SARS-CoV-2 vaccine related adverse events in Zimbabwe: the need to strengthen pharmacovigilance in a resource-limited settings

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Dear Editor

The emergency use approvals (EUAs) of vaccines for the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) by the World Health Organisation (WHO) starting in December 2020 was a welcome event [1]. Since their discovery, vaccines have become one of the best public health interventions to reduce the spread, morbidity and mortality associated with rapidly progressive infectious diseases. Unlike previous vaccines, SARS-CoV-2 vaccines were rapidly developed and authorised at a time when their safety and effectiveness data were not widely publicly available [2]. Understandably, given the unprecedented socio-economic burden, the COVID-19 pandemic presented [3], expedited solutions for control, including vaccination, were warranted. Whilst part of the Janssen vaccine clinical trials were conducted in South Africa, safety and effectiveness studies for the majority of the vaccines were conducted outside the Sub-Saharan Africa region. Therefore, safety data specific for this population group were lacking.

Zimbabwe approved the Sinopharm and Sinovac vaccines in February 2021, and subsequently, the Covaxin, Sputnik V and the Janssen vaccines. The first four have been in use in the country. The Janssen vaccines are not yet available. The Sputnik V vaccines are still to be approved by the WHO. The Medicines Control Authority of Zimbabwe (MCAZ), in line with its mandate prescribed by the law, approved the first four before they had EUAs from the WHO and before there were any publicly available safety data. This resulted in scepticism and safety concerns in the general population, which may have contributed to vaccine hesitancy. As of 1 October 2021, Zimbabwe had administered approximately six million doses of COVID-19 vaccines, the majority of which are Sinopharm and Sinovac vaccines. In this opinion piece, we highlight the need to maintain robust processes for pharmacovigilance by the MCAZ to identify and monitor potential SARS-CoV-2 vaccine-related adverse events.

Anecdotal evidence exists of adverse events (AEs) such as headaches, skin reactions and flu-like illnesses, with even deaths having been attributed to vaccination without adequate evidence-based for. For example, in Zimbabwe, early in the vaccination programme, there were widespread rumours and vaccine hesitancy after a healthcare worker had reportedly died after being vaccinated [4]. Other widely circulated myths relate SARS-CoV-2 vaccines to reproductive and menstrual cycle disturbances, which have raised concerns and propagated hesitancy among women of reproductive age [5]. Reports of miscarriage, erectile dysfunction and other undesirable AEs following vaccination have emerged [6]. These are just some examples of unsubstantiated AEs that have been circulated widely for which neither evidence of causation or association is present or has been noted in any clinical settings. Though widely circulated on social media even beyond Zimbabwe, no convincing clinical evidence to substantiate the claims exists to date. If not addressed adequately, misinformation propagates widespread vaccine hesitancy, making it difficult for the country to reach its 60% vaccination target by December 2021,
and the herd immunity threshold difficult to attain. The major significant problems currently include lack of safety and effectiveness data of the currently available vaccines in Zimbabwe for the local population, and lack thereof in comparable settings. This is compounded by a lack of clear and effective communication strategies by the relevant COVID-19 control pillars in the MoHCC. The space is overtaken by social media, which now exists in several forms, and it is easily accessible to the wider population, providing fertile ground for circulating unsubstantiated rumours, myths, falsehoods and misconceptions, blowing events out of proportion and propagating widespread vaccine hesitancy.

Zimbabwe has an established system of pharmacovigilance through the MCAZ. Unfortunately, because the pharmacovigilance system in existence is unknown to many, the tendency to run to social media for quick answers is increasing. The critical public question at this point is the strength of this system in its current form to monitor the safety of the SARS-CoV-2 vaccines specifically. Given that none of the clinical trials of the vaccines currently being used in Zimbabwe was conducted in this population, or in Sub-Saharan African countries, it is important that the relevant authorities set up a prospective drug safety follow up programme in the country to reassure the population that their safety concerns are carefully monitored. Strengthening the system, and putting in place a strong evaluation system to provide strategic information regarding the effect and implementing a solid evaluation system to provide strategic information regarding the effect of SARS-CoV-2 vaccines is as critical as availing the vaccines to the population. Taking advantage of existing platforms of the Zimbabwe Expanded Programme of Immunisation (ZEPI), which has been in existence since 1982, could provide an important baseline for the country [7].

As part of the solutions for these reasons, more sophisticated reporting and capturing systems for adverse events are urgently needed. Similar to reporting systems-procedures for suspected SARS-CoV-2 infection, which included 24 hours toll-free numbers and emergency call centres, and electronic data capturing and transmission systems for timely reporting, having specific ways of following up vaccines’ adverse events is important. All service providers involved in vaccination and healthcare must be adequately made aware of these systems, so they can work closely with the MCAZ and other relevant stakeholders to make such a system functional, effective, efficient, and informative. The public must be made aware of the existence of this system so that they can also fully utilise it, by making enquiries for any concerns for suspected vaccine-related AEs. The MoHCC and MCAZ need to put in place precise mechanisms for reporting these suspected AEs and circulate them widely sufficiently to allow greater population reach. Additionally, the risk communication and community engagement (RCCE) pillar, which

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is responsible for disseminating health promotion and protection messages in Zimbabwe must becoming actively involved in engaging the community regarding AEs related to SARS-CoV-2 vaccination, allaying the population’s anxiety, identifying key barriers of vaccine uptake and propagators of vaccine hesitancy. As the vaccination programme is extended to special populations such as children and pregnant and breastfeeding women, the MCAZ must up its pharmacovigialce game, whilst the RCCE pillar and other relevant stakeholders strengthen their messages to increase vaccine uptake momentum and propel the country towards its herd immunity threshold.

It has become apparent that the COVID-19 pandemic is likely to become protracted, and several SARS-CoV-2 variants of concern continue to emerge. As Zimbabwe and many other countries prepare for further inevitable waves of the COVID-19 pandemic, it is important to promote vaccination, and prevention and control strategies. Fighting vaccine hesitancy is indispensable, and providing prospective vaccine recipients with reassurance and meaningful safety data is critical [8]. There is therefore an urgent need to strengthen pharmacovigilance systems in the country, and operational research to provide the much needed answers regarding the question of vaccine-related adverse events. To this end, the MCAZ and other relevant stakeholders involved in public health and pharmacovigilance must converge urgently to improve, optimise and strengthen the existing systems. There is an urgent need for the strengthening of pharmacovigilance in Zimbabwe, a limited resource country, to use tailored, robust reporting or other pharmacoepidemiological methods that systematically collect and analyse adverse events associated with the use of COVID-19 vaccines, identify signals or emerging problems, and communicate how to minimise or prevent harm.

**Competing Interests:** None to declare.

**References**


