

(VE) for the treatment of otomycosis and analyse its effectiveness and safety.

**Material and methods** Antifungal ear drops are not commercially available. The otolaryngology service requested a broad spectrum topical antifungal for recurrent otomycosis. After a literature review, a sterile aqueous formulation of voriconazole 10 mg/mL was considered, ensuring the absence of ototoxic effects, with an optimal pH of 6.3 that allowed contact with the external channel. We assigned a beyond use date of 14 days refrigerated, 45 days frozen and protected from light.

Baseline data were collected from the clinical history. Patients reported their outcomes in interviews with the pharmacists: humidity, otorrhoea, earache, itching, loss of hearing before/after treatment and possible adverse events (AE) were recorded. Patients were informed and consent was requested for participation. Statistical analysis was made with SPSS and STATA. The results were analysed using the McNemar test of paired data.

**Results** Following the macroscopic finding of hyphae, microbiological culture was requested in 55.5% of cases, and *Candida* (33%) and *Aspergillus* (22%) isolates were found. All patients were treated previously with topical drugs (94.4% antibiotics, 55.5% antifungals) and 83.3% also with oral agents (83.3% antibiotics, 22.2% antifungals), without improvement. Eighteen patients (58.8% women, median age 67 years (range 44.5–75)), were treated with VE for an average of 4 weeks (SD 1.8), administering 1–2 drops 2–3 times a day.

Interviews were conducted in 14 patients: 93.3% reported a general improvement in symptoms and 86.7% associated it with VE. Patients experienced a significant improvement in humidity (pre 88.2%, post 13.3%,  $p < 0.05$ ), otorrhoea (pre 100%, post 6.7%,  $p < 0.05$ ), earache (pre 41.2%, post 0%,  $p < 0.05$ ) and itching (pre 41.2%, post 6.7%,  $p < 0.05$ ), and 36.4% perceived an improvement in hearing loss ( $p > 0.05$ ). Only one AE (mild tingling) was recorded.

**Conclusion and relevance** Our observations showed that voriconazole ear drops were an effective and safe option that significantly reduced symptoms in patients with recurrent otomycosis which failed to respond to other therapeutic alternatives. Further prospective studies are needed to confirm these findings.

#### REFERENCES AND/OR ACKNOWLEDGEMENTS

No conflict of interest.

3PC-010

#### EVALUATION OF THE PRODUCTION ACCURACY AND ERROR RATE IN THE AUTOMATED COMPOUNDING OF CYTOTOXIC PREPARATIONS BY A ROBOT

<sup>1</sup>BR Kujau, <sup>2</sup>J Raffaelli\*, <sup>1</sup>C Klaas. <sup>1</sup>Universitätsklinikum Münster, Zentrale Einrichtung Apotheke, Münster, Germany; <sup>2</sup>Loccioni Deutschland GmbH, Humancare, Calw, Germany

10.1136/ejhp-2020-eahpconf.57

**Background and importance** In chemotherapy compounding, the accuracy of the preparation is related to patient safety. A fully automatic production through a robotic system should ensure not only complete documentation and minimisation of the risk of pharmacy personnel being exposed to toxic drugs, but also greater accuracy of the compounding, consequently improving patient safety.

**Aim and objectives** The study aimed to verify the production accuracy of APOTECaChemo as well as the error rate of the robot during compounding.

**Material and methods** Using the statistical software 'APOTECAm@A', which allows regular checking of the performance of the robot, the pharmacy production of 20 anticancer active ingredients was monitored from January to October 2018, focusing on the dosage accuracy (%) of the preparations automatically compounded and the robot error rate.

The results of the analysis will define the performance of the automation in terms of preparation quality and safety, and production efficiency in the daily routine of the pharmacy.

**Results** During the study period, 8478 automated preparations were compounded with APOTECaChemo by the pharmacy. The error rate of the robot was ~1% of the total automated production. Regarding the accuracy of the successful preparations compounded by APOTECaChemo, 97.5% of the preparations had a dosage accuracy between 0 and  $\pm 3\%$ . The remaining 2.5% of the preparations produced with the robotic system were within the  $\pm 5\%$  tolerance limits defined by the pharmacy as acceptable.

**Conclusion and relevance** The analysis carried out by APOTECAm@A showed high dosage accuracy in combination with a low percentage of errors in the automated production. The data show high quality as well as high reproducibility of safe production using APOTECaChemo.

#### REFERENCES AND/OR ACKNOWLEDGEMENTS

No conflict of interest.

3PC-011

#### CLOSED SYSTEM TRANSFER DEVICE BASED ON AIR FILTRATION: THE DRUG VAPOUR CHALLENGE

G Levin\*. *Simplivia Healthcare, Research and Development, Hod Hasharon, Israel*

10.1136/ejhp-2020-eahpconf.58

**Background and importance** Chemotherapy drugs were shown to form hazardous vapours that pose a health risk to pharmacists and nurses. One of the aims of using a closed system transfer device (CSTD) is to prevent this harmful exposure. The vapour containment efficiency of air filtration CSTDs is perceived as less obvious compared with that of physical barrier based CSTDs, and therefore should be proven throughout the shelf life of these devices in order to support the claims of its instruction for use (IFU).

**Aim and objectives** The aim of the study was to test the drug vapour containment capacity of Chemfort, a new air filtration CSTD. The objective was to investigate if the air filter remained fully functional at the end of the shelf life (3 years). According to the IFU, the device can be used on a drug vial for a period of 7 days, and thus the study also tested the filter functionality after it was exposed to vapours of a hazardous drug for 7 days.

**Material and methods** The study was performed by Nextar Labs (Nes Ziona, Israel). Vial adaptors (VA) were applied on drug vials (cyclophosphamide, 5-fluorouracil (5-FU)). Extreme conditions were used to generate vapours—heating to 50°C and having a nitrogen gas flow (250 mL/min) into the vial for 5 hours via the VA fluid pathway. A closed test chamber was employed for capturing drug vapours. Vapours released through the air filter were trapped, recovered and quantified using validated LC/MS/MS methods. As a positive control,