Towards a “prescribing license” for medical students: development and quality evaluation of an assessment for safe prescribing

David R. M. Jansen1 • Carolina J. P. W Keijsers2 • Michiel O. Kornelissen3 • Marcel G. M. Olde Rikkert1 • Cornelis Kramers4 • (on behalf of the education working group of the Dutch Society for Clinical Pharmacology and Biopharmacy)

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Abstract

Purpose This report describes the development and validation process of an assessment with national consensus in appropriate and safe pharmacotherapy.
Methods A question-database on safe prescription based on literature of pharmacotherapy-related harm was developed by an expert group from Dutch medical faculties. Final-year medical students concluded a 2-year education program on appropriate and safe prescription by one of nine assessment variants of 40 multiple-choice questions each. An expert panel of professionals (n = 10) answered all database questions and rated questions on relevance. Questions were selected for revision based on lack of relevance or poor test and item characteristics.
Results A total of 576 final-year medical students of the Radboud University was assessed. There was no significant difference in performance between students and content expert group (p = 0.7), probably due to learning behavior. Out of 165 questions, 59 were selected for revision.
Conclusion Joint national effort from a team of experts in prescription and pharmacotherapy is an appropriate way to achieve a valid and reliable last-year student drug prescription assessment.

Keywords Prescribing • Medication safety • Education

Introduction

Teaching junior doctors to be prepared to write appropriate and safe prescriptions requires integration of knowledge on drug selection, dosage, side effects, interactions with other drugs, co-morbidities, and individual (genetic) variability in pharmacokinetics. Many studies report a high rate of inappropriate prescribing as a potential cause of preventable morbidity and mortality [1, 2]. In this light, it is relevant to note that for the hospital setting, most drugs are prescribed by junior doctors with limited clinical experience [3]. For example, in the United Kingdom, half of the prescriptions were made by foundation Y1 doctors, so the doctors with the least experience [3]. Although the cause of a prescribing error is probably multifactorial, and related to for example time pressure or interruptions, it is also shown that junior doctors tend to copy the drug treatment choices of their supervisors instead of basing their choices on their own independent analysis of the [4]. One way or the other, it is prerequisite that junior doctors are well prepared for the high-risk task of prescribing, before
entering the ward. Therefore, sufficient education on appropriate and safe prescribing is conditional and an assessment on whether this education had led to sufficient knowledge could be recommended. On top of that, it is shown that assessments drive the learning of the students [5]. Unfortunately, appropriate learning outcomes for such an education program are lacking in many (European) countries [6, 7].

It is known that the majority of adverse events are caused by a relatively small group of drugs, i.e., pain medication (non-steroidal anti-inflammatory drugs (NSAIDs) and opioids), (combination of) antithrombotics, antibiotics, cardiovascular agents, and drugs which are renally excreted [8–12]. Experts from seven out of eight Dutch University Medical Centers, brought together via the Dutch Society for Clinical Pharmacology and Biopharmacy (further referred to as ‘expert working group’), reached consensus on the learning goals as shown in Table 1, based on the drugs causing the most drug-related harm in clinical practice.

For each of these drugs, the expert working group decided specifically what learning goals should be mastered to ensure appropriate and safe prescribing. These included mechanism of action, the side effects that may harm the patient, risk factors for these side effects (both patient factors and co-medication), measures to prevent these side effects, and how to act if these side effects occur. Since drug-related harm was the starting point, the learning goals concern mainly safety. However, the most important indications of the drug causing drug-related harm had to be mastered. For antibacterial drugs, the learning goals mainly cover efficacy and not safety. The expert working group has taken the initiative to develop an educational program in pharmacotherapeutics which is followed by a formal assessment of students’ comprehension and application of knowledge on safe prescribing. On the Blooms taxonomy on cognitive domains, learning domains ranging from recall of information to applying and synthesis of information should be incorporated in this educational program and assessment on safe pharmacotherapy [13].

The educational program consists of a reader with literature review, practice assessments, and interactive lectures, which can be seen at https://www.youtube.com/user/fteindtoets/ (in Dutch). For better understanding of the scope of the reader, the learning goals for the domain of anticoagulants is added as an e-Appendix.

After the educational program, students could obtain a “medical prescribing license” by passing the formal assessment [14], which is part of the final-year medical examination program. By passing this assessment, the students demonstrate a certain basic level of mandatory knowledge for appropriate and safe prescribing analogous to the driving license theory test.

It is the ultimate vision of the expert working group to develop a compulsory assessment on all medical faculties in the Netherlands, so that medical students cannot graduate if they do not control both the knowledge and the skills needed for appropriate and safe drug prescription. Therefore, we describe in this paper the development and validation of a formal assessment in appropriate and safe pharmacotherapy by joint national effort and report its external validation process by using an independent group of experts for assessing content validity (further referred to as “content expert panel”).

## Methods

This study was designed to develop and study the quality of a newly developed 40 questions assessment on pharmacotherapy knowledge for last-year medical students.

### Development of the assessment: content

First, national consensus was achieved on the knowledge base required for the medical students by an expert working group (n = 7), consisting of representatives from seven out of eight Dutch medical faculties [14]. All of the members of the expert

| Table 1 Consensus-based domains for pharmacology training in last-year medical students across The Netherlands |
|---------------------------------|---------------------------------|
| **Domains**                     | **Subjects per drug-related group** |
| Drug-related topics             |                                  |
| Analgesics                      | 1. Medications and mechanisms of action |
| Anticoagulants                  | 2. Main indications               |
| Cardiovascular agents           | 3. Relevant kinetic data          |
| Drugs for diabetes mellitus     | 4. Main problems or side effects: including mechanism of action and main clinical presentation per agent |
| Antidepressants                 | 5. Patients who are most at risk per problem or side effect |
| Benzodiazepines                 | 6. Key interactions and mechanisms contributing to problem or side effect |
| Antibiotics                     | 7. Measures to prevent problems or side effects |
| General knowledge               | 8. Measures to take if a problem arises or side effect occurs |
| Basic pharmacokinetics          |                                  |
| Drug allergies                  |                                  |
| Laws and regulations            |                                  |
| Appropriate drug use            |                                  |

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working group were physician or pharmacist, all registered clinical pharmacologists.

In order to design the curriculum to prepare the students for applying the underlying knowledge in this assessment, a stepped approach was chosen: (1) identify the drug groups which cause most harm, (2) identify the learning goals per group, and (3) identify additional important knowledge to be assessed.

For the first step, the expert working group identified the drug groups known to cause the majority of preventable serious adverse reaction based on the Dutch HARM-Wrestling Task Force targeting outpatient drug safety [15]. The following drug groups were identified and clustered by indication into seven drug-related domains, namely, (1) analgesics, (2) anticoagulants, (3) cardiovascular agents, (4) drugs for diabetes mellitus, (5) antidepressants, and (6) benzodiazepines. By consensus, the expert working group added (7) antibiotics as domain. Antibiotics are widely used and, in the opinion of the expert working group, knowledge about good antibiotic practice is relevant because injudicious use may not only cause harm in individual patients but may also induce antimicrobial resistance at the societal level.

Second, the expert working group systematically assessed the following categories to identify learning goals connected to these drug groups: (1) medications and mechanisms of action, (2) main indications, (3) relevant kinetic data, (4) main problems or side effects: including mechanism of action and main clinical presentation per agent, (5) patients who are most at risk per problem or side effect, (6) key interactions and mechanisms contributing to problem or side effect, (7) measures to prevent problems or side effects, and (8) measures to take if a problem arises or side effect occurs.

Additionally, general knowledge domains concerning related general pharmacotherapy were defined by the expert working group, based on expert opinion and added to the previously defined seven drug groups, i.e., (1) basic pharmacokinetics including simple dosage calculations; (2) drug allergies; (3) laws and regulations including medicine and driving, and drug prescribing; and (4) appropriate drug use consisting of topics like the WHO-six-step method [16], pharmacotherapy in case of pregnancy, and lactation and transfer of information to pharmacist and primary care physician.

For the final step, Table 1 shows the domains and subjects per drug group as a result of expert consensus. Table 2 shows the test matrix of the assessment based on these domains. After three steps of content development, a total of 11 assessment domains were identified by expert consensus: core prescription knowledge on seven drug groups and on four related general pharmacotherapy domains. For details about drugs to be studied by the students per domain, see e-Appendix.

Development of the assessment: from questions to assessment

The expert working group created a database of multiple-choice questions covering basic and applied knowledge of the selected drugs and general topics. All experts contributed questions which were subsequently reviewed by experts from two other medical faculties. Second, all questions were appraised by an assessment expert which ultimately resulted in a database with 165 peer-reviewed and approved questions.

Next, from this database, assessments were drawn consisting of 40 questions each, covering the previously mentioned 11 domains with 2 to 6 questions per domain. In Table 1, the number of questions per category in each assessment is shown. These assessments were checked for duplicate questions and contamination between the questions, e.g., if one question could be helpful to answer another question. If needed, small adjustments were made to the assessment and questions were replaced by other questions. From the database, nine different assessments were drawn, which were alternately used for the test taking.

Patients and data collection procedure

In the period of March 2015 to December 2016, the assessment took place monthly in groups of on average 30 final-year medical students from the Radboud University Faculty of Medical Sciences. In total, 576 students were assessed, during 1-h sessions. The assessments consisted of 40 multiple-choice questions per assessment with on average four (range: two to five) alternatives resulting in at least 150 answer options per assessment, which is considered as lower limit for reliable psychometric analysis.

The assessment was summative as part of their regular curriculum; students had to pass it to become a physician. Those who failed the assessment had to repeat the assessment until they acquired a sufficient score. The pass rate was set at 80% correct answers after correction for guessing.

To study concurrent validity, a content expert panel, consisting of physicians with regular prescribing activities
and pharmacists, all registered clinical pharmacologists, was composed. The experts were approached by e-mail in October 2016 with the request to participate. The goal of this content expert panel was twofold: first to verify whether these experts considered the questions a precondition to be able to perform safe pharmacotherapy in clinical practice and second for external validation by comparison of the students’ results with the experts’ scores. The content experts examined the full set of 165 questions. All questions were answered and rated on relevance by choosing “essential,” “useful, but not essential,” or “not necessary” according to Lawshe [17]. Additionally, there was an option for the expert to note comments per question. The mean values per assessment for the experts were reduced from the total set of 165 questions.

The answers of the students, the answers of the experts, and experts’ relevance scores and comments were anonymously collected and analyzed.

**Data analyses**

**Construct validity**

In order to study construct validity of the assessment as a whole, the following analyses were performed:

Concurrent validity was studied by comparing the means of the medical students with the experts from the content expert panel by unpaired t tests for the overall comparison.

Content validity was tested according to Lawshe [17]; all experts rated all questions as “relevant” (value 1) or either one of the other two options: useful, but not essential, or not necessary, and in that case scored the item as “not relevant”: value −1. If the mean score was positive, an item was scored as “precondition to be able to perform safe pharmacotherapy”. Content validity was tested with a focus on the most frequent occurring drug-related problems, by which the larger part of safe prescribing was tested.

Criteria for content validity, discriminant analysis and test characteristics were combined to evaluate the appropriateness of all items separately (see below).

**Reliability**

Different aspects of reliability were analyzed, namely, the internal consistence of the assessment as a whole and several parameters per question. To start with, the internal consistency was measured by Cronbach’s alpha [18] per assessment, which should be 0.50 or higher, as proposed for multidimensional assessments and relatively short test lengths [19, 20]. Internal consistency was only analyzed in the student group because the internal consistency differs per test population, and the students were the population aimed for. Next, for all questions, discriminant analysis was performed per item calculating the “Difficulty Index” with correction for guessing on multiple choice; probability (p′), reflecting the percentage of students who answered the item correctly, including correction for guessing (p′ = p − (1 − p) / (number of options − 1)). The scores were interpreted as follows: > 0.9: easy question, 0.9 to 0.5: medium difficulty, < 0.50: difficult question for three-option multiple-choice questions, and for four-option multiple-choice questions: > 0.9, between 0.9 and 0.44, and < 0.44, respectively to correct for chance. Next, item-rest correlation (Rir) was calculated, which is the correlation between the question score and the overall assessment score (excluding that question), referring to how well a question differentiates between participants who master the material and those that do not know [21]. Rir scores were categorized as: < 0.19: poor, 0.20–0.29: adequate, 0.30–0.39: good, and > 0.40: very good) [22, 23]. Standard deviation was calculated for all assessments for students and content expert panel as a group.

For adequate interpretation of the quality of the questions, each question should preferably at least be used 100 times [23]. To accomplish this, we reduced the amount of assessment variants to the first four (of a total of nine). The other test variants were used less often, and so, for the questions in these variants, this criterion was not met.

**Selection procedure of potentially inappropriate items**

Finally, potentially inappropriate questions were selected for revision. A question was considered potentially inappropriate if at least one of the following criteria was met: (1) difficulty index (p′ value) < 0.8 for experts, (2) rated “irrelevant” by a majority of the content expert panel, or (3) p value < 0.65 for students and item-rest-correlation ≤ 0.

All assessments were entered in the online assessment tool TestVision (Teelen Kennismanagement, The Netherlands). Psychometric analysis was performed with Microsoft Excel 2007 (Microsoft) and IBM SPSS 22.0 (IBM). Statistical significance was set at p < 0.05.

**Results**

From March 2015 to December 2016, 576 unique students (66% female) were assessed, as well as 10 independent experts from the content expert panel (40% female). The students completed a total of 673 assessments.

Table 3 shows the main results. The vast majority (87.3%) of the students passed the assessment at the first attempt. A total of 73 students (12.7%) had to retake the assessment (8.9% once, 3.5% twice). The mean score of the students was 90.5% and of the experts 90.7%.
Validity

Three domains of validity were studied. First, the experts were compared with the students for concurrent validity. There was no significant difference between the students and the experts (mean score 36.18 out of 40, (SD 2.97) versus mean 36.29 (SD 2.03) (t(761) = −0.352, p = 0.73). Overall, the mean rating of the questions was positive meaning “essential” by a majority of the experts. Still, 24.4% (n = 40) of the questions were considered “useful, but not essential,” or “not necessary.” All these 40 questions were marked as potentially inappropriate.

Reliability

First, the analysis of complete assessments was considered. Internal consistence as measured by the Cronbach alpha (a measure of internal consistency) how closely related a set of items are as a group. It is considered to be a measure of scale reliability ranged from 0.54 to 0.77 and was in the range 0.5 or above per test, which is adequate to good.

Next, discriminant analyses on the level of the individual questions were performed. The majority of the questions (87.9% (n = 145)) turned out to be easy (p’ > 0.90) based on students results. Ten questions (6.1%) were medium and five items (3.0%) turned out to be a difficult question. The item-rest correlation was very good for 13.9% (n = 23) of the questions, good for 16.4% (n = 27) of the questions, adequate for 23.6% (n = 39) of the questions, and poor for 43.0% (n = 71) of the questions.

Qualitative analysis of potentially inappropriate questions

Figure 1 shows which questions were included for what reason in further adjustments based on potential inappropriateness. A total of 59 out of 165 questions (36%) were marked for revision based on at least one of the three defined criteria; 38 questions based on difficulty for the experts (p’ < 0.8), 40 questions based on lack of relevancy as rated by the majority of the experts, and seven questions based on student criterion. The vast majority of the questions was marked for revision based on the first two criteria. The third criterion only added one unique question to the set.

As a result of this discriminant analysis, all marked items were revised for future use by the expert working group.

Discussion

This report describes the development of an assessment on pharmacotherapy knowledge for medical students. Based on...
literature on frequently occurring drug-related problems, learning goals were set by a nationwide expert working group, resulting in 11 domains. Using a database of 165 questions, assessments consisting of 40 questions were tested in over 500 students and 10 content experts. It appeared that the overall performance of last-year medical students on this assessment on appropriate and safe prescription was comparable with the performance of experts in the field of appropriate and safe pharmacotherapy. The assessment had an acceptable internal consistence shown by a mean Cronbach alpha of 0.66. In the analyses of the individual questions by the experts of the content expert panel, still, many questions should be considered for revision namely 35.8%, with rather large overlap between the questions as considered non-relevant by the experts and those with low classic psychometric parameters based on expert data. This is an example of a stepwise approach of the development of a prescribing assessment.

A relevant finding is the fact that students score near the level of experts. Generally, summative assessments are designed to discriminate between those who do have enough knowledge and those who do not. The caesura, or cut of point to pass, is adjusted depending on the mean score of the subjects tested to assure a weighted distribution of scores independent of the difficulty of the questions. In other words, a student has to be in the upper half of scores to pass. For this assessment, however, the national expert working group set a fixed relatively high caesura of 80%, to be sure the students demonstrated a basic standard of competence in safe pharmacotherapy, in analogy with the driving license theory test. Consequently, a fair amount of 87.9% of the questions was considered easy as measured by the easiness index and moderate item-rest correlation index. Also, the high caesura might have stimulated the external motivation to study hard on this topic, with high assessment scores as a result [24]. However, whether and how these knowledge levels may again start to differ over time during a medical career remains to be studied [25].

More initiatives are taken worldwide to improve education on safe pharmacotherapy and to assess pharmacotherapy knowledge or skills [26]. For example, recently, there was a report about a nationwide initiative for an online prescribing
assessment for final-year medical students in the UK, published by Maxwell et al. [27]. However, a widely accepted consensus is lacking on (1) the optimal strategy in education of future prescribers to reduce prescription errors in primary care and hospitals and (2) appropriate learning goals for this education program and concluding assessment. This study shows how a national collaboration can lead to a nationally accepted assessment, which will gradually be introduced in all medical faculties in the Netherlands.

Despite consensus in the expert working group who initially developed the assessment, it was surprising that there was a lack of consensus between the experts that designed the assessment and the content expert panel as used in this study for external reference. This content expert panel rated 24.2% of the individual questions as non-relevant though these questions were developed by expert colleagues. Since prescribing is a complex skill, this often results in more than one adequate prescription and in more than one correct answer for a clinical problem, depending on the circumstances. This lack of concordance between the expert working group and the content expert panel is a clear indication that context affects assessment.

The assessment in this study shows to be a valid and reliable assessment on pharmacotherapy knowledge for medical students. However, before nationwide application, some considerations should be taken into account. One could discuss whether the Cronbach’s alpha is high enough. Although it is certainly acceptable, for an assessment with such a high caesura of 80%, a higher alpha may be required. The moderate alpha level may be explained by the fact that these assessments had (1) many knowledge domains, (2) relative limited number of items per assessment, and (3) a rather uniform performance of the group or ceiling effect due to learning behavior [20]. All three aspects however were chosen as a part of the design and therefore higher Cronbach’s alpha will probably not be obtained as long as students intensively study for their assessment, which is aimed for. An alternative would be to extend the assessment by adding more questions [19]. Since the end of data inclusion for this study, the assessment is extended to 60 questions per assessment. Regarding the item-analyses, it would have been preferable to have had more assessments in the analysis. The student numbers however were set, and within the time frame of the study, more assessments were not possible.

It should be noted that the development of this assessment is the result of a nationwide Dutch collaboration. However, because of differences in laws, regulations, and guidelines, learning goals are not one-on-one transferable to other countries. Nevertheless, we believe that the procedure described here, with joint national effort from a team of experts in the field, may be feasible for implementation in other countries.

Conclusions

This study describes a valid and reliable assessment, developed by national effort to assess final-year medical students’ comprehension and application of knowledge on appropriate and safe prescribing. With a high cut-off point to pass, students seem to be very motivated to pass. After an education program on appropriate and safe prescribing, students achieved a score comparable to experts in the field. In analogy of the “driving licence theory test” an addition practical test “on the road” should be obligate: this should assess whether students can apply clinical pharmacology knowledge in clinical practice. Whether such an assessment ultimately improves safe prescription and overall patients’ outcomes has yet to be investigated.

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Author contributions DJ, CKe, MK, MO, and CKr wrote the manuscript; DJ and CKr designed the research; DJ and MK analyzed the data.

Compliance with ethical standards

The Dutch Ethics Review Board of Medical Education (NVMO-ERB) approved conduct of this study.

Conflict of interest The authors declare that there are no conflicts of interest.

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