Summary Brief

A Wake-Up Call to Improve Country-Wide Reporting of Adverse Drug Reactions in Sierra Leone

Key Messages

- This first country-wide assessment of reporting of Adverse Drug Reactions (ADRs) to antimicrobials in Sierra Leone shows inconsistency, with reporting delays and incomplete data. This implies that ADRs are being underreported, and this can compromise clinical care and drug safety.

- 90% of all reports were from active reporting from mass drug administration campaigns, while 10% came from voluntary (passive) reporting from health facilities.

- This is a wake-up call to improve monitoring by introducing compulsory (active) reporting and setting performance targets for completeness and timeliness of ADR data as part of national pharmacovigilance guidelines.

What is the problem and why is it important?

The World Health Organization (WHO) advocates for vigilant monitoring of adverse drug reactions (ADRs) to antimicrobials as they can cause life-threatening illness, permanent disabilities, and death.

Since 2007, Sierra Leone has established a national database for monitoring ADRs (VigiFlow). However, there is insufficient insight into the quality of these data and “what works and what needs to be improved”. Poor reporting would compromise data utility in terms of clinical practice, drug safety and setting up drug regulation systems.

How did we measure it?

We assessed countrywide ADR reporting on antimicrobials using individual case safety reports (ICSRs) entered into VigiFlow between 2017–2021.

Reference:


Contact: e-mail: fthomas@pharmacyboard.gov.sl
What did we find?

- Of 566 ICSRs, inconsistent reporting was seen, with peaks in 2017 and 2019 (mass drug campaigns for deworming), zero reporting in 2018 (reasons unknown), and only a handful of reports in 2020 and 2021 (since COVID-19).
- 90% of all reports were from active reporting from mass drug administration campaigns, while 10% came from voluntary (passive) reporting from health facilities.
- There were reporting delays, with 90% of reports taking over 30 days to be entered into VigiFlow (maximum threshold = 30 days).
- Data was incomplete: 57% of reports had variables not filled in.
- Final patient outcomes in 36% of reports were not available.

Implications

- This first country-wide study from Sierra Leone shows inconsistent reporting with delays and incomplete data that can be improved.
- Possible ways forwards to improve monitoring include:
  1. Introduction of active compulsory ADR reporting at all health facilities.
  2. Enforcing performance targets for monitoring, including:
     - Serious ADRs are to be reported within 7 days and other ADRs within 30 days
     - All key variables are to be completed in ADR forms
  3. Optimizing the use of the already existing online electronic ADR reporting system using mobile phones and/or computers.
- Improved monitoring and reporting of ADR data will have major benefits for the clinical management of patients, which in turn could prevent life-threatening illness, permanent disabilities, and death.