McKesson Europe Policy Position

How pharmacy sales of OTC Medicines contribute to better public healthcare

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Introduction

OTC medicines\(^1\) play a vital and possibly overlooked role in general population healthcare. When taken correctly, OTCs are highly accessible and relatively inexpensive and enable patients to manage symptoms of common illnesses. This provides important benefits not only for patients, but also for healthcare systems and economies. McKesson’s EU-owned pharmacies and those of our customers safely sell OTCs to millions of patients across Europe every day.

At McKesson Europe we believe that many common illnesses and symptoms can and should be addressed in a licensed pharmacy, where qualified pharmacists and their teams can select the right medicine and provide supporting professional advice. Critically, where a pharmacist or their trained staff do not consider an OTC intervention appropriate, they may instead offer an alternative intervention such as signposting the patient for non-pharmaceutical care. This kind of professional pharmacy intervention sets pharmacy transactions apart from non-medicine sales or OTC sales outside of a licensed pharmacy.

Recognising that medicines require special treatment, in this paper we call on national policy makers to ensure that their regulatory environment protects the public from harm whilst ensuring availability of OTC medicines when they are needed.

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\(^1\) For the purpose of this document, Over the Counter (OTC) medicines are defined as non-prescription medicines (NPMs) which the public can purchase from a licensed pharmacy and other non-pharmacy outlets (i.e. drug stores, petrol stations, etc.). Across Europe no agreed common definition of OTC exists.
Medicines are not ordinary products

Medicines can bring many benefits, but can also be harmful when not used properly. They must not be considered as ordinary commercial goods, but as highly regulated health products – an approach which has been explicitly recognised in EU legislation.  

OTC medicines are typically used to manage symptoms of common illnesses. They play an important role in the broader self-care agenda, which includes other non-pharmaceutical support to deliver better overall health and well-being. Although beneficial however, OTC medicines are not without risks, as highlighted in the table below.

Table 1: OTC medicines – the risks and the benefits

<table>
<thead>
<tr>
<th>Benefits</th>
<th>Risks</th>
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<tbody>
<tr>
<td>Low cost</td>
<td>Unforeseen side-effects and complications due to interaction with other medicines</td>
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<tr>
<td>Readily available - patients do not require GP appointment / prescription</td>
<td>Repeat sales less likely to be picked up if sold outside of a licensed pharmacy</td>
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<td>Reduced pressure on healthcare professionals re minor complaints – freeing time for more serious ones</td>
<td>Inappropriate dosage and / or incorrect self-diagnosis</td>
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<td>Highly accessible: quick relief of the medical problem</td>
<td>Medicine dependency and / or deliberate misuse</td>
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<td>Individual can exercise more autonomy and choice</td>
<td>Unwanted hospital admissions due to Adverse Drug Reactions (ADRs)</td>
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<td></td>
<td>Additional GP or hospital visits if used incorrectly</td>
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Access to OTC medicines must be controlled to deliver all the benefits and manage the risks listed above. We believe that the rules regarding sales of OTC medicines should adequately reflect the special status of medicines. Accordingly, switching medicines from prescription status to another category needs to address potential abuse and misuse concerns as well as ADRs. Patient safety must remain the top priority.

Our key messages

- Medicines are unlike ordinary commercial goods: they provide enormous benefits to patients but can be harmful if not used properly
- Pharmacy personnel are best placed to provide professional, personalised advice to patients on the most appropriate use of medicines
- Sales of OTC medicines through non-pharmacy channels can, in certain cases, put patients at risk through lack of supervision and professional advice
- If Rx medicines are reclassified to non-prescription status, we recommend creating a Pharmacy Medicines category

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2 This includes the Falsified Medicines Directive (Directive 2011/62/EU) Recital 22, which recalls that the Court of Justice of the EU ‘has recognised the very particular nature of medicinal products (including NPMs)’ and TFEU Article 168 paragraph 4c.
3 Such as colds, coughs, pains or digestive conditions
4 For example, medical devices, fitness and diet advice and coaching
5 ADRs result in 6.5-10.9% hospital admissions and mortality rates of 0.15-2.9%. *Adverse Drug Reactions related to mortality and morbidity: Drug-drug interactions and overdoses; Angella Angiji, Xendo, 2018*
The professional and personal value pharmacies offer

McKesson Europe believes that the sale of OTCs is best conducted within the framework of licensed pharmacies under the professional supervision of qualified pharmacists or their teams operating under the pharmacy’s quality control system, e.g. pharmacy technicians. Pharmacists have been practising a form of personalised medicine for years. Only they are best placed to:

- Advise patients on appropriate medicine and dosage, based on their personal needs / medication history;
- Provide special advice for OTC use by vulnerable patients;
- Warn about side-effects and interactions with other medicines, and the necessary precautions to take;
- Provide holistic advice to a patient about managing their medical condition.

Medicine Reclassifications – A Pharmacy Medicine

McKesson Europe welcomes the reclassifications of medicines from prescription only to non-prescription status where sufficient evidence exists to support patient safety claims.

In such cases, we recommend that these medicines should be considered across the EU as a special category – a Pharmacy Medicine (P Med). This category already exists in some Member States where such medicines are not immediately accessible to patients without the supervision of a pharmacist. This status ensures that patients get professional support with their medicine purchases. We would encourage at Member State level a review and the possible translation of current OTC medicines and products into this newly defined P Med category. Some Member States may already consider their current OTC categorisation as sufficiently robust. In other countries the terms OTC and Pharmacy Medicines are interchangeable. We believe that these are unsustainable situations, as there is increasing pressure from non-healthcare outlets for liberalisation of the OTC sector. Their arguments are based predominantly on price and competition. The European Commission also broadly supports OTC liberalisation.

We would strongly urge pharmacy regulators to create the Pharmacy Medicine category as a recognised subset of what is today called OTC medicines.

Uncontrolled sales through non-pharmacy channels

As discussed, Member States are reviewing OTC medicines currently thought to be of lesser risk, with a view to making them available through non-pharmacy channels, i.e. outside the pharmacy regulation framework. Recognising that staff at retail outlets have no medical training, that patients may be unaware of any risks, may be over-confident in their ability to navigate purchases on their own, or may return later for repeat sales, we believe that stronger controls are required to protect the public.

We recommend that where OTC medicines are reclassified, or sales are made possible through non-pharmacy channels, that these should be for a very limited number of medicines for immediate and short-term use only. Furthermore, we believe that these medicines should not be available for self-selection and should be stored out of children’s reach. We would also recommend that outlets selling these products should also be subject to additional controls to protect public health.

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6 Such as children, pregnant women, the elderly or carers
7 European Commission Communication and Staff Working Document: A European retail sector fit for the 21st century, April 2018
8 Typically convenience stores or petrol stations
9 In packs of four or less, while multiple pack sales should not be allowed thereby reducing the chance of accidental or deliberate overdosing.
Maintaining a comprehensive community pharmacy network

Licensed community pharmacies provide a public service role – not only do they supply medicines and free health advice, but they also relieve pressure on other primary and secondary care providers by providing advice immediately and without an appointment. This is very convenient for 98% of Europeans who live within 30 minutes of a community pharmacy\(^{10}\) – often their first healthcare contact point.

In many countries, OTC medicine sales have historically formed a key part of a traditional ‘bricks and mortar’ community pharmacy’s sales, ranging from 5% to over 50% of a typical pharmacy’s income.\(^ {11}\) Removing these sales could undermine the financial sustainability of many community pharmacies, particularly in less populated areas. We strongly recommend that policy makers think cautiously about liberalising for short-term gains.

Our recommendations

- Policy makers, and in particular competition and anti-trust authorities, in all countries should recognise that OTC medicines are not ordinary commercial goods.
- McKesson Europe believes that increasing access to medicines is best enabled by increasing the reclassification of medicines from prescription-only status to pharmacy-only sales.
- Where sufficient evidence exists to support a medicine’s reclassification from prescription to non-prescription status, we believe that such medicines must remain in licensed pharmacies under a Pharmacy Medicine (P Med) category. In this way, they remain under the appropriate supervision of pharmacists and their teams. Where this category does not exist, we are calling on pharmacy regulators to make the necessary changes to guarantee public access and safety through the creation of this category.
- McKesson Europe does not believe it is necessary for non-healthcare policy makers to facilitate increased access to P Meds and OTC medicines as we consider this a public health matter. Such decisions should remain the competence of medicines authorities. However if changes are sought to increase availability, we strongly advocate that this applies only to very small packs for limited symptomatic relief medicines with proven safety profiles.

About McKesson Europe

McKesson Europe is a leading international wholesale and retail company and provider of logistics and services to the pharmaceutical and healthcare sector. With about 39,000 employees, the group is active in 13 European countries. Every day, the company serves over 2 million customers – at more than 2,100 pharmacies of its own, at about 300 managed pharmacies and at over 5,700 participants in the brand partnership schemes. With 110 own and seven managed wholesale branches in Europe, McKesson Europe supplies more than 55,000 pharmacies and hospitals every day with up to 130,000 pharmaceutical products.

Facts and Figures


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\(^{10}\) Survey of Chain of Trust Project, under EC Public Health Programme (Grant Agreement N° 2009 11 13)

\(^{11}\) From PGEU internal survey.
Definitions

OTC Medicines. In the course of our research it became clear that there is no agreed definition of Over the Counter (OTC) medicines. For the purpose of this Paper, OTCs are defined as non-prescription medicines (NPMs) which the public can purchase from a licensed pharmacy and other non-pharmacy outlets (i.e. drug stores, petrol stations, etc.).

Pharmacy Medicines (P Meds). This definition will be new to some readers. Medicines in a P Med category are subject to stricter rules and sales criteria designed to ensure the public gets the most appropriate treatment for them. In addition P Med sales are conducted safely under the supervision of a pharmacist.

Some typical attributes of the P Med category

- This applies to the safe and professional sale of recognised medicine products or treatments as defined under national legislation.
- This is a product specifically referred to as a P Med in national legislation, for example, a product in a certain package / containing certain amount of pills / tablets.
- Medicines not the P Med category would be either prescription medicines, or non-medicinal products.
- If sold through an online pharmacy, strict questions and standards of sale will equally apply.
- A set of criteria typically applies to all P Med sales:
  - Sales conditions – Questions about whether the medicine was previously taken, symptoms, who the medicine is for etc.
  - Quantity criteria – Only one or two packs; for any more the pharmacist must use professional judgement
  - Display conditions – Not directly self-selectable without supervision, but can be displayed in the pharmacy in secure units
  - Safety conditions – P Meds will always include recently reclassified prescription medicines
  - Additional or fewer criteria may apply depending on the country in question.

For detailed examples please review the legislation applying to Pharmacy Medicines in the UK or Ireland.
General OTC safety concerns

Europe: A recent study on availability of paracetamol and reported enquiries to Poisons Information Centres found that:

‘A significantly lower median frequency of paracetamol-related enquiries was found in countries without non-pharmacy outlet sales compared to those with such sales (median difference 2.2%, p = 0.02).’ ¹

Denmark: A recent study from Danish researchers found that the use of ibuprofen and other non-steroidal anti-inflammatory drugs (NSAIDs) was associated with increased health risks, including risk of cardiac arrest. ²

Ireland: The pharmacy regulator (PSI) published updated guidelines for pharmacists in October 2017 on the safe supply of non-prescription medicines containing codeine, stating:

‘The safety concerns around the misuse of non-prescription medicinal products containing codeine are well established. Consumption of quantities of these medicines in excess of the recommended dose, or over a prolonged period of time, may cause tolerance and dependence, as well as the risk of other adverse effects. Furthermore, the consumption of excessive quantities of ‘combination products’, i.e. those containing codeine and another analgesic such as paracetamol, aspirin or ibuprofen, also increases the risk of harm from these other medicinal products. This risk applies to both short and long-term use.’ ³

National examples of misuse of OTCs

Finland: A survey of over 700 pharmacists for the Association of Finnish Pharmacies published in January 2018 found that half of the respondents reported customers buying OTC medicines for the wrong purposes on a weekly basis, and one in four reported misuse at least once a day. Medicines which the respondents named as commonly misused included cough medicines, nasal decongestants, painkillers, drinks for treating flu symptoms, laxatives, cortisone lotions, antibiotic ointments, motion sickness medication and aspirin. ⁴

Ireland: In 2016, paracetamol was used in 1,234 reported poisoning cases, out of a total of almost 5000 involving drugs. ⁵

In 2014 the police asked pharmacists to practice caution in dealing with requests for pseudoephedrine, an active ingredient in non-prescription cough and cold medicines, as it was being used by criminal gangs for the production of methamphetamine. ⁶

¹ Availability of Paracetamol Sold Over the Counter in Europe: A Descriptive Cross-Sectional International Survey of Pack Size Restriction; Morthorst BR et. al, Basic Clin Pharmacol Toxicol. 2018 Jan
³ Non-Prescription Medicinal Products Containing Codeine: Guidance for Pharmacists on Safe Supply to Patients, Pharmaceutical Society of Ireland, October 2017 http://thepsi.ie/Libraries/Folder_Pharmacy_Practice_Guidance/01_3_Safe_supply_of_Codeine_to_patients.sflb.ashx
⁵ Poisons Information Centre of Ireland, Annual Report 2016
**Norway**: A survey by the Norwegian Pharmacy Association conducted in 2015 on attitudes to paracetamol among 15-24 years showed that around half of the 400 respondents had a different opinion of medicines bought in groceries to those bought in a pharmacy – they perceived the former to be ‘milder’.

**Poland**: A 2016 study based on a survey of 680 pharmacists found a significant problem:

‘The misuse of OTC drugs is increasing in Poland from pharmacists point of view. The most popular substance was PSD followed by COD and DXM. The main reason of misuse of these drugs could be related to the use of Internet and free access to these medications. In respondents (58.2%) opinion OTC drugs containing analyzed substances should be moved into the prescription status.’

**Sweden**: The medicines supervisor (Läkemedelsverket) ordered the removal of paracetamol pills from supermarkets in 2015 after an increase in cases of misuse – poisoning cases had increased by 40% since sales were liberalised in 2009.

In 2014 oral medication with fluconazole was switched from Rx to OTC and only sold in pharmacies. In 2016 the Läkemedelsverket ordered the removal of oral medications with fluconazole from pharmacies as from June 2017, switching it back to Rx from OTC. The order was issued due to the correlation between use of antibiotics and antibiotic resistance.

**UK**: According to the research published in the Journal of Public Health, a fifth of the UK population may have misused NPMs in some way during their lifetime, either by taking a higher dose than recommended, using a product more often than recommended or by using a medicine beyond the recommended time limit.

**Public support for P Med category**

Ireland: In January 2018, the Irish Pharmaceutical Healthcare Association (IPHA) and the Irish Pharmacy Union (IPU) published a market research report which they had jointly commissioned detailed market on attitudes towards self care in Ireland. This included the following observation regarding attitudes to pharmacy:

‘Focusing on those who made an OTC medicines purchase, arguably the group most focused on self care, the research highlights that they were much more likely to assess the following facets as being important contributors to their decision on what to buy: staff attitude, interaction with staff, past relationship with the pharmacy and advice from the counter staff or pharmacy assistants. All of these elements were more contributory for this group than they were for others.

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6 Vigilance in the Sale of Pseudoephedrine- Garda Request, The Pharmaceutical Society of Ireland, June 2014  
http://www.thepsi.ie/gns/Pharmacy_Practice/PracticeUpdates/VigilanceinthesaleofPseudoephedrine.aspx  
7 Misuse of OTC drugs in Poland, Zaprutko T et al, Health Policy. 2016 Aug;120(8):875-81  
8 Paracetamol tablets only available in pharmacies, Swedish Medical Products Agency, 30 October 2015  
https://lakemedelsverket.se/english/All-news/NYHETER-2015/Paracetamol-tablets-only-available-in-pharmacies-  
Increased availability of paracetamol in Sweden and incidence of paracetamol poisoning: using laboratory data to increase validity of a population-based registry study, Gedeborg R et al., Pharmacopsidemiol Drug Saf. 2017 May;26(5):518-527  
10 SELF CARE Taking charge of your health, January 2018  