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What Future Healthcare Professionals Need to Know About Pharmacovigilance: Introduction of the WHO PV Core Curriculum for University Teaching with Focus on Clinical Aspects

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Abstract  Adverse drug reactions (ADRs) can cause serious health problems, as shown in studies about drug-related hospitalizations. To build knowledge of and raise awareness about ADRs among healthcare professionals, more education in the field of ADRs and pharmacovigilance (PV) is needed. No standard exists for teaching PV at universities for medical, pharmacy, dentistry and nursing students, so a core curriculum needs to be developed to teach important aspects of PV to students. In September 2016, a stakeholders’ meeting was initiated on behalf of the World Health Organization (WHO) and organized by the Netherlands Pharmacovigilance Centre Lareb. This meeting addressed and agreed on the PV competencies students need to develop and what key aspects of the subject should be taught. Five key aspects were identified: understanding the importance of PV in the context of pharmacotherapy, and preventing, recognizing, managing and reporting adverse drug reactions. Since time and resources for PV education are limited, elements of the WHO PV core curriculum for university teaching were designed to be integrated into existing courses but can be used as a stand-alone programme. The basis of and outline for the WHO PV core curriculum for university teaching are addressed in this paper. It is expected that PV competencies for students are vital for their contribution to safe use of medicines in the future. In addition, this article aims to stimulate discussion on this subject and promote collaboration between universities, national PV centres and other stakeholders to integrate key aspects of PV in their educational programmes.

Key Points

The World Health Organization (WHO) pharmacovigilance (PV) core curriculum for university teaching focuses on clinical aspects and can be integrated into existing courses such as pharmacology and pharmacotherapy or used as a stand-alone course.

Key aspects of the WHO PV core curriculum for university teaching are understanding the importance of pharmacovigilance and preventing, recognizing, managing, and—finally—reporting adverse drug reactions.
1 Introduction

Medicines help us treat many diseases, but adverse drug reactions (ADRs) cause serious health problems. Studies indicate that ADRs account for approximately 5% of all acute hospitalizations [1–3]. About half of these hospitalizations could have been prevented by more effective drug therapy monitoring, timely recognition of the symptoms, and knowledge about certain risk factors for occurrence of these ADRs [1, 2]. From the patients’ perspective, all ADRs are important. All ADRs can pose a problem and may negatively influence quality of life and drug adherence, irrespective of seriousness and whether they arise from medication errors [4]. Sufficient knowledge of and skills around the safety of medicines and safe use of drugs in daily practice are important for all healthcare professionals (HCPs) who are directly involved in pharmacotherapy, such as doctors, pharmacists, dentists and nurses, to reduce drug-induced patient harm [5].

Competence in handling ADRs in clinical practice is important, not only for patient safety in individual patient care but also for drug safety monitoring at a population level. The science and activities relating to the detection, assessment, understanding and prevention of adverse effects or any other drug-related problem is called pharmacovigilance (PV) [6]. Although medical and pharmacy students may recognize the importance of ADR reporting and express the intention to report ADRs, they are insufficiently prepared to handle ADRs and have inadequate PV competencies [7–11]. This hampers optimal patient care and the safe use of drugs now and in the future [7]. For future HCPs to be competent in handling ADRs in clinical practice and reporting them, more education is needed [12].

HCPs often consider ADR reporting to be an additional activity rather than routine practice [13]. Beyond improvement of PV education at the university level, continuous educational interventions for practicing HCPs have proven to be effective in PV competencies [14]. Besides education, inter-professional collaboration improves the culture in healthcare in terms of recognizing ADRs and the awareness of ADR reporting [15].

To acknowledge the importance of safety issues in drug treatment, PV education should be introduced at an early stage, along with patient care as part of the Hippocratic Oath “first, do no harm”. Since students are accustomed to acquiring new knowledge and skills, it can be expected that PV education of students at university level will improve knowledge about safety of medicines and skills for a safer use of medicines at an early stage of their career [12, 16].

2 Development of a Practical Pharmacovigilance (PV) Curriculum for University Teaching

To teach important aspects of PV to undergraduate students, a core curriculum that describes desired competencies and learning outcomes and provides practical materials needs to be developed. Although patient safety, safe prescribing and PV are increasingly recognized as important topics in university educational frameworks worldwide, in many countries this education depends on individual professors or active PV centres [5, 12, 17–19].

Although there are initiatives to support education in the field of patient safety and PV in general, there is no standard programme about PV for university teaching. In 2011, the World Health Organization (WHO) established the WHO Patient Safety curriculum for undergraduate students. Although this curriculum contains medication errors and related adverse events, some aspects related to PV are still missing [20]. The WHO–International Society of Pharmacovigilance (ISoP) curriculum for PV teaching, designed for people interested in PV in any setting, consists of a comprehensive list of topics that cover almost all aspects of PV. Relevant topics for university teaching can be selected by users, but practical information about implementation of relevant sections of this curriculum and ready-to-use-teaching materials and an undergraduate course description should be developed [21, 22]. The European programme in Pharmacovigilance and Pharamcoepidemiology (EU2P) also offers education and training in pharmacovigilance for individuals with a special interest in PV [23]. For the broad audience of medical, pharmacy and nursing students in general, a manageable curriculum or programme is needed.

In September 2016, a stakeholder’s meeting was initiated on behalf of the WHO, under its Programme of International Drug Monitoring, and organized by The Netherlands Pharmacovigilance Centre Lareb WHO Collaborating Centre for Pharmacovigilance in Education and Patient Reporting. Meeting participants had backgrounds in medical or pharmacy teaching, PV and governmental positions and came from various continents (for details see the co-author section). This meeting addressed and agreed on the competencies in the field of PV students at university level need to develop and the key aspects that should be taught.

In this article, we outline the key aspects and competencies that form a PV core curriculum for university education. The key aspects of PV can be integrated in existing courses for medical, pharmacy, dentistry and nursing education. The curriculum can also be used as a stand-alone programme. To our knowledge, the WHO PV
core curriculum for university teaching is the first to be described in the literature.

### 3 Key Aspects of PV Education

PV entails various aspects of science and activities related to the safety and safe use of medicines throughout the entire lifecycle of a drug, from clinical research to post-authorisation phase. It is strongly associated with both regulatory aspects and clinical pharmacology. Since most students will work in clinical practice and provide patient-care services after graduation, our aim is to focus on the clinical aspects of PV and the use of authorised drugs as desired learning outcomes. The proposed key aspects, formulated as learning outcomes of undergraduate PV education, are as follows:

1. **Understanding the Importance of PV**
   - Drug-induced harm and hospital admissions
   - Historical examples

2. **Preventing ADRs**
   - General risk factors
   - Individual risk factors
   - Treatment guidelines and safety information

3. **Recognizing ADRs**
   - ADR classification
   - Risk factors
   - Confounding factors
   - Epidemiology

4. **Managing ADRs**
   - ADR classification
   - Seriousness
   - Severity

5. **Reporting ADRs**
   - Limitations of premarketing phase
   - Relevance of ADR reporting
   - Documentation of ADRs

The PV key aspects are summarized in Table 1 and explained in more detail in the following sections, see also Fig. 1.

### 3.1 Understanding the Importance of PV in the Context of Pharmacotherapy

Awareness of the impact of ADRs on individual patients is important for undergraduate students to understand patients’ perspectives on safety issues. The stories of patients who have experienced ADRs will elucidate the impact of ADRs to students. Both serious and non-serious ADRs may negatively influence patients’ quality of life and treatment satisfaction and hamper drug compliance [24].

Students also need knowledge of the extent of drug-induced harm at a population level, which can be visualized using drug-related hospital admissions [1, 2] and with historical examples of drug-induced disasters such as limb deformities following maternal use of thalidomide [25]. These examples gave rise to the present-day regulatory framework and the way in which the safety monitoring of drugs is organized.
3.2 Preventing Adverse Drug Reactions (ADRs)

Safe and rational prescribing and dispensing of drugs entails therapeutic reasoning and decision making, such as choosing the right drug for each individual patient. The occurrence of an ADR to previous treatment, as well as the individual patient’s risk factors should influence this choice. Students need to learn how to consider all circumstances that influence the occurrence of ADRs in a patient. This implies knowledge of general risk factors, such as the increased incidence of ADRs in the elderly, as well as individual risk factors related to a certain drug or to a patient’s medical condition, such as comorbidity and genetic factors. Drug–drug interactions can be expected when patients use concomitant medication, especially in polypharmacy [26], or when patients use herbal or other traditional medicines [27]. Food–drug interactions may influence drug absorption or drug metabolism; for example, grapefruit juice inhibits drugs metabolized by cytochrome P450 (CYP)-3A4 [28]. Information on these risk factors is generally available in drug labelling, treatment guidelines and medical literature, which students must learn to use and interpret.

3.3 Recognizing ADRs

It is important that students learn how to recognize symptoms of ADRs in patients. Identifying ADRs can be difficult, because sometimes they are hard to distinguish from underlying disease or comorbidity. This requires knowledge of clinical pharmacological principles of ADRs, such as types of ADRs, dose-relatedness, pharmacological actions, hypersensitivity reactions, time relationship and risk factors [29, 30]. Training in clinical reasoning with a focus on causality reasoning is important to distinguish a suspected ADR from other medical conditions or background incidence.

Observations of ADRs and designing individual treatment plans for managing ADRs will strongly contribute to the development of awareness of the impact of ADRs in patients and on the importance of PV at a population level.

3.4 Managing ADRs

Patients who experience an ADR need special and individual attention. The impact of ADRs ranges from absent or mild influence on quality of life to life-threatening situations, such as anaphylaxis or intracranial haemorrhages. Furthermore, ADRs may negatively influence a patient’s compliance with therapy. Therefore, depending on the indication, seriousness, perceived severity and prognosis, patients and HCPs must design an individual treatment plan. Actions for treatment include drug discontinuation, dose adjustment or additional treatment of the symptoms. Sometimes, alternative treatment for the initial disease should be chosen. After experiencing a severe ADR, patients might fear drug treatment in general [31]. Students need the skills and knowledge to classify an ADR and the competence to respond when they come across serious or severe ADRs, which can be difficult.

Communication skills are also needed to better explain safety data to colleagues and patients and encourage and understand feedback from those involved in PV [32].

Students should furthermore develop the ability to explain ADRs to patients and their families, advising about future drug use and the chances of the ADR recurring [33].

Finally, relevant elements of the ADR should be recorded properly into the patient’s healthcare record.

3.5 Reporting ADRs

Beyond the awareness of ADRs in individual patients, PV also focusses on the improvement of public health regarding the safety and safe use of medicines. Students need to understand that information on ADRs from the premarketing phase is limited. HCPs have a responsibility to share their clinical practice experience in the real world.
with regulatory authorities. Spontaneous reporting systems continue to be an effective method of signal detection [34]. ADR reports from HCPs and patients often trigger the detection of potential new medicines safety signals. They improve general information about ADRs. Reporting an ADR to PV centres or other relevant organisations is a skill that can be developed easily during clinical internships [4, 32].

### 4 Integration of PV Key Aspects in Existing Educational Programmes

#### 4.1 PV Competencies for Future Healthcare Professionals

Competency-based educational models are becoming more dominant than the traditional content-driven and teacher-centred learning methods in university teaching [36, 37]. A competency can be described as an individual’s ability to make deliberate choices for handling situations and tasks in specific contexts of professional practice [36]. In the context of pharmacotherapy, competencies in PV are the timely recognition, prevention and management of ADRs.

To acquire competencies, different learning outcomes with skills, knowledge and attitudes can be defined [35]. Competencies in PV were first described by Edwards et al. [32] in 2006. In the broad context of professional PV, healthcare professionals in clinical practice must ensure the safe use of drugs and contribute to the reporting of ADRs.

We summarized learning outcomes regarding these competencies as key PV aspects (Table 1). Students develop different levels of competence throughout their educational career, which require adjusted and aligned learning activities. Figure 2 illustrates how the related key PV aspects can be divided into levels of competence throughout university education. By offering key PV aspects with increasing complexity at different stages of education, students will acknowledge the importance of the subject and will be challenged and motivated to learn.

#### 4.2 Practical Aspects

The importance of PV can be taught from the first year of education, since limited specific knowledge is required. The clinical aspects of PV can be considered logical extensions of general subjects such as pharmacology and toxicology. The actual recognition, management and prevention of ADRs therefore requires broad knowledge and skills from pharmacology, pharmacokinetics, pathophysiology, pharmacotherapy, drug development, pharmaceutical regulations, epidemiology and general science. This implies the integration of key PV aspects becomes feasible later on in the Bachelor or Master phases, depending on the...
local structure of educational programmes. The actual reporting of an ADR needs a real clinical context, which is offered at the end of most educational programmes.

As demonstrated by Hartman et al. [12] and Schutte et al. [7], the time for education on PV appears to be limited. The amount of contact hours for explicit PV education in medical, pharmacy and nurse education was about 2–5 h in the entire curriculum [12]. The limited time and the applicability in clinical practice forces the PV core curriculum for university teaching to be confined to clinical aspects. The multidimensional character of PV education requires integration of key aspects in existing pharmacotherapy courses or methods such as prescribing safety assessments (PSA) [38]. In PSA, the evaluation of preventing, recognizing and managing ADRs should be incorporated as defined by the Guide to Good Prescribing [39].

4.3 PV in Educational Frameworks

Educational frameworks and medical codes of conduct provide a formal basis on which to integrate PV aspects into existing curricula [18]. For pharmacy and medical students, preventing and handling ADRs, as well as reporting ADRs to national PV centres, are listed in the frameworks [17, 40–43]. In Europe, a set of 252 learning outcomes for clinical pharmacology and therapeutics (CPT) education was published recently to harmonize medical education. The key PV aspects are covered in the description of the CPT learning outcomes [19].

Preventing, recognizing and managing ADRs and PV elements are lacking in educational frameworks for nurses everywhere, although patient safety in general is regarded as an important topic [44, 45]. Furthermore, it has been shown that nurses are thoroughly capable of recognizing and reporting ADRs [5, 46].

5 Examples of Undergraduate PV Education

The ways in which PV is taught differs according to country [12]. In some European countries, PV has been integrated at a high level in therapeutics courses or programmes such as PSA [18]. In developing countries, pharmacotherapy education has its own challenges, and PV is less well developed, both in general and in education [47]. A few examples of PV educational methods at universities are described in the literature and summarized in Table 2. Most of these examples show active learning elements, such as developing a reporting form, reporting an actual ADR, assessing causality of ADR reports, problem-based learning and group discussions. Active and contextual educational methods have proven to be effective ways of transferring knowledge [48]. However, education is currently almost exclusively aimed at ADR reporting. Studies have demonstrated that educational interventions for HCPs about PV positively influenced ADR reporting in the short term [14, 49–52]. There are no long-term studies of the PV competencies of university students after graduation.

| Table 2 Examples of methods in pharmacovigilance education at university level, ordered in type of students, educational phase and kinds of learning activities |
|---|---|---|---|---|
| Method | Students | Phase | Learning activities | Countries |
| ADR reporting | Pharmacy and medical | Master/internship | Collecting ADRs, questioning patients, Reporting ADRs | Turkey [16] USA [53] USA [54] Denmark [55] Iran [56] India [57] |
| PV course | Pharmacy | Bachelor 3rd year | Lectures, causality assessment, design reporting form, group discussion | Nepal [58] Malaysia [59] |
| ADR assessment | Medical | Bachelor/Master | ADR report analysis | Netherlands [60] |
| ADR problem solving | Medical | Bachelor | Problem solving | Netherlands [61] |
| Safe prescribing | Medical and nurse | 4th year | Interprofessional learning | USA [15] |
| Pharmacology course: risk perception of ADRs | Medical | Bachelor 3rd year | Clinical training | France [62] |
| PV in pharmacoepidemiology course | Biology and pharmacology | Not provided | Problem-based learning | Canada [63] |

ADR adverse drug reaction, PV pharmacovigilance
6 Conclusion and Future Steps

This paper provides an outline of the WHO PV core curriculum for university teaching, in which competencies and key aspects of PV for future HCPs have been formulated. In today’s busy clinical practice, drug-related problems such as the prevention, recognition and management of ADRs do not always get the attention they need. When students are competent at preventing, recognizing and managing ADRs, they can and will improve the safe use of drugs.

Since PV is important for all HCPs in clinical and public health disciplines, key PV aspects should be integrated into existing programmes and courses for medical, pharmacy, dentistry and nursing education. We emphasise that the key PV aspects are intended to be add-in elements. Therefore, we expect that integrating PV elements into existing courses requires a limited time investment.

As a first step, universities would need to recognize possible gaps in their educational programmes and provide opportunities for teachers to integrate elements of the PV core curriculum for university teaching. National PV centres or other PV stakeholders can stimulate and facilitate PV education for students. For example, they can provide ‘teach the teacher’ courses to train university teachers about PV. Second, in countries where reporting ADRs is a challenge, PV education at a university level can take place in parallel with improving the reporting system.

Further discussions are needed to develop materials and share experiences to integrate key PV aspects in university teaching. A second stakeholder meeting was held in 2017 to discuss materials, teaching methods and strategy for sharing and implementation. Participants from the second meeting came from various parts of the world and are actively involved in PV education at universities, as PV staff or as university teachers. The main outcomes were to develop a web portal for sharing materials and to construct a ‘teach the teacher’ programme to promote PV education to university staff who are less familiar with PV.

To meet PV competencies in clinical practice, a cultural change is required in healthcare institutions in which PV is not yet common practice. The number of pharmaceutical products is increasing, as is the number of patients across the world with access to these treatments. The central role for PV, aiming at the safety and safe use of medicines, is fundamental for future HCPs. By developing PV competencies at universities, we may take the first step in this journey of change.

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Compliance with ethical standards

Conflicts of interest Rike van Eekeren, Lea`n Rolfes, Andries Koster, Lara Magro, Gurumurthy Parthasarathi, Hussain al Ramimmy, Tim Schutte, Daisuke Tanaka, Eugène van Puijenbroek and Linda Härmark have no conflicts of interest that are directly relevant to the content of this study.

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