Antiretroviral prescriptions with potential drug-drug interactions from general practitioners and specialists

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To the Editor: It has been estimated that globally about 38.6 million people were infected with HIV by 2005,1 with about 5 million of them living in South Africa. The World Health Organization estimated that 4.7 million people living in sub-Saharan Africa urgently needed antiretroviral therapy (ART). In that year SA implemented prescribed minimum benefits (PMBs) for HIV/AIDS in the private health care sector.2 Despite the increased availability and affordability of ART in SA, only 60 000 people were receiving ART through medical aid schemes by mid-2005.3

Antiretrovirals (ARVs) have transformed HIV/AIDS into a chronic disorder that can be managed effectively. The right of all HIV-infected adults and children to receive standard care is endorsed by the SA HIV Clinicians Society (SAHIVCS),4 with ART guidelines recommending different combinations. The rapid approval of new drugs resulted in an increased risk of prescribing errors, dispensing of incorrect dosages/dose frequencies, and incorrect reporting of drugs by the patient to the prescribers,5 all leading to treatment failure.

Drug–drug interactions (DDIs) are an under-recognised consequence of medication prescription errors, resulting in significant health care costs.6 Since DDIs determine positive and negative consequences of treatment for HIV-infected patients, recommendations to avoid some drug combinations and to adjust dosages of some co-administered drugs were formulated by both the SAHIVCS4 and the Department of Health.7 The private health care environment recommends the Aid for AIDS ART guidelines, which are similar to those of the SAHIVCS (Table I).

We investigated the prevalence of prescriptions with potential DDIs between ARVs from general practitioners (GPs) and specialists (SPs) for patients in different age groups in the private health care sector, and evaluated the prescribed daily dosages (PDDs).

Methods
A non-experimental, retrospective, quantitative study was performed on data from a South African Pharmacy Benefit Management (PBM) company from 1 January 2005 to 31 December 2007. ARV drug names were classified according to the pharmacological groups in the Monthly Index of Medical Specialities (MIMS) and the Regulations to the Medicine and Related Substances Act (Act 101 of 1965). Prescribers of ARVs were divided into GPs and SPs.

We evaluated potential DDIs between ARVs using PDDs. It is important that the PDD be related to the diagnosis made for the prescribed medication. The reference guides used to evaluate PDDs were according to the recommended ARV dosing.2,5

Basic descriptive statistics, i.e. frequencies, the arithmetical mean (average), standard deviations and effect sizes (Cohen’s d) were used to characterise the study sample, and were calculated using the computer software Statistical Analysis System (SAS) for Windows 9.1. The age groups used were as follows; group 1 – <0≤12 years, group 2 – >12≤19 years, group 3 – >19≤45 years, group 4 – >45≤59 years, and group 5 – >59 years.

Confidentiality of information was maintained, as no patient, medical practice, pharmacy or medical scheme could be identified. Permission to conduct the study was granted by the PBM company, and approval was obtained from the research and ethics committees of North-West University, Potchefstroom campus, (ethical number 07M01) and Walter Sisulu University, Mthatha campus.

Results
The study was performed on 49 995, 81 096 and 88 988 ARV prescriptions prescribed to 7 664, 10 162 and 10 061 HIV patients in 2005, 2006 and 2007, respectively. ARV prescriptions represented 0.59% (N=49 995), 0.90% (N= 81 096); and 1.11% (N= 88 988) of the claims for the 3 years.

In the 3 years, GPs prescribed more ARV prescriptions than SPs, and these increased from 2005 to 2007. The highest proportion of prescriptions was for three ARV items, followed by two ARV items. No practically significant differences were found between the average numbers of ARV medicine items per prescription (d<0.8) claimed per year for the different years.

ARV prescriptions from GPs with potential DDIs and incorrect PDDs increased from 12.33% in 2005 to 24.26% in 2007; those from SPs increased from 15.46% in 2005 to 33.16% in 2007. The highest numbers of incorrect PDDs with DDIs were identified in ARV combinations of lopinavir/ritonavir 1 066.4 mg/264 mg with efavirenz 600 mg, and lopinavir/ritonavir 1 066.4 mg/264 mg with nevirapine 400 mg, followed by indinavir 1 600 mg with ritonavir 800 mg, ritonavir 600 mg with efavirenz 600 mg, and saquinavir 800 mg with efavirenz 800 mg, for both GPs and SPs. The highest number of prescriptions were prescribed by GPs to patients in the age group >19≤45 years, followed by the age group >45≤59 years.

Discussion
We aimed to determine the prevalence of ARV prescriptions with potential DDIs and to evaluate their PDDs with specific reference to prescriber and age group. DDIs are of particular concern in HIV patients receiving highly active antiretroviral therapy. Although combination ARV treatment is potent and effective for HIV infection, ARVs frequently interact among themselves as many are metabolised...
The ARV regimens most identified with incorrect PDDs were between lopinavir/ritonavir (PI) prescribed at PDDs 1 066.4 mg/264 mg, 4 500 mg/3 999 mg and 1 599.6 mg/264 mg with efavirenz (non-nucleoside reverse transcriptase inhibitor) (NNRTI), then the same with nevirapine. These were followed by indinavir (PI) with ritonavir (PI), saquinavir (PI) with efavirenz, and saquinavir with ritonavir. All these combinations were prescribed to patients aged >19-45 years. The standard adult dose for lopinavir/ritonavir is 400 mg/100 mg twice daily or 800 mg/200 mg once daily. A limitation of this study was that information about whether patients were HIV naive or experienced was not available. Once-daily dosing for lopinavir/ritonavir is only recommended for treatment-naive patients, and not for patients receiving efavirenz, nevirapine or nelfinavir. When lopinavir/ritonavir is given with efavirenz or nevirapine, the recommended dose for treatment-experienced patients is 600 mg/150 mg twice daily with food.

Furthermore, the results revealed that GPs prescribed lopinavir/ritonavir for patients in the age group 0-12 years. Another limitation of this study was that the weights for children were not available and it was not clear to which weight and age categories of patients this co-formation was prescribed. The safety, efficacy and pharmacokinetic profiles of lopinavir/ritonavir in children under the age of 6 months have not been established. The paediatric dose for lopinavir/ritonavir as recommended in the SA ART guidelines is 12 mg lopinavir/kg twice daily for children weighing <15 kg and 10 mg lopinavir/kg for children ≥15 kg. In this case lopinavir/ritonavir was prescribed in a higher PDD, considering one capsule of lopinavir/ritonavir to be 133.3 mg/33.3 mg, with the maximum dose not exceeding 399.9 mg/99.9 mg. If given in higher PDDs the patient may therefore experience side-effects such as diarrhoea, nausea, vomiting, hyperlipidaemia and glucose intolerance.

We observed that ritonavir was given with other PIs in PDDs of 3 000 mg, 1 200 mg, 800 mg and 600 mg. The recommended dosages of 100 mg capsules or 600 mg/7.5 ml solutions in adults are 600 mg every 12 hours (when ritonavir is used as sole PI). As a pharmacokinetic booster for other PIs, the recommended dosage is 100 - 400 mg per day in two divided doses. Boosted PI regimens that utilise a low dose of ritonavir (100 - 200 mg) appear to offer the best balance of efficacy and tolerability.

Limitations

Limitations of this study include non-availability of patient clinical data to do in-depth clinical evaluation of the potential DDIs and PDDs. The extent to which potential DDIs actually occurred could therefore not be determined. Information on HIV-naive or experienced patients, CD4 values and viral loads were not available from the database. The clinical relevance of the identified potential DDIs was evaluated according to criteria in the literature. Information about the pharmacists who dispensed the prescriptions was not available. Pharmacists in collaboration with the prescriber could have detected and corrected prescriptions with potential DDIs and incorrect doses, this being one of the primary responsibilities of specific pharmacists. However, the results emphasised the possibility of DDIs that could have caused severe problems. A further limitation of the study was that no differentiation could be made between the prescription patterns of dispensing doctors versus pharmacy-dispensed prescriptions.

Conclusion

Our results emphasise the important role of prescribers and pharmacists in ensuring the appropriate use of ARVs. The importance of educating prescribers and dispensers by medical schemes or PBM companies in the private health care sector is also emphasised.

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References


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