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A European approach to categorizing medicines for fitness to drive: outcomes of the DRUID project

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Keywords
accidents, automobile driving, drug prescriptions, drug utilization, risk assessment, traffic

WHAT IS ALREADY KNOWN ABOUT THIS SUBJECT

• Some commonly prescribed medications can be a hazardous to traffic safety.
• Fifteen categorization systems are currently available in Europe. However, none of these systems clearly reports the methodology that was followed in order to categorize medications that impair driving.
• None of the existing categorization systems are currently implemented at European level.

WHAT THIS STUDY ADDS

• This study describes standardized and harmonized criteria to categorize medications according to their potential to impair fitness to drive.
• This study proposes a European categorization system of medications that impair driving that covers all the most frequently prescribed medications.
• The proposed categorization system can be seen as a tool to improve prescribing and dispensing procedures of medications that impair driving as well as an instrument to make patients aware of the role medications play in traffic safety.

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AIMS

To illustrate (i) the criteria and the development of the DRUID categorization system, (ii) the number of medicines that have currently been categorized, (iii) the added value of the DRUID categorization system and (iv) the next steps in the implementation of the DRUID system.

METHODS

The development of the DRUID categorization system was based on several criteria. The following steps were considered: (i) conditions of use of the medicine, (ii) pharmacodynamic and pharmacokinetic data, (iii) pharmacovigilance data, including prevalence of undesirable effects, (iv) experimental and epidemiological data, (v) additional data derived from the patient information leaflet, existing categorization systems and (vi) final categorization. DRUID proposed four tiered categories for medicines and driving.

RESULTS

In total, 3054 medicines were reviewed and over 1541 medicines were categorized (the rest were no longer on the EU market). Nearly half of the 1541 medicines were categorized 0 (no or negligible influence on fitness to drive), about 26% were placed in category I (minor influence on fitness to drive) and 17% were categorized as II or III (moderate or severe influence on fitness to drive).

CONCLUSIONS

The current DRUID categorization system established and defined standardized and harmonized criteria to categorize commonly used medications, based on their influence on fitness to drive. Further efforts are needed to implement the DRUID categorization system at a European level and further activities should be undertaken in order to reinforce the awareness of health care professionals and patients on the effects of medicines on fitness to drive.
Introduction

Driving a motor vehicle is a multifaceted task and it requires appropriate cognitive and psychomotor skills (e.g. alertness, concentration, reaction time, visual acuity) [1–3]. Medication can adversely affect these driving-related skills, and, consequently, be a hazard to traffic safety [4, 5].

The European Council Directive 83/570/EEC of October 1983 established that the summary of product characteristics (SmPC) has to contain information on medicines’ effects on the ability to drive and to use machines [6]. In October 1991 the European Committee for Medicinal Products for Human Use (CHMP) provided a Note for Guidance for the SmPC in which it was stated that section 4.7 of medications registered from 1 January 1992 had to indicate, on the basis of the pharmacodynamic profile, reported adverse drug reactions (ADRs) and/or impairment of driving performance or performance related to driving based on three different levels of impairment with respect to the ability to drive and/or operate machines [7, 8]. However, this rule has never been implemented [9].

In September 2009, a new SmPC guideline was issued, which established that ‘on the basis of the pharmacodynamic and pharmacokinetic profile, reported adverse reactions and/or specific studies in a relevant target population addressing the performance related to driving and road safety or using machines, specify whether the medicinal product has: (i) no or negligible influence, (ii) minor influence, (iii) moderate influence or (iv) major influence on these abilities’ [10]. These new guidelines were partly based on the proposal sent to the European Medicines Agency (EMA) by DRUID Work Package (WP) 4 partners during the consultation phase for the revision of the SmPC guidelines, in March 2008.

Despite the above-mentioned regulations, at this moment, a European categorization system has not yet been established, and warning systems for medications that potentially impair driving have mainly been developed and/or implemented at national levels [8, 11].

Existing categorization systems on medicines and driving

A review of the existing classification/categorization and labelling systems for medicines and driving was performed in 2008 and 15 different approaches were identified [12]. The categorization/labelling systems differed significantly and were not standardized, making them difficult to understand. In most cases [13–17], the categorization systems were developed by different and unrelated bodies, societies or researchers and were, in general, aimed at improving the prescription and dispensing of medicines to the patients and drivers. The identified categorization systems often included a limited number of medicines belonging to a few different therapeutic groups (e.g. antihistamines, anxiolytics, etc) and were not legally binding. However, the review also identified a couple of categorization/labelling systems that were developed by regulatory bodies, included the use of pictograms and were legally binding [18, 19].

In 1973, the Netherlands became the first country to introduce a list of medications that can impair driving abilities. Besides the list, the use of a yellow warning sticker on medication boxes was established and implemented [20]. In 1981, Denmark, Finland, Iceland, Norway and Sweden adopted a warning label. The label consisted of a red triangle printed on packages of ‘especially dangerous’ medications, and it is currently still in use in Denmark, Finland, and Norway. Most recently, France [18] and Spain [19] developed a categorization/labelling of all available medicines using technical interdisciplinary groups formed from their respective national medicines regulatory agency [21, 22]. The introduction of pictograms (three-tier labelling system in France and two-tier in Spain) to be added on the packages of certain medicines became legally binding in both countries.

It is important to point out that although different categorization systems are currently available across Europe, the criteria for the establishment of a categorization system for potentially impairing medications has neither been clearly described nor published nor been officially adopted at European level [12].

The DRUID project and its categorization system on medicines and driving

The Driving under the Influence of Drugs, Alcohol and Medicines (DRUID) project is an integrated project funded by the European Commission. The main aim of DRUID is to give scientific support to European Union (EU) transport policy by establishing guidelines and measures that combat impaired driving [23].

The DRUID WP4 aims to provide the basis and the methodology for the development of a European classification/labelling system for medications with respect to their impact on fitness to drive. Furthermore, it also focuses on the development of a classification of relevant therapeutic groups that are currently on the market in Europe as well as new medications approved by the European Medicines Agency (EMA) in the years 2007–2009 [23].

Aims of the study

This paper illustrates: (i) the criteria and the development of the DRUID categorization system, (ii) the number of medicines that have currently been categorized and the distribution of the DRUID categories across the Anatomical Therapeutic Chemical (ATC) index, (iii) the importance of this system, its implications for health care professionals (HCPs) and patients, and its strengths and limitations and (iv) the next steps in the implementation of the DRUID system and some general recommendations.
Methods

The development of the DRUID categorization system was based on the criteria that were established by a group of experts in the field of medicines and driving, involved in the DRUID WP4, and based on their consensus [24].

The four DRUID categories on medicines and driving

In 2006, the DRUID group established and agreed that, according to its influence on fitness to drive, a medicine could be categorized as follows (Figure 1):

- **Category 0** (no or negligible influence on fitness to drive),
- **Category I** (minor influence on fitness to drive),
- **Category II** (moderate influence on fitness to drive),
- **Category III** (severe influence on fitness to drive).

The proposed categorization is in line with the recently approved SmPC guidelines, which were adopted in September 2009 by the EMA [10].

Furthermore, the DRUID experts decided to develop, for each category, practical information to be used by HCPs for patient counselling purposes as well as simple warning labels that could be easily understood by patients (labeling) (Figure 1).

The DRUID categorization of medicines and driving

The ATC classification list [25] was used as a starting point for the selection of the relevant groups of medicines to be categorized. The aim was to categorize all available medicines on the European Union market for each selected ATC group.

Figure 2 shows the process that was followed in order to identify all those medications that are currently available on the EU market. In general, a medicine was considered available on the EU market if it was commercialized in at least two of the following European countries: Belgium, France, Germany, Greece, the Netherlands, Spain, United Kingdom and Ireland. If the above-mentioned criterion was not fulfilled, the medication was not included in the categorization process.

After a meeting with the French Health Products Safety Agency (AFSSAPS) experts in categorizing medications affecting driving performance, the DRUID WP4 group...
decided to adopt a procedure similar to the one used in France and more specifically to evaluate the following information and data:

1 Conditions of use of the medicine in the EU market
2 Pharmacodynamic and pharmacokinetic data
3 Pharmacovigilance data (including prevalence of undesirable effects reported in the SmPC)
4 Experimental and epidemiological data
5 Additional data derived from the patient information leaflet (PIL) and existing categorization systems and information from other sources
6 Synthesis and final categorization.

Figures 3 and 4 summarize the methodology that was followed in order to assign a category to a selected medicine.

The conditions of use of the medicine, pharmacodynamic, pharmacokinetic and pharmacovigilance data (including prevalence of undesirable effects) were derived from the SmPC [10], whereas point 4 (experimental and epidemiological data) was based on a scientific literature search.

The SmPC and PIL of the selected medications were found online, in one of the following websites: Medicines and Healthcare products Regulatory Agency (MHRA) [26], Electronic Medicines Compendium (eMC) [27], or Irish Medicines Board (IMB) [28], or retrieved from national medicines regulatory agencies as needed. In case of recently approved active substances, the SmPC was found on the EMA website [29]. The selection of the above mentioned medicines regulatory affairs agencies was simply based on the fact that the required information had to be available.
available either in English or in a language that could be fully understood by DRUID WP4 partners.

Specific sections of the SmPC and PIL were used to retrieve details on the active substance presentations and strength, indications, posology, route of administration (step 1), pharmacodynamic and pharmacokinetic profile (step 2), effects on the ability to drive and use machines (step 5) and undesirable effects related to driving and operating machines (step 3).

With respect to the undesirable effects, their occurrence was considered as a key point, especially if experimental and epidemiological data were lacking or limited. This type of information was found in section 4.8 of the SmPC and, when not available, was retrieved from the available literature.

Generally speaking, only those adverse reactions that could affect the ability to drive and that were reported as common (>1/100, <1/10) or very common (>1/100) were considered to be relevant, as in accordance with the most recent EMA categorization on frequency of undesirable effects, side effects or adverse reactions. In cases of rare or very rare undesirable effects, or if certain severely impairing effects occur, for example sudden sleep attacks, the DRUID partners recommended that this should be mentioned in the PIL.

Table 1 reports the criteria used for assigning a medicine to a specific category whenever experimental or epidemiological data were lacking or limited.

Table 2 lists the undesirable effects that could impair the ability to drive, and, therefore, were taken into account in the categorization process.

Data sources for the scientific literature evaluation included the electronic databases Medline, Science Direct and PsycINFO. The search was performed by using these combinations of keywords: ‘active substance name and psychomotor performance’, ‘active substance name and automobile driving’ and ‘active substance name and traffic accidents’. The final data selection was limited to full text articles published in English and other languages that included references to side effects, experimental and pharmacoepidemiological studies and case reports on each active substance to be categorized and its possible driving impairment. No restrictions concerning the publication year were applied.

Additional steps consisted of reviewing section 4.7 of the SmPC ‘Effects on ability to drive and use machines’ and the PIL section on ‘Driving and using machines’ as well as reviewing the previous categorization (if available) of the medicine in Belgium, France, the Netherlands, Spain and the International Council on Alcohol, Drugs and Traffic Safety (ICADTS) list.

In the cases of severely impairing medicines, recently approved medications, or medicines belonging to the ATC groups N and R06, all the collected data were compiled in fact sheets with a standardized lay-out, which were used...
Figure 4
Flowchart representing the methodology that was followed during the DRUID categorization process. Legend: SmPC, Summary of Product Characteristics; PIL, Patient Information Leaflet; EMA, European Medicines Agency; MHRA, Medicines and Healthcare products Regulatory Agency; eMC, Electronic Medicines Compendium; IMB, Irish Medicines Board.
during the active substance evaluation procedure and the approval of its final category.

After evaluating all the available data, a provisional category was assigned to each active substance. The provisional category was proposed and discussed during WP4 meetings, where a final and definitive category was assigned and approved by all WP4 partners.

It is important to note that the DRUID methodology on the categorization of medicines affecting driving fitness allows not only to categorize an active substance but also to revise a previously assigned category, in cases where new evidence emerges, by following the same 5 step approach (Figure 4).

**Medicines to be categorized**

The following ATC groups were considered in the categorization process:

- A – Alimentary tract and metabolism
- B – Blood and blood forming organs
- C – Cardiovascular system
- D – Dermatologicals
- M – Musculoskeletal system
- N – Nervous system
- R – Respiratory system
- S – Sensory organs

**Table 1**

<table>
<thead>
<tr>
<th>Declaration of undesirable effects that can potentially impair the fitness to drive safely</th>
<th>DRUID category</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very common (&gt;1/10)</td>
<td>Category II or III</td>
</tr>
<tr>
<td>Common (&gt;1/100, &lt;1/1000)</td>
<td>Category I</td>
</tr>
<tr>
<td>Rare (&gt;1/10 000, &lt;1/10000) or very rare (&lt;1/10 000)</td>
<td>Category 0</td>
</tr>
</tbody>
</table>

**Table 2**

<table>
<thead>
<tr>
<th>System organ class</th>
<th>Selection of side effects that can impair the ability to drive safely</th>
</tr>
</thead>
</table>
| Nervous system disorders | • Somnolence, dizziness, drowsiness  
• Confusion – cognitive disorder- disorientation  
• Involuntary movement disorders: ataxia, tremor, parkinsonism, acute dystonic (dyskinesia) and dyskinetic reactions (dystonia)  
• Convulsions – seizures |
| Psychiatric disorders | • Perception disturbances (hallucination, visual hallucination, auditory hallucination, illusion)  
• Psychotic reactions and psychotic disorder (including paranoia psychosis)  
• Other: emotional lability, mood swings, aggression, nervousness, irritability, personality disorders, thinking abnormal, abnormal behaviour, euphoric mood, restlessness (emotional state of excitement), depersonalization |
| Eye disorders | • Diplopia or double vision,  
• Blurred vision  
• Accommodation disorders  
• Visual acuity reduced  
• Photophobia  
• Other: visual field defect, peripheral vision loss, altered visual depth perception, oculogyric crisis |
| Ear and labyrinth disorders | • Vertigo  
• Hearing loss  
• Other: buzzing, tinnitus |
| Metabolism and nutrition disorders | • Hypoglycaemia |
| Vascular disorders | • Hypotension |

**Results**

Three thousand fifty-four medicines were considered for inclusion into the categorization process. Of these 3054 medicines, 1513 were not categorized because they were not available on the EU market.

The distribution of the 1541 categorized medicines (see supplementary data) was as follows (Figure 5): Category 0 50.3%, Category I 26.0%, Category II 11.2%, Category III 5.8%, Multiple categories 4.4% and Depending on the medication in combination 2.3%. This figure shows that the majority of medications belong to either category 0 or category I (Figure 5).

It is important to note that the term ‘multiple categories’ refers to the fact that a certain medication could be included in more than one category. There could be several reasons for this, such as different routes of administration of the same active substances (e.g. topical, oral, parenteral, etc), different pharmaceutical formulations (e.g. aqueous vehicle, cream, drops or ointment, etc), different dosages administered, etc.

With respect to the terminology ‘depending on medicines in combination’, it is relevant to observe that this approach was used when the categorization depended on the combination of the medication under evaluation with another active substance. In these cases, since the ATC classification [18] often did not report the medicine used in combination, it was decided not to use a final category but to follow the above-mentioned approach.

Table 3 gives an overview of the distribution of the medicines in each category, stratified by ATC group. It is apparent from this table that the N group contains the highest number of category III medications. A detailed description of the category distribution within the N group is depicted in Table 4. The N05 sub-group shows the highest number of category III medicines, followed by the...
N01 sub-group. The N05 sub-group also contains the highest number of medications assigned to more than one category.

**Discussion**

The current DRUID categorization system establishes and defines standardized and harmonized criteria to categorize commonly prescribed medicines based on their influence on fitness to drive. To date, this system nearly embraces the full ATC index and it intends to provide a complete coverage of the most commonly prescribed medications in Europe. This categorization procedure is developed by a European group of experts and is meant to go beyond the national context to address a broader European scenario and involve different facets of health care practice.

The categorization system could be seen as a tool to improve prescribing and dispensing procedures both at a national and European level and, therefore, as an instrument to inform and involve HCPs better [11, 30]. In this respect, it is important that HCPs know the fundamentals of the categorization system and use it properly in order to inform fully their patients about the risks of driving under the influence of impairing medicines. Furthermore, HCPs should be able to distinguish between the four levels of impairment and, if possible, choose the least impairing medication within the same therapeutic group. Moreover, this system should encourage HCPs to update their knowl-
edge on medicines and driving in order to be prepared to answer questions that patients might have on this topic [8, 11].

The DRUID categorization system should also be used as a tool to motivate HCPs to provide patients with clear information, communicate to patients the risk associated with driving under the influence of medicines and catalyze health care professional-patient discussions, leading to both safer prescriptions and patients who are more conscientious about their decision on whether or not to drive [8, 11, 30].

This classification could be a useful tool in helping patients be more involved in the decision-making process, understand the hazards of some medications to traffic safety and remind them to use caution while driving until their individual responses to their therapy have been well established.

To our knowledge, this is the first time that the European Commission assigned an expert group in the field of medicines and driving the task of establishing the criteria for a European classification system and developing a categorization system for relevant therapeutic groups of medications with respect to their impact on driving skills. The categorization efforts were carried out by an international group of DRUID partners, coming from six different institutions in Europe, and gathered all their scientific competence, knowledge, expertise, and experience in the field of road safety research and practice. All the available data from multiple sources were collected according to a standardized step-by-step procedure, which allows for the future maintenance and/or revision of the current DRUID categorization system as new evidence emerges in the future, and it also allows for the constitution of a consistent evidence-based classification methodology to categorize new medications prior to their market authorization. Last but not least, as reported above, the DRUID categorization system encompasses the entire ATC list. Therefore, it is the first categorization system to provide a nearly complete overview of the influence of frequently prescribed medications on the ability to drive. Additionally, in the cases of severely impairing medications (e.g. medicines from the N group), the system is integrated with fact sheets which concisely emphasize the key points of the categorization and can be easily used as a support mechanism in HCPs’ daily practice [24].

Lastly, some limitations of the DRUID categorization system should be considered. In particular, special attention should be paid to the fact that a category is attributed

<table>
<thead>
<tr>
<th>ATC group</th>
<th>N – nervous system</th>
<th>Not evaluated or not available in EU market</th>
<th>Druid categorization</th>
<th>Depending on the medicine in combination</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>N01 ANAESTHETICS</td>
<td>31</td>
<td>3</td>
<td>3</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>N01A Anaesthetics, general</td>
<td>20</td>
<td>11</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N01B Anaesthetics, local</td>
<td>11</td>
<td>3</td>
<td>3</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>N02 ANALGESICS</td>
<td>93</td>
<td>2</td>
<td>7</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td>N02A Opioids</td>
<td>31</td>
<td>2</td>
<td>7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N02B Other analgesics and antipyretics</td>
<td>52</td>
<td>2</td>
<td>6</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>N02C Antimigraine preparations</td>
<td>10</td>
<td>1</td>
<td>9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N03 ANTIPILEPTICS</td>
<td>23</td>
<td>14</td>
<td>1</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>N03A Antiepileptics</td>
<td>23</td>
<td>14</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N04 ANTIPARKINSON</td>
<td>16</td>
<td>3</td>
<td>16</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>N04A Anticholinergic agents</td>
<td>10</td>
<td>4</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N04B Dopaminergic agents</td>
<td>6</td>
<td>3</td>
<td>12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N05 PSYCHOLEPTICS</td>
<td>107</td>
<td>4</td>
<td>16</td>
<td>26</td>
<td>12</td>
</tr>
<tr>
<td>N05A Antipsychotics</td>
<td>31</td>
<td>13</td>
<td>8</td>
<td>9</td>
<td></td>
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<tr>
<td>N05B Anxiolytics</td>
<td>23</td>
<td>1</td>
<td>3</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>N05C Hypnotics and sedatives</td>
<td>53</td>
<td>3</td>
<td>11</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>N06 PSYCHOANALEPTICS</td>
<td>62</td>
<td>2</td>
<td>10</td>
<td>20</td>
<td>7</td>
</tr>
<tr>
<td>N06A Antidepressants</td>
<td>37</td>
<td>1</td>
<td>7</td>
<td>12</td>
<td>7</td>
</tr>
<tr>
<td>N06B Psychostimulants, agents used for ADHD and nootropics</td>
<td>22</td>
<td>3</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N06C Psycholeptics and psychonaleptics in combination</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N08D Anti-dementia drugs</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N07 OTHER NERVIOUS SYSTEM DRUGS</td>
<td>14</td>
<td>2</td>
<td>3</td>
<td>9</td>
<td>1</td>
</tr>
<tr>
<td>N07A Parasympathomimetics</td>
<td>6</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N07B Drugs used in addictive disorders</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>N07C Antivertigo preparations</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N07X Other nervous system drugs</td>
<td>4</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>346</td>
<td>9</td>
<td>30</td>
<td>86</td>
<td>53</td>
</tr>
</tbody>
</table>
to the single medicine, given to an adult, for its main indication, in a normal dosage, and at the start of the treatment [7, 8, 17]. Therefore, if a medication is not prescribed according to these conditions, it is crucial to bear in mind that the categorization system can only be used as background information, and it is necessary to carefully assess all the individual risk factors and avoid strict adherence to the medication classification. Furthermore, the system is focused on the effects of medications on fitness to drive and, consequently, the role of the disease, which could also influence fitness to drive, is not considered and certainly needs further attention while counselling the patient [7, 8].

Finally, the categorization system should always be associated with proper patient counselling in order to avoid any misunderstandings from the patient’s side and to ensure that the patient receives adequate information allowing him/her to make a consistent decision with the message given by the medication category.

**Next steps and recommendations**

The categorization system presented in this manuscript was developed within the DRUID project and, therefore, in a European context. As a consequence, the DRUID partners agreed that the European regulatory authorities should to be informed about this categorization process. This should lead to discussion and consensus on the criteria hereby proposed and special efforts should be carried out to implement the current system at both international and national level, with consideration country specific circumstances.

In this respect, it is important to underline that the DRUID consortium [31] previously approached the EMA Pharmacovigilance Working Party (PhVWP) in order to obtain its contribution in relation to the development of the categorization/labelling system for medications that impair driving [32]. In June 2011, the PhVWP agreed that any information on the influence of medicines on driving ability should be simple and helpful to the patient and, therefore, be reflected in the package leaflet. Furthermore, the PhVWP recommended including in the package leaflet a two-tier risk classification system differentiating between medicinal products with a potential for relevant influence on driving (moderate or major influence) and medicinal products without a potential for relevant influence (no or minor influence). Finally, the PhVWP recognized that this two-tier risk classification system could be further divided to include a maximum of four categories at the discretion of Member States [32]. This consensus is an important step in the harmonization of information on the potential for a medicine’s impairing effects on fitness to drive. However, it would be desirable for Member States to be provided with further discretionary activities, which could be used to reinforce the awareness of HCPs and patients on the effects of medicines on fitness to drive.

Since the categorization requires constant revision, it is also advised that an expert working group on medicines and driving be established to keep the system functional, up-to-date, and reliable.

Furthermore, it is recommended that special attention be paid to educating those who might play an active role in traffic safety. In this respect, medical and pharmacy schools should develop targeted educational programmes covering the issue of medication use and driving. Police officers and driving instructors should be adequately trained on this topic so that they are able to transfer knowledge about the effects of certain medications on a person’s ability to drive to potential patients who may drive in traffic.

Finally, a guideline should be developed to explain the use of the categorization system to HCPs and to serve as a support mechanism in the decision making process. On the other hand, since the PIL is the most accessible source of information for patients, it would also be advisable to develop an effective strategy to communicate the risk related to the use of medicines and driving. For instance, a straightforward grading system could be included in the patient package leaflet and warning labels in the form of pictograms could be printed on the medication box to provide clear instructions about the use of the medication and driving to patients.

**Disclaimers**

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2 This document reflects only the authors’ view. The European Community is not liable for any use that may be made of the information contained therein.

**Competing Interests**

The authors declare that they do not have any competing interests.

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