Results  Drugs dispensed by ADSs represent 60% and 71.6% respectively of total medicines consumed in ICU and ED. Four years after implantation:

- The quantity of drugs dispensed and drug stock has decreased in both units.
- The pharmacy department knows the type and amount of medicines to be found in each unit and in real time.
- The information it provides has helped to improve patient safety in relation to a better quality of prescription.

Since the implementation of ADSs, consumption has decreased compared to 2007:

<table>
<thead>
<tr>
<th>Year</th>
<th>2007 Consumption</th>
<th>2008 Consumption</th>
<th>2009 Consumption</th>
<th>2010 Consumption</th>
<th>2011 Consumption</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICU</td>
<td>$-12.62%$</td>
<td>$-16.98%$</td>
<td>$-20.25%$</td>
<td>$-40.65%$</td>
<td></td>
</tr>
<tr>
<td>ED</td>
<td>$-1.49%$</td>
<td>$-14.94%$</td>
<td>$-29.18%$</td>
<td>$-40.79%$</td>
<td></td>
</tr>
</tbody>
</table>

Conclusions  The implementation of ADS has meant an estimated saving of 938,330 €.

The ADSs have increased drug control by the pharmacy department, have achieved a better rationalization of resources and have improved efficiency in drugs use.

No conflict of interest.

DSL-012  EVALUATING THE STANDARDS OF HOSPITAL PHARMACIES IN THERAPEUTIC CENTRES AFFILIATED WITH KERMANSHAH UNIVERSITY OF MEDICAL SCIENCES, IRAN

doi:10.1136/ejhpharm-2013-000276.255

1. Kermanshah University of Medical Sciences, Pharmaceutics, Kermanshah, Iran; 2. Tabriz University of Medical Sciences, Pharmaceutics, Tabriz, Iran; 3. Tabriz University of Medical Sciences, clinical pharmacy, Tabriz, Iran

Background  Nowadays pharmaceutical care departments located in hospitals are amongst the important pillars of the healthcare system.

Purpose  To evaluate the quality of hospital pharmacies affiliated to the Kermanshah University of Medical Sciences.

Materials and Methods  In this cross-sectional study a validated questionnaire was used which enquired about all the necessary and standard requirements of an ideal hospital pharmacy. The questionnaire was filled in by one of the researchers in all seventeen hospital pharmacies located in the teaching and non-teaching hospitals affiliated to the Kermanshah University of Medical Sciences. Data analysis was done using SPSS (version 17).

Results  The results show that in the hospitals observed, 24% of pharmacy environments, 25% of pharmacy store and storage conditions, 49% of storage procedures, 25% of drugs ordering and supplies, 75% of supplies reception (proper procedures followed for receiving supplies), 35% of supplies reception (prompt action taken if deterioration of drugs received is suspected), 23.35% of drugs supplied to patients and finally 0% of stock cards used for inventory control met these standards in full. Several instances of incorrect processes of ordering, receiving, storing and delivering medicines to the patient were detected that had led to wasted money in hospitals and considerable decrease in the quality of medical services.

Conclusions  Non-standard space allocation, incorrect ordering, receiving, storing processes and delivery of medicines in the patient were revealed by the questionnaire. These issues may reduce the efficiency and safety of pharmaceutical services and drug administration in hospitals.

No conflict of interest.

DSL-011  EFFICACY AND SAFETY OF EPOETIN ZETA IN DIALYSIS PATIENTS

doi:10.1136/ejhpharm-2013-000276.254

1. B Bajraktar, 2. B Lazareva, 3. E Najdovska, 4. Mihalova, 5. Institute of nephrology, Hospital Pharmacy, Struga, FYROM; 6. Clinical Hospital, Hospital Pharmacy, Stip, FYROM; 7. Clinical Hospital, Hospital Pharmacy, Bitola, FYROM

Background  Anaemia in chronic kidney disease (CKD) remains one of the predictable and modifiable non-traditional cardiovascular risk factors. Epoetin zeta, which is a biosimilar product, is used in the treatment of anaemia associated with chronic kidney disease.

Purpose  This study was performed to evaluate the efficacy and safety of the biosimilar product epoetin zeta to maintain stable haemoglobin levels in dialysis patients.

Materials and Methods  This study was conducted in 2 dialysis centres with 35 patients. Before the study 30 of the 35 patients were treated with biosimilar Epoetin zeta and were followed for 6 months. The initial weekly doses as well as the frequency of use per week were kept constant (1–3 times/week). During the follow-up, haemoglobin levels, iron status, dialysis efficiency, body weight and adverse events were monitored at least once a month.

Results  33 patients were treated with biosimilar Epoetin zeta (27 men and 6 women); average age 59.1 (28–76) years; the frequency of use was 1–3 doses/week subcutaneously, over a period of 6 months. Dosing was to be adjusted to keep the Hb levels within 10.5–11.5 g/dl ± 0.5 g/dl. The main AEs (adverse events) were in 1 patient hypotension (3%), in 1 patient in-dialyzer clotting (3%) and SAE (serious adverse event) was in 1 patient thrombosis of arterovenous fistula (AVF) (3%). No anti-epoetin antibodies and no clinical signs of pure red cell aplasia (PRCA) were observed in any patients on the study.

Conclusions  Treatment of anaemia with Epoetin zeta was shown to be effective and safe. The mean Epoetin zeta doses remained stable in patients switched from all pre-study ESAs. The observed adverse events profile was in line with expectations for the study population.

No conflict of interest.

DSL-013  EVALUATION OF THE LIMITS OF AUTOMATION AND IMPACT ON DRUG MANAGEMENT AT MOHAMMED V MILITARY TEACHING HOSPITAL PHARMACY, RABAT, MOROCCO

doi:10.1136/ejhpharm-2013-000276.256


Background  Nowadays, hospitals tend to automate medicines management to increase quality, efficiency and safety of drug dispensing. At Mohammed V Military Teaching Hospital (MVMTH), a decentralised Automated Drug Dispensing System (ADDS) was installed at the duty pharmacy. We expect this experience will be decentralised to all hospital services.