Purpose We aimed to describe the incidence of neuropsychiatric disorders in a cohort of HCV-infected patients treated with interferon and ribavirin, and their impact on treatment adherence and viral response rate (SVR).

Materials and Methods Data from a cohort of HCV patients who visited an outpatient pharmacy service (OPS) included all adult patients mono-infected with HCV who had completed treatment in 2010. Monitoring of neuropsychiatric disorders was assessed at weeks 0, 4, 12, 24, 48, and 72 through the self-administered questionnaires Hospital Anxiety and Depression Scale (HADS) and General Health Questionnaire (Goldberg). Adherence to treatment was assessed by counting drugs dispensed and patient reporting. Virological response was determined by the physician according to standard criteria.

Results Of the 76 patients included, 19 (25%) had a pre-existing psychiatric disorder, mostly depression and anxiety. The incidence of medically-confirmed neuropsychiatric disorders was 33% (n = 25), with a peak of abnormal results in the tests in week 12. Patients with and without pathological scores did not differ in baseline characteristics, except for pre-existing psychiatric disorder (60.0% vs. 7.8%, respectively (p < 0.001)). Antidepressants and/or anxiolytics were prescribed to 48% of patients with medically confirmed disorders (n = 12). Overall, 45% of patients achieved an SVR. Strict adherence (96% vs. 90%; p = NS) and SVR (59% vs. 52%; p = NS) were similar in patients with or without medically confirmed disorders.

Conclusions Patients often develop neuropsychiatric disorders during interferon therapy. Neuropsychiatric side effects had a non-significant effect on adherence to treatment and attainment of SVR. Multidisciplinary monitoring provided during the treatment of hepatitis C can contribute to early detection and management of neuropsychiatric disorders and to improve integrated patient care.

No conflict of interest.

Background Non-formulary drugs are prone to cause medication errors due to their less common use in the daily routine on the ward. Therefore non-formulary drug (NFD) management in the hospital pharmacy includes checking the dose and indication which is usually very time consuming. In 2010 the drug information centre had to deal with 12,903 prescriptions for NFDs.

Purpose Loss of relevant drug information at the interface between pharmacy and ward has been observed in some cases. Therefore a survey was performed to detect information gaps. Did the pharmacist’s recommendation reach the medical staff?

Materials and Methods During a period of four weeks all NFD prescriptions were documented concerning the type of medicine. If a treatment-relevant intervention (e.g. dose correction) was made the trainee pharmacist visited the ward to clarify if the pharmacist’s advice was received. In addition the medical staff were interviewed about the general procedure of information transfer within the ward staff.

Results 1158 NFDs were ordered. Out of these 261 required extensive action with pharmacist intervention. 256 interventions were accepted on the ward and only 5 were rejected. In only one case out of these the pharmacist’s information had to be resupplied to the ward as it had not reached the staff. The survey showed a very high acceptance (98.1%) of the drug information provided. 53 drugs within the ATC Code “antibiotics for systemic use” were particularly counselling-intense. Dosing problems were the most frequent drug-related problem (52). Information transfer within the ward turned out to be highly inhomogeneous.

Conclusions The pharmaceutical advice offered to the ward was accepted to a very high percentage. To prevent information loss on the ward a standardised system for information transfer amongst the staff needs to be established.

No conflict of interest.
carbohydrates (polynuclear iron (III)-oxohydroxy cores stabilised by carbohydrates), glutarimoids (polypeptides) and liposomal drugs [1]. Like biological MPs, NBCDs are complex MPs consisting of non-HEMO molecular, partially nanoparticle, structures. Composition, in vitro and in vivo characteristics are defined by manufacturing. Subtle changes of the manufacturing modify quality, efficacy and safety of the MP. NBCDs are not fully characterised physicochemically. In contrast to biosimilars, a regulatory framework is not established.

**Purpose** Intended copies of NBCDs such as the iron sucrose similars have been approved in several countries by the classical generic pathway. Growing scientific evidence in the non-clinical and clinical setting has raised doubts about interchangeability and/or substitutability.

**Material and Methods**

Science-based statements for comparability of intended copies and reference MPs were discussed among experts from regulatory science, clinicians, hospital pharmacists and industry in a Workshop at FIP 2012. The conclusions were used to propose regulatory requirements for NBCDs.

**Results** The FIP 2012 consensus meeting confirmed the lack of an appropriate regulatory market authorization of intended copies of NBCDs. For liposomes, physicochemical equivalence testing seems to be more likely to be achievable, but clinical efficacy trials are needed on a case-by-case basis (EMA). Nanoparticle iron sucrose similars show almost no comparability and therapeutic equivalence has to go through quality, efficacy and safety assessments [2]. Glutarimoids, with a not-understood mode of action, also need a broad, as yet to be defined, regulatory approach. Nanoparticle assessment includes sizing and morphology (FDA) and also evaluation of in vivo biodisposition (EMA). The upcoming Terminology and a White Paper will integrate these conclusions.

**Conclusions** For NBCDs and their specific characteristics a regulatory pathway is needed to assess comparability and eventually therapeutic equivalence of originator and intended copy MPs. In multiprofessional medicines management specific attention to the limits of interchangeability and substitutability is mandatory.

**References**


No conflict of interest.

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**Observational prospective study on pulmonary arterial hypertension and drug exposure**

*doi:10.1136/ejhp-2013-000276.125*

A Calinet, S Günter, A Rieutord, D Montani, MC Chaumais, Hôpital Antoine Béclère, Pharmacy, Clamart, France; Hôpital Bicêtre, Pneumology, Kremlin Bicêtre, France

**Background** Pulmonary arterial hypertension (PAH) is a rare disease characterised by an elevation of the pulmonary vascular resistance leading to right cardiac failure and death. Among different aetiologies of PAH, association with drug exposure was proved forty years ago with aminorex and more recently with benfluorex. Other drugs such as dasatinib or interferons seem to be associated with PAH development and/or severity. Pharmacovigilance is critical to improve our knowledge of PAH associated with drug exposure.

**Purpose** To confirm the feasibility of collecting the drug exposure history in PAH patients during hospitalisation by a systematic interview.

**Materials and Methods** This pilot study was performed in the French national PAH reference centre. Patients with idiopathic, heritable PAH, PAH known to be associated with drug exposure and pulmonary veno-occlusive disease were included. A standard questionnaire to collect the past and current medicines history was designed and approved by pharmacists and pneumologists. For each patient, this questionnaire was systematically assessed by a pharmacist after patient consent had been obtained.

**Results** Interviews were performed in 57 PAH patients. The median time of interview was 30 minutes. 16% of patients had a history of anorexigen exposure which led to 5 pharmacovigilance reports. The remaining four other patients were already known to the pharmacovigilance centre. Twenty seven patients (47%) had been exposed to psychoactive drugs, two patients to cytotoxic agents and one patient to interferon. Interestingly, a quarter of all patients had a history of nasal vasoconstrictor exposure.

**Conclusions** This pilot study demonstrates the feasibility of collecting the history of drug exposure in PAH patients during hospitalisation. Our observations match those reported in the literature except for the nasal vasoconstrictors, for which no epidemiological data have been published yet. Further studies are warranted to investigate the potential harmfulness of nasal vasoconstrictors.

No conflict of interest.

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**Off-label prescriptions in the neonatal intensive care unit at Marseille North Hospital**

*doi:10.1136/ejhp-2013-000276.126*

A E Fageur, M Desbourdes, N Colombini, V A Vial, M Bules-Charbit, Hôpital Nord, Pharmacy, Marseilles, France; Hôpital Nord, Pediatric ICU, Marseilles, France

**Background** The availability of drugs specifically assessed for use in neonates is limited as evaluation is more difficult in neonates than in adults. The result is a widespread off-label use of drugs, especially in neonatal intensive care units. Such practice is an essential part of their care and should be based on the best available evidence.

**Purpose** To describe and analyse the off-label use of medicines in a neonatal intensive care unit.

**Materials and Methods** Prospective observational study conducted over three months, from 27 February 2012 to 27 May 2012. All the drugs prescribed were analysed with regard to their licence status for the: indication, dose, route of administration, mode of administration, age category, formulation (compounding of capsules, oral suspensions, eye drops), contraindications and warnings specified in the summary of product characteristics of the marketing authorization.

**Results** In total, 638 prescriptions, comprising 59 different medicines were written, 107 newborn babies were admitted (60 male, 47 female). Their age varied from 0 to 27 days (average: 2 days), their mean gestational age was of 34 weeks of amenorrhea (65% premature), their weight ranged from 630 g to 4700 g (average: 2230 g). A total of 487 prescriptions were written off-label (76%), with 101 patients (94%) receiving at least one drug used off-label. Drugs were prescribed off-label mostly concerning the indication (48%), then came off-label use for the dose and the age category. The medicine most often prescribed off-label was caffeine citrate.

**Conclusions** Critically ill neonates are exposed to numerous medicines, a significant proportion of which are not yet approved for use in this vulnerable group of patients. Despite European initiatives aiming to promote greater awareness and research in the paediatric population, there is still a high percentage of unlicensed or off-label drug use in neonatal intensive care. This study underlines the need for clinical research and approval of the clinical data acquired within the neonatal population.

No conflict of interest.
GRP-124 Non-Biological Complex Drugs and Their Regulatory Approach – of Concern For Hospital Pharmacists and Medicines Management?

G Borchard, JB Rottembourg, B Flühmann and S Mühlebach

Eur J Hosp Pharm 2013 20: A44-A45
doi: 10.1136/ejpharm-2013-000276.124

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