaccines and therapeutic monoclonals may be jeopardized by these new subvariants, which carry further mutations in their spike proteins. BA.4/5 is 4.2-fold more resistant, and as a result, more likely to cause vaccine breakthrough infections [1]. Meanwhile, novel therapeutic agents have emerged over time, serving to reduce the severity of the disease, admission and mortality, especially among high-risk populations. Oral nirmatrelvir/ritonavir (Paxlovid®) combination, a protease inhibitor, was introduced in early 2022. It was reported that this combination could reduce the risk of disease progression significantly by 89%, with a considerably safe profile [2].

Pharmacists played multiple roles in handling the COVID-19 pandemic, including innovating medication delivery and monitoring systems; inventory management and procurement; development of treatment guidelines; supporting clinical trials; and ensuring the continuity of pharmaceutical care and counselling [3–6]. The roles

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of pharmacists in handling novel drugs have not been reported previously. This article highlights the roles of pharmacists, challenges, and potential innovations in managing Paxlovid®, a novel therapeutic agent within the Malaysian context.

**Inventory management and distribution of medicines within the hospital**

The Perak state consists of five publicly funded tertiary hospitals with specialists, 11 secondary hospitals without specialists, and 11 district health offices. Pharmacists’ leadership was crucial in the face of drug shortages during COVID-19 [7]. To ensure the balance between supply and demand for Paxlovid®, pharmacists in these facilities were tasked to constantly monitor the inventory levels of Paxlovid®. Online spreadsheets were used to collect and monitor the usage of Paxlovid® across different facilities. Pharmacists at the ground level were required to fill up the electronic inventory spreadsheet, which detailed input movement and the current stock level of Paxlovid® on a daily basis. From the central level, pharmacists at the State Health Division closely monitored Paxlovid® levels and activated an inventory mobilization plan, which involved the transit of Paxlovid® from the major hospitals to the health district offices once the pre-determined minimum stock level was triggered. Meanwhile, liaison pharmacists in these facilities were required to update the patient administration registry and return medication registry, comprising the age, ethnicity, COVID-19 category, regimen, as well as the start and end date of therapy on a weekly basis.

In the initial stage, Paxlovid® was available for patients who visited publicly funded hospitals and health clinics. Towards the end of June 2022, the supply of Paxlovid® has been expanded to benefit patients who seek treatment from private facilities, including private hospitals and general practitioners [8]. Once the letter of intent and application form were received, the liaison pharmacists in charge of inventory management from public health clinics were required to mobilize inventory to the private health facilities. During the transportation of Paxlovid®, pharmacists were required to maintain the temperature between 15 and 25 °C to ensure stability [9]. Simultaneously, site pharmacists filled up the electronic Paxlovid® inventory spreadsheet, which was subsequently reported to the Pharmacy State Health Division.

**Screening of patients**

Screening of patients is routinely performed by pharmacists [10]. Paxlovid® was reserved for a certain population who met the clinical eligibility criteria based on a set of scoring systems [11]. It was developed through reviewing of current literature and consensus among clinicians, including family medicine specialists, infectious disease specialists, and medical specialists [12]. Briefly, patients who were symptomatic for less than 5 days with a clinical stage of 2 or 3 were considered potential candidates for Paxlovid® [7]. This population of patients was further stratified based on age, immunity, comorbidities, obesity, smoking, as well as vaccination status. Older adults, immunocompromised patients, patients with multiple comorbidities, obese and smoking patients, and those who did not complete COVID-19 vaccination were prioritized, given the fact that this population inherited a greater risk of progressing to severe COVID-19 stages [11, 12].

To ensure patients’ safety, pharmacists played an essential role in screening and evaluating the clinical history of patients before the initiation of Paxlovid® [10]. Pharmacists were tasked to identify patients with contraindications, such as those less than 18 years old, with symptoms onset more than 5 days, patients who required oxygen, with severe hepatic impairment (Child–Pugh Class C), severe kidney impairment (eGFR < 30 ml/min), pregnancy or breastfeeding. For patients with moderate renal impairment (eGFR 30–60 ml/min), renal adjustment dosage was recommended. Furthermore, pharmacists screened for hypersensitivity to the Paxlovid® active ingredients or excipients, contraindicated medications, as well as other drug interactions based on information on the Infectious Diseases Society of America COVID-19 treatment and management guideline [13] and online drug interaction checker [14].

Pharmacists also act as an intermediary to facilitate communication between prescribers and patients [15]. This was crucial as Paxlovid® was newly introduced and not prescribed routinely. In this case, patients can exert an informed decision after counselling by the pharmacists. They can choose whether to take Paxlovid®. Patients were informed by the pharmacists that they were suitable candidates, and their decision would be communicated by the pharmacists to the prescribers [15]. Patients who rejected Paxlovid® were reviewed by the prescribers. Prescribers then decide whether to omit the treatment or provide an alternative based on the risk–benefit ratio, i.e., judging on whether the patient will progress to a more severe COVID-19 stage.

**Dispensing**

Before dispensing to the patients, a pharmacist’s core responsibility is to assure that the prescriptions are accurate so that patients receive the correct medication and the appropriate dosage [10]. At this point, pharmacists will assess the appropriateness, efficacy, and safety of Paxlovid® for each patient. Using the “five rights” practise [16] necessitates a pharmacist extensively reviewing each
In our setting, pharmacists were required to follow-up all patients started on Paxlovid® via phone call on Day 3 and Day 5 post-initiation, examining both adherence and ADR. To improve call response, patients were informed before discharge or during their last encounter that they would be contacted by pharmacists on Day 3 and Day 5 following Paxlovid® initiation. The schedule was rationalized as follow-up patients after the 1st week of discharge could be beneficial in reducing readmissions [26]. Pharmacists recorded Paxlovid® treatment adherence and ADR into the existing patient administration registry based on patients’ self-reports. In non-adherence cases, pharmacists explored the reasons and provided additional counselling. If patients insisted on stopping Paxlovid® treatment, the pharmacist advised them to return the medication to the nearest pharmacy unit.

Pharmacists are also responsible for reporting Paxlovid®-related adverse drug reactions to a central database hosted by the National Pharmaceutical Regulatory Agency, through either the pharmacy information system, online web form, or manual submission [27]. Patients were advised by pharmacists to seek medical assistance at the nearest health clinic if they experienced Paxlovid®-related ADR, especially if the event was severe or unusual.

Challenges
We experienced several challenges in managing Paxlovid®, including follow-up, workload, medication storage, and adherence (Fig. 1). One of the major obstacles faced by pharmacists in performing follow-up was the limitation of the manual phone call method. Calling patients manually entails a high workload, involving a team of designated pharmacists. Each Paxlovid®
follow-up phone call lasted about 10 min and each patient required three follow-ups. This was compounded by the pharmacists’ existing workloads, who were expected to meet the patient waiting time target as well as complete other clinical and administrative duties. At the time of writing, there were more than 650 patients started with Paxlovid® in Perak state.

Moreover, unanswered phone calls complicate the follow-up efforts. Some patients did not pick up calls after discharge, whilst some provided invalid phone numbers. Based on the local protocol, pharmacists were required to call the same number at least three times before excluding the patient from Paxlovid® phone call follow-up. The traditional phone call follow-up method entailed a high drop-out. In the United States, almost half of the patients could not be reached by phone call after discharge [28]. Furthermore, a minority of underprivileged population from lower socio-economic backgrounds did not own a phone, posing a different challenge in Paxlovid® phone call follow-up.

Besides, stringent storage and transport requirements must be met to ensure the stability of Paxlovid®. According to the FDA [9], Paxlovid® must be stored at a controlled room temperature between 20 and 25 °C. Any excursion below 15 °C and exceeding 30 °C may cause the drug to no longer be usable. Pharmacists were required to monitor the temperature of Paxlovid® closely throughout the transit, storage, and dispensing process.

Ensuring adherence to Paxlovid® therapy was another challenge faced by pharmacists. Some patients deliberately stopped their medication, rationalizing that their symptoms had improved over time. While pharmacists strive to explain the importance of treatment adherence, this could not be achieved in every patient. From a clinical point of view, this may compromise the efficacy of Paxlovid® therapy. From the economic perspective, the omission of doses causes wastage, as the average cost of one complete Paxlovid® regimen was estimated at USD 250 [29].

Research and innovation
Innovative measures could be considered to overcome administrative, technical, and clinical challenges (Fig. 1). In the face of the COVID-19 pandemic, virtual patient follow-up via phone and video calls was crucial to enhance patients’ access to healthcare [30]. However, the manual phone call follow-up to monitor Paxlovid® adverse events was resource intensive. A previous study found that calling patients using an Interactive Voice Response System was not inferior to human-manual calls [31]. This could largely reduce the pharmacist’s burden, provided that the system is mature enough to
be distinguished from a spam call. Meanwhile, as some patients might feel intruded by a telephone call, alternative communication methods such as email and texting could be offered at the point of discharge to improve the follow-up response rate [32].

The Malaysian public hospitals serve a diverse population, including hard-to-reach and marginalized people, who have not responded positively to our Paxlovid® follow-up phone call. Active engagement of pharmacists through network and relationship building with key stakeholders such as the Department of Orang Asli Development (JAKOA), existing community organizations, and health volunteers could be instrumental in designing effective and culturally sensitive Paxlovid® follow-up programmes to improve access for this population [33–35].

Pharmacists play an instrumental role in optimizing patients’ adherence to Paxlovid®. Timely review and integration of adherence-based mobile application into medication counselling serves as an important measure to improve Paxlovid® adherence [36]. Preliminary evidence suggests that mobile applications may improve patients’ adherence to medications [37]. Integration of the Paxlovid® adherence monitoring function into the widely used MySejahtera application could be one cost-effective option [38]. Moving forward, the feasibility and clinical impact of integrating artificial intelligence into these mobile applications should be explored [39].

Traditionally, ADRs are reported by healthcare professionals, including pharmacists. However, underreporting of ADR was common due to administrative and technical barriers [23]. Implementation of an electronic ADR reporting system could be instrumental in facilitating and promoting Paxlovid®-related ADR reporting, especially when such a system is integrated into the hospital information system [40]. Furthermore, the public should be educated on ADR reporting mechanisms to reduce ADR reporting workloads among health professionals. To facilitate reporting, a user-friendly interface should be introduced, for example by integrating such a function into the MySejahtera application.

Conclusion
Pharmacists are holding the ground firmly during the COVID-19 pandemic. The discovery of novel treatment agents such as Paxlovid® holds pharmacists accountable for inventory management, screening, dispensing, counselling, monitoring, and reporting, among others. Wearing multiple hats, a pharmacist’s involvement in research is critical to innovative communication and pharmacovigilance systems. Advancements in technologies warrant digitalization in pharmacy services. Transformation could begin by shifting from manual follow-up and reporting to electronic and automation models, encompassing artificial intelligence in the near future.

Acknowledgements
We would like to thank the Director-General of Health Malaysia for his permission to publish this article.

Author contributions
All authors contributed equally to this work including conceptualization, data curation, writing and review of manuscript. All authors read and approved the final manuscript.

Funding
No external source of funding received for this work.

Availability of data and materials
The datasets used in the current study are not publicly available, but are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
This study was registered in the Malaysia National Medical Research Registry (NMRR-ID-22-01555-LOS). The work was conducted according to the Malaysian Guidelines Good Clinical Practice (4th edition) and other relevant guidelines for research.

Consent for publication
Not applicable.

Competing interests
The authors declare no conflict of interest.

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Received: 7 October 2022   Accepted: 13 October 2022

Published online: 23 October 2022

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