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Accompanying the document
Proposal for a DIRECTIVE OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL on the approximation of the laws, regulations and administrative provisions of the Member States concerning the manufacture, presentation and sale of tobacco and related products
(Text with EEA relevance)

Delegations will find attached Commission document SWD(2012) 452 final (Part 1).

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Part 1

COMMISSION STAFF WORKING DOCUMENT

IMPACT ASSESSMENT

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Proposal for a

DIRECTIVE OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL

on the approximation of the laws, regulations and administrative provisions of the Member States concerning the manufacture, presentation and sale of tobacco and related products

(Text with EEA relevance)

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ABBREVIATIONS USED IN THE IMPACT ASSESSMENT

ENDS Electronic Nicotine Delivery System
FCTC WHO Framework Convention on Tobacco Control
FMC Factory Manufactured Cigarettes
NCP Nicotine Containing Products
NRT Nicotine Replacement Therapy
PA Policy Area
PO Policy Option
PoS Point of Sale
RYO Roll-Your-Own tobacco
STP Smokeless Tobacco Products
SKU Stock Keeping Unit
TEU Treaty of the European Union
TFEU Treaty on the Functioning of the European Union
TNCO Tar, nicotine and carbon monoxide
TPD Tobacco Products Directive
TVM Tobacco Vending Machine
WHO World Health Organisation
WTO World Trade Organisation
GLOSSARY OF TERMS

Additive – substance contained in a tobacco product, its unit packet or its outside packaging with the exception of tobacco leaves and other natural or unprocessed parts of tobacco plants.

Characterising flavour - a distinguishable aroma or taste other than tobacco, resulting from an additive or combination of additives, including but not limited to fruit, spice, herb, alcohol, candy, menthol or vanilla observable before or upon intended use of the tobacco product;

'Cheap whites / illicit whites' - cigarettes produced (often legitimately) in their country of origin at very low cost, destined to be illicitly sold in other jurisdictions and not respecting the legal requirements in the jurisdiction of destination.

Chewing tobacco - a smokeless tobacco product exclusively designed for the purpose of chewing.

Cigar - a roll of tobacco consumed via a combustion process and further defined in Article 4(1) of Council Directive 2011/64/EU of 21 June 2011 on the structure and rates of excise duty applied to manufactured tobacco.

Cigarette – a roll of tobacco consumed via a combustion process and further defined in Article 3(1) of Council Directive 2011/64/EU.

Cigarillo – a small type of cigar with a diameter of up to 8 mm.

Contraband - products which have been diverted into illicit trade, not respecting the legal requirements in the jurisdiction of destination.

Counterfeit – brand protected products which have been falsified without consent of the brand owner and are not respecting the legal requirements in the jurisdiction of destination.

Electronic cigarette (Electronic Nicotine Delivery Systems, ENDS) – electronic device typically consisting of a mouth piece (containing an electronic evaporator) and a cartridge (typically replaceable) and designed to deliver nicotine to the lung through inhalation of a mixture of air & vapours into the respiratory system.

Factory manufactured cigarette (FMC) – a cigarette, produced by a tobacco manufacturer, capable of being smoked as such.

FCTC commitments – political commitments to implement the non-binding guidelines developed under the FCTC to assist Parties in meeting their implementation obligations under the FCTC.

FCTC obligations – obligations to implement the legally binding FCTC and the Illicit Trade Protocol.

Flavouring – an additive that imparts aroma and/or taste.

Herbal products for smoking – a product based on plants or herbs which contains no tobacco and is consumed via a combustion process.

Ingredient – an additive, tobacco (leaves and other natural, processed or unprocessed parts of tobacco plants including expanded and reconstituted tobacco), as well as any substance present in a finished tobacco product including paper, filter, inks, capsules and adhesives.

Illicit trade – any practice or conduct prohibited by law and which relates to production, shipment, receipt, possession, distribution, sale or purchase, including any practice or conduct intended to facilitate such activity.

Nasal tobacco - a smokeless tobacco product consumed via the nose.

1 The purpose of this glossary is to provide the reader with a better understanding of the terms used in the document. It should in no way prejudge the terminology defined in the legal proposal.
Nicotine containing products (NCP) – a product usable for consumption by final consumers via inhalation, ingestion or in other forms and to which nicotine is either added during the manufacturing process or self-administered by the user before or during consumption.

Nicotine Replacement Therapies (NRT) - remedial administration of nicotine to the body by means other than tobacco, usually authorised under the pharmaceutical legislation as part of smoking cessation. Common forms of nicotine replacement therapy are nicotine patches and nicotine gum.

Novel tobacco product - a tobacco product other than a cigarette, roll-your-own tobacco, pipe tobacco, water-pipe tobacco, cigar, cigarillo, chewing tobacco, nasal tobacco or tobacco for oral use placed on the market after entry into force of the Directive.

Pipe tobacco – tobacco consumed via a combustion process and exclusively designed for the purpose of being used in a pipe.

Plain packaging – full standardisation of the packages, including brand- and product names printed in a mandated size, font and colour on a given place of the package; standardised package colour; standardised size and appearance of the package; display of required (textual and pictorial) health warnings and other legally mandated product information, such as tax-paid stamps and marking for traceability and security purposes.

Promotional / Misleading element – any element promoting a tobacco product by a means that is false, misleading, deceptive or likely to create an erroneous impression about its characteristics, health effects, hazards or emissions, any element suggesting that a tobacco product is less harmful than others or has vitalising, energetic or other positive health effects, any element referring to flavour or taste or the absence thereof, or any elements resembling a food product. Such elements can take the form (but are not limited to) texts, signs, pictures or other graphical elements, references to natural or biological characteristics or to certain flavours or flavourings or other additives, inserts and other additional material, e.g. adhesive labels, stickers, onserts, scratch-offs, sleeves.

Roll-your own tobacco (RYO) – tobacco which can be used for making cigarettes by final consumers or retail outlets.

Smokeless tobacco products (STP) – a tobacco product not involving a combustion process, including tobacco for oral use.

Tobacco for oral use/oral tobacco - all products for oral use, except those intended to be inhaled or chewed, made wholly or partly of tobacco, in powder or in particulate form or in any combination of those forms, particularly those presented in sachet portions or porous sachets.

Traditional use – Continuous use of a smokeless tobacco product in a Member State or part thereof for at least 30 years.
MAIN REPORTS/STUDIES USED FOR THE IMPACT ASSESSMENT

- Special Eurobarometer 332; 2010 (Eurobarometer 2010): http://ec.europa.eu/health/eurobarometers/index_en.htm
- Matrix insight. Economic analysis of the EU market of tobacco, nicotine & related products 2012 (Matrix 2012)
1. INTRODUCTION

1.1. CONTEXT


More than ten years have passed since the adoption of the TPD. In line with market, scientific and international developments it has become necessary to update and complete the TPD. A revision is explicitly foreseen in Article 11 of the current TPD and was repeatedly called for by the Council and the European Parliament³. The initiative to revise the TPD is included in the Commission's Work Plan 2012.⁴

The overall objective of the revision is to improve the functioning of the internal market. In particular, the proposal aims to:

- Update already harmonised areas to overcome Member States' obstacles to bring their national legislations in line with new market, scientific and international developments.⁵

- Address product related measures not yet covered by the TPD insofar as heterogeneous development in Member States has led to, or is likely to lead to, fragmentation of the internal market.⁶

- Ensure that certain provisions of the TPD are not circumvented by placing on the market of products not compliant with the TPD.⁷

It is also important to ensure a harmonised implementation of FCTC obligations and a consistent approach to non-binding FCTC commitments if there is a risk of diverging national transposition.

In line with Article 114(3) TFEU a high level of health protection has been taken as a basis for this impact assessment when choosing between different policy options. In this context, the revision seeks to regulate tobacco products in a way that reflects their specific characteristics (nicotine has addictive properties) and the negative consequences of their

² OJ L 194, 18.7.2001, p. 26–35
⁵ Without an update, Member States cannot, for example, increase the size of the health warnings, change their location of the package or replace the display of tar, nicotine and carbon monoxide levels.
⁶ For example, at this stage, eight Member States have adopted pictorial health warnings and the regulations of ingredients differ between Member States.
⁷ For example, measures on cross-border distance sales and traceability will facilitate legal activity and thus prevent sale of tobacco products not complying with the TPD (e.g. health warnings and ingredients).
consumption (health risks such as various cancer types, cardiovascular problems, increased risk of blindness, impotence, lower fertility, impact on the unborn child etc.). Their treatment costs more than 25 bEUR per year. Furthermore, tobacco is the most significant cause of premature deaths in the EU, responsible for almost 700,000 deaths every year (see section 2.1.2). The revision focuses on initiation of tobacco consumption, in particular by young people, taking into account that 70% of the smokers start before the age of 18 and 94% before the age of 25 years. This is also reflected in the selection and focus of the policy areas proposed and the products primarily targeted (FMC, RYO and STP).

From a broader perspective, the revision will contribute to the overall aim of the EU to promote the well-being of its people (TEU Article 3) and the Europe 2020 strategy as keeping people healthy and active longer, and helping people to prevent avoidable diseases and premature death, will have a positive impact on productivity and competitiveness. An unintended, but welcome side effect of the measures against trade of products not complying with the requirements of the TPD might be that the tax revenues of Member States are better protected as the products often also circumvent national tax legislations.

The revision of the TPD focuses on five policy areas: (1) STP and extension of the product scope (i.e. NCP and herbal products for smoking), (2) packaging & labelling, (3) ingredients/additives, (4) cross-border distance sales and (5) traceability and security features. When preparing this impact assessment report economic, legal and scientific considerations were taken into account. Particular attention was given to the Fundamental Rights Charter and international obligations (FCTC, WTO-TRIPS, TBT).

This impact assessment report presents the analysis and all relevant results of the impact assessment work. Due to space limitations, citations are limited to key publications and, as appropriate, relevant studies illustrating the current evidence base. More detailed information and supporting materials are also included in the five technical annexes accompanying the main report. The first four annexes provide more detailed information on stakeholders' views, the tobacco market, the regulatory framework and the assessment criteria used when assessing the impacts as well as scoring tables. Annex 5 outlines the socio economic impacts and explains in detail how a reduction in tobacco consumption will impact on stakeholders (indirect impacts).

1.2. APPLICATION, IMPLEMENTATION AND ENFORCEMENT OF THE CURRENT TPD AND Notifications FROM MEMBER STATES

1.1.1. Content of the existing TPD

The existing TPD was adopted to recast two previous internal market Directives. According to its Article 1, the TPD aims at approximating certain national rules regarding tobacco products, e.g. tar, nicotine and carbon monoxide (TNCO), health warnings, ingredients and misleading description of tobacco products. The current TPD is limited to products containing tobacco (i.e. NCPs and herbal products for smoking are not subject to the TPD). It applies to all categories of tobacco: FMC, RYO, pipe tobacco, cigars, cigarillos, STP and other forms of tobacco.

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8 Eurobarometer 2012
9 This includes also novel tobacco products which are primarily expected to fall within the category of STP.
11 Directives 89/622EEC and 90/239/EEC
Article 3 of the TPD sets the maximum levels for **tar, nicotine and carbon monoxide** (TNCO) and Article 4 explains how the TNCO levels should be measured. Regarding **ingredients**, Article 6 foresees reporting obligations for the industry (including available data on toxicology and addictiveness). Article 12 invites the Commission to submit a common list of ingredients authorised for tobacco products. The Commission has not suggested such a list taking into account a shift in regulatory priorities, including in the context of the FCTC and the adoption of partial guidelines on ingredients related to attractiveness.

Article 5 contains **labelling** requirements. It stipulates that all tobacco products except STP must carry a general health warning, (e.g. "Smoking kills"), covering not less than 30% of the front side, and a specific text warning (e.g. "Smoking causes fatal lung cancer"), covering not less than 40% of the back side. The TPD also requires that all STP carry a health warning ("This tobacco product can damage your health and is addictive."). Moreover, the TPD empowers the Commission to adopt rules for the use of additional pictorial warnings that Member States have to comply with if they decide to require those warnings. In addition, the packages should display the levels of tar, nicotine and carbon monoxide (TNCO). Article 7 of the TPD prohibits the use of trademarks and texts suggesting that a particular product is less harmful than others (e.g. "mild" or "light"). To ensure product identification and traceability, the tobacco products should be marked by batch numbering enabling the place and time of manufacture to be determined (Article 5 (9) TPD). The Commission was invited to provide technical details, but has not responded to the invitation in the light of new international developments the Commission concluded that the information on time and location of manufacturing was not sufficient to ensure full traceability and reduce illicit trade effectively.

Article 8 prohibits the placing on the market of **oral tobacco** (snus) outside Sweden.

Articles 9, 10 and 11 of the TPD contain **comitology** provisions and **reporting** obligations. Article 13 sets out the conditions under which Member States can take stricter provisions.

### 1.1.2. Application of the TPD

Article 11 of the current TPD requires that the Commission reports regularly on the application of the Directive. Two such reports have been issued, a first one in July 2005 and a second one in November 2007. Subsequent reports were not issued in the light of the pending revision/impact assessment process.

The **First Report on the Application of the TPD** concluded that the Commission should consider further the development of labelling, such as the wider use of quit line telephone numbers. As regards reporting of ingredients, it was stressed that Article 6 on the reporting of ingredients needs to be developed, that information transmitted from the industry varies greatly and that there is lack of capacity to analyse the data.

In its **Second Report on the Application of the TPD**, the Commission concluded that it should examine the possibilities with regard to an increased size of the warnings, mandatory pictorial warnings on both sides of the packets and the replacement of TNCO levels by other information. The Commission also stressed that it should explore the possibilities of generic

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standardised packaging. As regards reporting of ingredients, the report refers to the wish from a number of Member States and the industry to make the formats developed on a voluntary basis compulsory throughout the EU. In terms of oral tobacco, it was concluded that the scientific opinion on the health effects of STP (SCENIHR 2008) should form the scientific basis for any future risk management decision of the Commission. In addition, it was concluded in the report that new tobacco and/or nicotine products should be studied with a view to ensure proper regulation. Both reports have provided input to the current impact assessment. Both reports also covered Article 12 of the TPD on ingredients. For the reasons outlined above, not concrete follow-up was given to this provision.

1.1.3. Implementation of the TPD

Member States have transposed the existing TPD and the Commission has used its powers to adopt rules for the use of colour photographs and other illustrations and to amend the textual health warnings in line with scientific and technical progress. However, the Commission has not made use of its powers to adopt measures for identification and traceability purposes nor has the Commission responded to the invitation from legislators to develop a common list of ingredients.

1.1.4. Enforcement of the TPD and legal challenges

In general, enforcement of the TPD has not been seen as a problem and only a limited number of infringement procedures have been launched. However, the current TPD is unclear as regards to the level of harmonisation. While Article 5(5) allows Member States a degree of discretion to adapt the labelling of tobacco products to the requirement of public health protection, Article 13(1) stipulates that they cannot, for considerations relating to health warnings, prohibit or restrict the import of tobacco products complying with the TPD provisions. This Article has been subject to several consultations of the Commission's Legal Service. The current situation implies that Member States are allowed to take certain actions, but only for domestically produced products while they cannot impose the same requirements on imported products. This does not make sense in a globalised market such as the tobacco market and - taking in account the significant cross border trade - can easily result in production moving to countries where less stringent rules are required. Only a revision can address these shortcomings. In addition, the development of the internet as a distribution channel for tobacco products and the definition of “oral tobacco” represent particular challenges in terms of enforcement. Also the current requirement for rotation of the health warnings has been subject to different interpretation and needs to be clarified.

The current TPD has been subject to several legal challenges since its entering into force. In 2001 British American Tobacco (BAT) and Imperial Tobacco, initiated legal proceedings in the British Courts on the validity and interpretation of the Directive. The case was referred to

14 SCENIHR 2008
16 Two judgments concerning the previous tobacco labelling Directive 89/622 also addresses Member States' possibilities of imposing stricter national rules: C-222/91 Ministero delle Finanze and Ministero della Sanità v Philip Morris Belgium SA and others. European Court reports 1993 Page I-03469 and The Queen v Secretary of State for Health, ex parte Gallaher Ltd, Imperial Tobacco Ltd and Rothmans International Tobacco (UK) Ltd, European Court reports 1993 Page I-03545
the ECJ (Case 491/01). The companies argued that the legal basis for the directive (Article 95 TEC, current article 114 TFEU) was inadequate, because it was a public health measure being introduced as an internal market measure. The companies also argued that the introduction of Article 133 TEC as a second legal basis invalidated the directive. The principles of proportionality and subsidiarity were also claimed to have been infringed. They also argued that the labelling provisions for yields and larger health warnings, and the ban on misleading descriptors breached trade mark and intellectual property rights, as well as being an infringement of the obligation to give reasons. In 2002, the Court upheld the validity of the Directive and confirmed the validity of its provisions.

The directive’s ban on the marketing of certain types of oral tobacco (snus) has also been challenged (Cases C-434/02 and C- 210/03). These challenges were brought by Swedish Match, a manufacturer of oral tobacco (snus), against the UK government, and a German wholesaler who brought a case against the German government. They claimed that the directive was in breach of the rules laid down in Articles 95, 133 and 253 TEC. A claim was also made that the ban on oral tobacco was a breach of the principle of subsidiarity. They also alleged the directives' provision to constitute a restriction as referred to in Articles 28 and 29 TEC which prohibits quantitative restrictions in trade between Member States. One of the claimants also alleged that the ban was in breach of the principle of the freedom to pursue an occupation. Also in this case, the Court rejected the arguments and upheld the validity of the directive.

1.1.5. Notifications

A large number of notifications under Directive 98/34/EC have been received from Member States in the area of tobacco including on pictorial health warnings, the maximum number of FMC sticks in the package, display of quit lines on the package, regulations on ingredients and rules for herbal products for smoking. In addition, fourteen RAPEX notifications have been received so far regarding electronic cigarettes (17 December 2012).

1.3. Consultations, expertise and other input

Stakeholder consultations

A public consultation was held between 24 September and 17 December 2010. The Commission received more than 85,000 contributions from a wide range of stakeholders. Citizen contributions accounted for 96% of the survey response, 57% of which are “duplicate”/repeated responses which appear to be the result of several citizen mobilisation

19 Case C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd. [2002] ECR I-11453.
20 The Court found, however, that Article 95 TEC was the only appropriate legal base, but that the addition of Article 133 TEC as a legal base was not a reason for declaring the Directive invalid.
21 Case C-210/03 The Queen, on the application of: Swedish Match AB and Swedish Match UK Ltd v Secretary of State for Health. [2004] ECR I-11893.
23 RAPEX is the EU rapid alert system that facilitates the rapid exchange of information between Member States and the Commission on measures taken to prevent or restrict the marketing or use of products posing a serious risk to the health and safety of consumers with the exception of food, pharmaceutical and medical devices: http://ec.europa.eu/consumers/dyna/rapex/rapex_archives_en.cfm (accessed 28 Nov 2012).
24 A response considered “duplicate” in the public consultation was a response fulfilling the following criteria: 1. At least six responses containing the same text. 2. Text box containing more than three words. 3. Text box not containing text directly copied from the consultation document.
campaigns that took place in some Member States. The actions and efforts of these campaigns seem to have affected the overall quantitative data of the public consultation, which indicates that most of the citizens responding to the consultation were against changes to the TPD. This outcome deviates significantly from the latest Eurobarometer survey, published in May 2012. Unlike public consultations, it is important to note that respondents in a Eurobarometer survey are selected randomly. Member States representatives and - even more so - health NGOs favour the introduction of strict tobacco control measures, while tobacco industry and retailers are against some of the stricter measures (for more details, see Annex 1 and separate sections under the assessment of each policy area). A report presenting the outcome of the consultation was published on 27 July 2011 and contributions have been published online.

**Targeted discussions** with stakeholders took place throughout the revision process. A first exchange of views with health NGOs, tobacco- and pharmaceutical industries took place on 3 and 4 December 2009 and on 19 and 20 October 2010 and discussions with NGOs, growers, FMC producers, other tobacco producers, distributors of tobacco products and upstream suppliers of tobacco products have continued throughout 2011 and 2012. A number of written contributions were also received, which were carefully considered in assessing the impacts of different policy options. In particular, the criticism received in relation to the external study from RAND Europe (see below) was taken into account. Alternative data submitted by stakeholders was also carefully studied. The Commissioner for Health and Consumer Policy met with Health NGOs and economic stakeholders in February-March 2012. The revision of TPD has also been discussed regularly in the TPD Regulatory Committee from 2009 to 2012.

The policy area "traceability and security features" was added to the revision in response to concerns put forward by some stakeholders that the selling of contraband and counterfeit products not complying with the requirements of TPD is already today a significant problem. For the purpose of this Directive the main concerns associated with illicit products is that these products are non-compliant with the safeguards of the TPD. A more detailed summary of stakeholders' positions in the context of the consultations can be found in Annex 1.

**Inter Service Steering Group (ISSG)**

An Inter Service Steering Group (ISSG) was established in March 2009 to support the work of DG SANCO. The following services were invited: SG, SJ, AIDCO, AGRI, COMP, DEV, EAC, ECFIN, ECHO, ELARG, EMPL, ENTR, ENV, MARE, INFSO, JLS, JRC, MARKT, SJ, AIDCO, AGRI, COMP, DEV, EAC, ECFIN, ECHO, ELARG, EMPL, ENTR, ENV, MARE, INFSO, JLS, JRC, MARKT, SJ, AIDCO, AGRI, COMP, DEV, EAC, ECFIN, ECHO, ELARG, EMPL, ENTR, ENV, MARE, INFSO, JLS, JRC, MARKT,

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25 For example, a campaign was organised by a group representing over 75% of Italian Tobacconists (European Voice, 10 February, 2011). This action was followed by over 30,000 submissions, including 99% duplicate responses from Italy
26 Eurobarometer 2012
27 Public Consultation Report 2011. In addition to the contributions received on-line, the contributions received through other formats from 20 Member States at the level of Governments or ministries as well as from two EFTA/EEA countries have been published on the same web site.
29 Idem.
30 The minutes from the meetings can be found at: [http://ec.europa.eu/health/tobacco/events/index_en.htm#anchor0](http://ec.europa.eu/health/tobacco/events/index_en.htm#anchor0) (accessed 28 Nov 2012).
31 It is important to underline that the preferred policy options do not – in the assessment of the Commission – lead to increased illicit trade. On the other hand, illicit trade accounts already for 8,25% of the current consumption (Euromonitor data as presented in Matrix 2012).
OLAF, REGIO, RELEX, RTD, TAXUD, TRADE and TREN. The Group held 8 meetings: 5 March 2009, 26 November 2009, 2 July 2010, 18 April 2011, 3 October 2011, 1 March (information meeting and hand-out of draft Impact Assessment Report), 12 March 2012 and 19 July 2012. A meeting with associated services also took place on 30 November to present the legal proposal in the context of the inter-service consultation.

**Expertise and input**

A number of external studies have been commissioned to provide input to this impact assessment. First, a study on liability and the health costs of smoking was presented in December 2009. This study provided valuable input as far as the socioeconomic impacts of tobacco control policies are concerned. An updated version of this study was prepared in 2012. Second, a study assessing the impacts of revising the TPD was presented in September 2010. This report (by RAND Europe) was criticised by many stakeholders for its actual and perceived inaccuracies. In this respect, it is important to stress that this study has provided input to, but has not formed the exclusive basis of this impact assessment. The information was verified on the basis of other sources. Third, a study on novel and emerging tobacco, nicotine or related products was commissioned in 2010. Fourth, a study on the economics of the EU market of tobacco, nicotine and related products was commissioned in September 2011 to fill some remaining data gaps.

The Commission’s independent Scientific Committee on Emerging and Newly Identified Health Risks (SCENIHR) has presented two opinions relevant to the impact assessment: one on smokeless tobacco in February 2008 and one on additives in tobacco products in November 2010.

Finally, the Eurobarometer surveys conducted in October 2009 and February 2012 have been used to provide better insight on tobacco and nicotine use in the EU and on attitudes towards tobacco control policies. In particular the last Eurobarometer showed increased public support for the policy measures envisaged in this impact assessment.

**Invitations from European Parliament and the Council**

Since October 2007, the European Parliament has repeatedly called on the Commission to present a proposal for an amendment of the TPD and consider measures on ingredients and sales arrangements. A large number of questions on the TPD revision have also been received from the European Parliament during the past years.

The Council has twice invited the Commission to consider strengthening the tobacco control legislation and, in this context, to consider product related measures aimed at reducing the...
attractiveness and addictiveness of tobacco products and to analyse the legal issues and the evidence base for the impact of plain packaging, including its effect on the functioning of the internal market.  

1.4. **IMPACT ASSESSMENT BOARD**

A first version of this impact assessment report was submitted to the Impact Assessment Board (IAB) on 21 March 2012. On 18 April 2012 DG SANCO representatives had a meeting with the Board and the written opinion of the Board was received on 20 April 2012. The opinion concluded that the draft report required further work and asked for resubmission. A second version was submitted to the Board in June 2012. The second opinion of the Board, of 12 July 2012, did not request a resubmission but made some further recommendations on how to improve it. The opinion criticised in particular the evidence base for the policy areas "display at PoS" and "STP", which were subsequently dropped/amended. The table below sets out the main comments of the second opinion of the Impact Assessment Board and describes how they have been reflected in this revised version of the report.

<table>
<thead>
<tr>
<th><strong>Main comments from the IAB</strong></th>
<th><strong>Revision of the draft IA Report</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Better present the problem:</td>
<td>-Clearer references to effectiveness, implementation, enforcement and non-harmonised issues were introduced in the problem identification (2.2.1-2.2.5).</td>
</tr>
<tr>
<td>-Present separately problems related to effectiveness, implementation, enforcement and currently non-harmonised areas.</td>
<td>-Internal market justifications were better explained under the problem identification (2.2.1-2.2.5). The policy areas on PoS and TVM were discarded (4.1). The preferred policy option for STP was amended. The section on the legal basis (2.4.1) was reviewed.</td>
</tr>
<tr>
<td>-Demonstrate recourse to Article 114 for non-harmonised areas, in particular for limitation of promotion at the PoS.</td>
<td>-The references to equality were replaced by references to the objectives of Article 3 TEU (promote the well-being of its people) where relevant (for ex. 2.2). Health inequalities were, however, maintained as assessment criteria.</td>
</tr>
<tr>
<td>-Clarify compatibility between inequality in health and Member States' competences in defining their health policies.</td>
<td></td>
</tr>
<tr>
<td>2. Develop a robust baseline scenario:</td>
<td>-National actions in the context of FCTC have been further explained under the baseline scenario (2.3.2).</td>
</tr>
<tr>
<td>-Explain foreseeable national actions in the context of FCTC.</td>
<td>-The likely take-up of STP is described under the baseline scenario (2.3.3) and assessment of option 1(5.2.1). Circumvention is described in 2.2.1.</td>
</tr>
<tr>
<td>-Analyse further the likely take-up of STP, better explain circumvention possibilities of oral tobacco and acknowledge the uncertainty in relation to STP.</td>
<td></td>
</tr>
<tr>
<td>3. Better demonstrate the proportionality of policy options</td>
<td>-A clarification of the options status in comparison with the current TPD and FCTC has been introduced in a table on TPD and FCTC commitments in Annex 3 (3.2).</td>
</tr>
<tr>
<td>-Present measures going beyond current TPD and FCTC.</td>
<td>-A number of discarded options have been added to section 4.</td>
</tr>
<tr>
<td>-Discuss a wider range of discarded options.</td>
<td>-The preferred option on cross-border distance sales have been further justified (5.5.4). New preferred options have been identified for STP (regulation of chewing and nasal tobacco, notification of novel tobacco products and maintaining the ban on oral tobacco, see 5.2.1), NCP (all NCP above a certain nicotine threshold are subject to the medicinal products legislation, see 5.2.2).</td>
</tr>
<tr>
<td>-Justify STP, NCP and cross-border distance sale preferred options and alternative ban-options.</td>
<td>-The definition of traditional use in the glossary was revised, but no longer part of preferred option (5.2.1). The policy area on TVM was discarded (4.1). Traceability obligations in the context of FCTC were clarified in section 2.1.4 and Annex 3.</td>
</tr>
<tr>
<td>-Explain STP derogation, TVM verification and traceability obligations.</td>
<td></td>
</tr>
<tr>
<td>4. Improve the assessment of impacts:</td>
<td>-Divergences were better explained, for example under 5.4 on ingredients.</td>
</tr>
<tr>
<td>-Better explain persisting divergences, in particular on ingredients.</td>
<td>-Discounted values are presented in section 5.7.2 and Annex 5.</td>
</tr>
<tr>
<td>-Present discounted values.</td>
<td>-Administrative costs for internet notifications are better explained under 5.5.1.</td>
</tr>
<tr>
<td>-Assess administrative costs for internet notifications.</td>
<td></td>
</tr>
</tbody>
</table>

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2. **PROBLEM DEFINITION**

2.1. **MARKET DESCRIPTION**

1.1.6. **Tobacco products market**

   **a) Supply side**

   The total value of the tobacco market at retail level, including taxes and excise duties, is 136.5 bEUR and the market currently consists of five main categories of products\(^41\): 1) Factory manufactured cigarettes (FMC), 2) Roll-your own tobacco (RYO), 3) Pipe tobacco, 4) Cigars and cigarillos and 5) Smokeless tobacco products (STP) (oral, chewing and nasal tobacco). The value and volume of sales as well as manufacturing methods and consumption patterns differ significantly between the product categories.

   ![Figure 1: Comparison of relative market value of tobacco products in 2010](source: Euromonitor and industry estimates)

   **FMC** represent almost 90% (121.3 bEUR) of the total tobacco market value and despite a decline in sales volumes over the last ten years, the overall market value has increased consistently over the same period. This is mainly due to the tax increases and continuous development of premium brands sold at higher prices. FMC manufacturing is increasingly in the hands of four large multinational companies (PMI, BAT, JT and IT) accounting for around 90% of the EU FMC market.

   The FMC production is also highly concentrated in geographical terms, with only a few Member States (the United Kingdom, Netherlands and Germany) accounting for more than

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\(^{41}\) This categorisation is in line with the Commission's merger practice. It is important to underline that the notion of a market in this impact assessment is not based on competition law terminology.
50% of the total EU production.\textsuperscript{42} American Blend (using a mixture of Burley, Oriental and Virginia leaves) accounts for 76% of the FMC market.\textsuperscript{43}

The RYO market has increased significantly in recent years and accounts today for about 6.8% (9.3 bEUR) of the total tobacco market. This market is also characterised by strong presence of the four biggest FMC manufacturers (70% of the market), but to a lesser extent than FMC. The markets for \textit{pipe tobacco, cigars, and cigarillos} are considerably smaller. These products as well as chewing and nasal tobacco products are to a large extent manufactured by SMEs.\textsuperscript{44} The pipe tobacco and the cigar markets have been continuously declining in the last decade. The increase in cigarillo sales was partly driven by flourishing sales of so called eco-cigarillos which are not typical cigarillos and were recently re-classified as cigarettes.\textsuperscript{45} The STP market is dominated by oral tobacco (snus) and concentrated to Sweden, the only Member State where the marketing of this product is allowed.\textsuperscript{46} Chewing and nasal tobacco account for less than 0.1% of the total tobacco market and sales are concentrated to about 12 companies, mostly SMEs, who also sell other tobacco products.\textsuperscript{47}

Recent years have seen an increasing interest from bigger cigarette manufacturers to enter the STP market.\textsuperscript{48} Novel categories of tobacco products are currently being developed but they are not yet placed on the EU market.\textsuperscript{49} Most of these products are expected not to involve a combustion process and therefore would fall in the product category of STP.

Figure 2: Market value and volume 2000-2010

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>FMC</td>
<td>90.7</td>
<td>121.3</td>
<td>+33.8</td>
<td>793.7 bs</td>
<td>608.8 bs</td>
<td>-23.3</td>
</tr>
<tr>
<td>RYO</td>
<td>4.2</td>
<td>9.3</td>
<td>+123</td>
<td>53.1 tt</td>
<td>75.5 tt</td>
<td>+42.2</td>
</tr>
<tr>
<td>Pipe</td>
<td>0.576</td>
<td>0.480</td>
<td>-17</td>
<td>6.33 tt</td>
<td>4.03 tt</td>
<td>-36</td>
</tr>
<tr>
<td>Cigars/cigarillos\textsuperscript{50}</td>
<td>4.62</td>
<td>4.65</td>
<td>+0.6</td>
<td>7.81 bs</td>
<td>9.92 bs</td>
<td>+27</td>
</tr>
<tr>
<td>STP\textsuperscript{51}</td>
<td>0.48</td>
<td>0.83</td>
<td>+73</td>
<td>5,3</td>
<td>5.9</td>
<td>+10</td>
</tr>
</tbody>
</table>

Source: Euromonitor. Nominal prices

In terms of \textbf{market development}, a significant diversification of products has taken place in recent years. For example, the main manufacturer of oral tobacco (snus) increased its portfolio from 22 to 180 brands between 2002 and 2008,\textsuperscript{52} distinctive flavoured FMC such as "pina colada" and "chocolate" have been put on the market\textsuperscript{53} and FMC with new technology

\textsuperscript{42} Eurostat 2008/2009
\textsuperscript{43} Euromonitor 2010 (Matrix 2012)
\textsuperscript{44} European Smoking Tobacco Industry: Facts & Figures for DG Sanco, ESTA 2011-2012, ECMA, European Cigar Manufacturers Association, Facts and Figures
\textsuperscript{45} Directive 2010/12/EC on the structure and rates of excise duty applied on manufactured tobacco. Germany and Hungary have been granted transitional periods until 2014.
\textsuperscript{46} Marketing of oral tobacco is banned according to Article 8 of the current TPD and has been banned since 1992. Sweden and Norway were granted a permanent derogation from the ban received in their Accession Treaty; OJ C 241, 29.8.1994 (see article 151 and Annex XV thereof)
\textsuperscript{47} European Smoking Tobacco Industry Facts & Figures for DG SANCO, ESTA 2011-2012
\textsuperscript{50} This covers also "eco-cigarillos" exempted from the definitions for cigars and cigarillos through Directive 2010/12/EC on the structure and rates of excise duty applied on manufactured tobacco.
\textsuperscript{51} The data is limited to chewing tobacco in Denmark and Slovenia and oral and nasal tobacco (according to Euromonitor terminology snuff) in Germany, Sweden and Denmark
\textsuperscript{52} Swedish Match magazine Inside #2 from May 2008
\textsuperscript{53} Matrix 2012
including capsules filled with flavourings embedded in the filter have been introduced in many Member States.\textsuperscript{54} New market strategies have also been developed, in particular as a result of the tobacco advertising ban/restrictions in Member States, including innovative packaging and promotion at point of sale.\textsuperscript{55}

\textbf{Illicit trade} in FMC currently accounts for 8.25\% of total trade in the EU and is estimated to increase by 1\% per year in the next five years.\textsuperscript{56} Three broad categories exist: contraband, counterfeit and cheap whites/illicit whites (see glossary). Typically, these products do not comply with the safeguards of the TPD.

In terms of \textit{employment}, there were 48,500 persons employed in tobacco manufacturing in the EU in 2009.\textsuperscript{57} In addition to this, Eurostat data indicates that there are 86,113 farmers.\textsuperscript{58} In 2010, there were almost 100 first processors\textsuperscript{59} of tobacco in the EU located close to the growers’ areas, while the market of second processors\textsuperscript{60} is in the hands of two global players\textsuperscript{61} with similar market shares. 45,900 persons are employed in wholesale.\textsuperscript{62} The retail sector differs widely between Member States, but according to the European retailer association (CEDT), there are almost 990,000 premises selling tobacco in the EU and around 230,000 of these are specialised shops which usually generate 45-50\% of their turnover from tobacco.\textsuperscript{63}

As far as \textbf{raw tobacco} is concerned, the EU was a net importer of around 430,000 tonnes (about 2/3 of the quantity needed) in 2010.\textsuperscript{64} The EU tobacco production amounted to 294,000 tonnes. The production is often limited to small regions, very specialised and with large family labour requirements. Despite the fact that Virginia is the leading tobacco variety (46\% of the EU production), most of the EU tobacco farmers (81\%) are involved in labour intensive Burley and Oriental farming which is used in American blend cigarettes.\textsuperscript{65} According to tobacco producers, adding certain substances, including sugar, is indispensable for the use of these tobacco varieties. This is because these varieties lose their sugar content during the drying process whereas other varieties (such as Virginia) keep it.

A number of \textbf{upstream actors} other than farmers and processors are also involved in the tobacco manufacturing process. These include suppliers of ingredients, cigarette papers, filters

\textsuperscript{54}German Cancer Research Center (DKFZ). Menthol Capsules in Cigarette Filters – Increasing the Attractiveness of a Harmful Product. Heidelberg: DKFZ; 2012.
\textsuperscript{56}Euromonitor data presented in Matrix 2012. According to OLAF indications, the ratio between categories of illicit trade is 30\% contraband, 50\% counterfeit and 20\% illicit whites (for definitions, see glossary).
\textsuperscript{58}Economic stakeholders UNITAB and COPA estimate a number of 400,000 workers involved in growing in total (including family and seasonal working force.)
\textsuperscript{59}The first processors collect the raw tobacco cured by farmers and make a first process before selling it to the industry producing FMC, cigars or manufactured tobacco. DG AGRI
\textsuperscript{60}The second processors subsequently purchase, process, blend, pack, store and ship tobacco to meet each specifications of manufacturers of FMC and other tobacco products. DG AGRI
\textsuperscript{61}Alliance One Int. and Universal Corporation
\textsuperscript{62}Eurostat 2010. Bulgarian farmers represent 50\% of the EU tobacco farmers, followed by Poland and Greece (both 17\%)
\textsuperscript{63}Tobacco Retailers Figures. CEDT (Confédération Européenne des Détailleurs en Tabac). Sent to DG SANCO in January 2012
\textsuperscript{64}Eurostat 2010
and packaging (see also figure 3). Based on industry reported data, the total value of those supplies is around 3 bEUR.

Figure 3: Composition of a FMC

Source: Background to Cigarette Manufacturing and Use of Additives. Prepared by Philip Morris International Management S.A. 8 January 2010

The multinational dimension of the tobacco market combined with the geographic concentration of manufacturers and growers results in significant cross-border trade, both within the EU and with third countries. About 268,000 tonnes of raw tobacco were subject to internal trade between Member States in 2010 and the overall value of manufactured tobacco products traded across Member States' borders in 2010 was 9.5 bEUR. Figure 4 illustrates the main intra-EU flows in the production of American blend cigarettes.

Figure 4: Trade flows between tobacco-growing, tobacco-producing and tobacco-consuming countries

More than one third of tobacco products manufactured within the EU are sold across borders. Figure 5 illustrates intra EU trade in tobacco, following the categories used by Eurostat.

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66 Eurostat (Procom) 2008
67 Matrix 2012 based on information from the industry
68 As indicated in Annex 5, the total value of the FMC and RYO is 18 bEUR (ex-manufacture price). Almost 8 bEUR (see figure 5) is subject to intra EU trade.
Figure 5: Intra-EU trade in tobacco 2010

<table>
<thead>
<tr>
<th>Category</th>
<th>Value 2010 (bEUR)</th>
<th>Vol. 2010 (100kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cigarettes (containing tobacco)</td>
<td>6.591</td>
<td>3,427,689</td>
</tr>
<tr>
<td>Cigars, Cheroots and Cigarillos (containing tobacco)</td>
<td>0.644</td>
<td>68,641</td>
</tr>
<tr>
<td>Cigars, Cheroots, Cigarillos and Cigarettes of tobacco substitutes</td>
<td>0.004</td>
<td>1,864</td>
</tr>
<tr>
<td>Smoking Tobacco (whether or not containing tobacco substitutes in any proportion)</td>
<td>1.058</td>
<td>970,327</td>
</tr>
<tr>
<td>Manufactured Tobacco, Extracts and Essences, N.E.S.</td>
<td>0.279</td>
<td>639,322</td>
</tr>
<tr>
<td>Total Manufactured Tobacco (whether or not containing tobacco substitutes)</td>
<td>8.576</td>
<td>5,107,843</td>
</tr>
</tbody>
</table>

Source: Eurostat intra-EU trade figures

As regards trade between the EU and third countries, the total import to the EU of manufactured tobacco is less than 200mEUR, whilst the export of FMC and other manufactured tobacco products outside the EU accounts for 2.5 bEUR, i.e. a positive trade balance of 2.3 bEUR. On the other hand, import of unmanufactured tobacco (both raw and processed) counts for 2.17 bEUR, while the export accounts for 670 mEUR, what results in a trade deficit of 1.5 bEUR. For more details on the tobacco market and the manufacturing process, see Annex 2.

b) Demand side

According to the latest Eurobarometer 28 % of all EU citizens and 29 % of young people (aged 15-24 years) smoked in 2012. FMC is the most widely used tobacco product. 70% of the smokers start before the age of 18 and 94% under the age of 25. The overall smoking prevalence differs widely between Member States and ranges from 13% to 40%. Menthol cigarettes account in the EU, for approximately 4% of the EU market, ranging from 25% in Finland to 0.1% in Greece. Cigars and pipes are smoked far less than FMC and tend to be smoked occasionally rather than daily and mostly by older individuals. STP use is most common in Sweden, where around 12% of the population uses STP (mainly oral tobacco/snus) on a regular basis, compared to 2% or less in other EU countries where marketing of oral tobacco is banned, but chewing and nasal tobacco is allowed. The figure is higher if one considers all citizens who have at least tried STP. Chewing tobacco appears to have most users in Denmark and nasal tobacco is mainly used in Germany. Some industry players see a significant growth potential for STP as well as NCP, also in the light of smoke free environments. Figure 6 indicates the trends in smoking prevalence 2006-2012 and STP use 2009-2012.

Figure 6: Smoking Prevalence 2006-2012 and STP use 2009-2012

<table>
<thead>
<tr>
<th>STP</th>
<th>2009</th>
<th>2012</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regular Users</td>
<td>Na</td>
<td>1%</td>
</tr>
<tr>
<td>Proportion who have tried STP</td>
<td>EU 27</td>
<td>6%</td>
</tr>
<tr>
<td></td>
<td>DK</td>
<td>12%</td>
</tr>
<tr>
<td></td>
<td>FI</td>
<td>20%</td>
</tr>
</tbody>
</table>

69 Eurobarometer 2012
More than six out of ten smokers in the EU have tried to quit, with two out of ten in the previous twelve months. Personal health concerns are the main motivation to quit. While the number of smokers in the EU, and with this also the smoking prevalence, has decreased in the past decades, some Member States have seen an **upwards trend** in young people since 2005. The Health Behaviour in School-aged Children (HBSC) study of WHO Europe from 2012 indicates an increase in smoking prevalence in 14 Member States for 15 year old boys and in nine Member States for 15 year old girls.

According to the 2012 Eurobarometer, smokers and non-smokers commonly perceive FMC with lower tar or nicotine levels to be less harmful. The most important factor in choosing a FMC brand is the taste of tobacco (84%), followed by the brand itself (69%) and price (65%), but packaging (23%) and specific tastes (such as menthol, spicy, fruity or sweet) (32%) are also important. Peer group pressure/behaviour is obviously the most important factor for smoking initiation. EU citizens, including smokers, are largely and increasingly in favour of various tobacco control measures. For example, three out of four citizens support putting pictorial health warnings on all tobacco product packages. A third of smokers and ex-smokers also say that health warnings had an impact on their attitudes and behaviours towards smoking.

Most EU citizens who smoke or have smoked purchase regularly their cigarettes in a specialised tobacco shop (37%), from a newsagent (26%) a supermarket (22%) or a convenience store (20%). 10% of EU citizens purchase their tobacco from TVM (mounting to 15% in Member States where TVM were accessible). Only 10 EU citizens in 100 report that they have purchased tobacco products in a country other than their country of residence in the past 12 months. Cross border purchases via the internet are also limited (according to self-reporting). However, taking into account the increased use of e-commerce in other sectors, it appears likely that the market segment will grow.

### 1.1.7. The role of tobacco in the society

Tobacco is a legal product on the EU’s internal market, but is **no ordinary commodity** in the sense that it is the largest avoidable health threat in the EU, responsible for almost 700,000 deaths in the EU each year (see Annex 5). Moreover, millions of people in the EU suffer from one or more of the six main disease categories associated with smoking: 1) Bronchitis and other lower respiratory infections, 2) Chronic obstructive pulmonary diseases, 3) Stroke, heart attacks, arterial obstructions (especially in the legs) and other cardiovascular diseases, 4)  

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70 Eurobarometer 2012  
72 Eurobarometer 2012  
73 Idem.
Asthma, 5) Lung cancers and 6) Other cancers, such as pancreas, oesophagus, and stomach. Studies show that around 50% of smokers die prematurely and if they do so they die on average 14 years earlier. In addition, smokers have more life years that are characterised by serious disease.\textsuperscript{74}

In addition to measures at EU and international level, strong \textit{tobacco control policies} are pursued by Member States to address the harmful effects of tobacco and to protect public health. It is commonly agreed among tobacco control regulators that only a comprehensive set of continuously updated and adapted measures (e.g. price/taxation, smoke-free environments, advertising bans, labelling, ingredients regulations, information campaigns) is effective to reduce smoking prevalence over time, including reduced uptake among young people.

Tobacco consumption also has important \textit{economic implications} for society and is associated with important expenditures/costs. Annual EU public healthcare expenditure on treating smoking attributable diseases is estimated around 25.3 bEUR and society loses 8.3 bEUR per annum due to productivity losses (including early retirements/deaths and absenteeism) linked to smoking. In addition, if monetised, the life years lost due to smoking correspond to 517 bEUR every year. Figure 7 summarizes the costs associated with smoking in table format.

Figure 7: Costs associated with smoking in EU27 in mEUR

<table>
<thead>
<tr>
<th>Monetary value of life years lost</th>
<th>Smoking induced health care costs</th>
<th>Smoking induced early retirements</th>
<th>Smoking-induced absenteeism</th>
</tr>
</thead>
<tbody>
<tr>
<td>516,713</td>
<td>25,300</td>
<td>6,081</td>
<td>2,162</td>
</tr>
</tbody>
</table>

On the other hand, revenue from excise duty on the sale of tobacco product in the EU exceeded 79 bEUR in 2010, contributing to almost 3% of the total Government revenue.\textsuperscript{75} For more detailed information on economic impacts on the society, see Annex 5 (A5.2.3).

1.1.8. \textbf{Non-tobacco products}

\textit{Nicotine Containing Products (NCP)}

In addition to the traditional tobacco market, recent years have seen the emergence of new nicotine containing products (NCP) marketed primarily as consumer/leisure products. Electronic cigarettes appear to be the most commonly available type.

The EU \textit{electronic cigarette} industry is still quite fragmented and the absence of reliable trade statistics makes it difficult to summarise and analyse in terms of market size and value.\textsuperscript{76} However, it can be concluded that the market is growing rapidly. An electronic cigarette supplier has estimates that the current value of the German market is around 100 mEUR and that the total value of the EU27 e-cigarette market (including devices and refills) is between 400 and 500 mEUR.\textsuperscript{77} The Electronic Cigarette Industry Trade Association


\textsuperscript{75} EC, DG Taxud. 2010 For the purpose of this impact assessment, VAT has been excluded from the analysis as money not spent on tobacco will be spent on other goods attracting VAT. In this respect, any measure should be VAT neutral.

\textsuperscript{76} See Matrix 2012

\textsuperscript{77} Matrix 2012
(ECITA) estimates that they represent 60-70% of the volumes sold on the UK market and reports that the market is growing 20-30% monthly.\textsuperscript{78}

The EU market is mainly composed of distributors rather than manufacturers and dominated by small enterprises, although there is a growing interest from bigger cigarette manufacturers (including the big four FMC producers) to enter the market.\textsuperscript{79}

Most of electronic cigarettes are produced in China. Once imported into the EU, they are subject to significant cross-border trade. For example, in the Netherlands vendors of electronic cigarettes are reported to operate as a hub, reselling most of the electronic cigarettes they import from China to the rest of Europe. Around 20\% of their sales are internal to the Dutch market, while around 60\% are sold to German vendors and the remaining 20\% to vendors in Denmark, Spain, France, Austria and Switzerland.\textsuperscript{80}

\textbf{Figure 8: Value chain of the e-cigarette market in Europe}

\textsuperscript{80} Matrix 2012. Based on information from the Dutch e-cigarette consumer association.
As for the demand side, the current use of NCP is growing quickly. 7% of EU citizens have reported in the latest Eurobarometer that they have at least tried electronic cigarettes. In the UK an increase in the number of electronic cigarette owners has been estimated from a small number in 2006 to over 1 million by 2013.

Electronic cigarettes are widely advertised on the internet. A study monitoring Google search queries from January 2008 to September 2012 reported that online interest in electronic cigarettes has surpassed that of oral tobacco (snus) and nicotine replacement therapies.

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81 Eurobarometer 2012
Electronic cigarettes are most often marketed by their producers as an alternative to FMC rather than a smoking cessation aid. There are limited data available at this stage why people use electronic cigarettes. However, according to a recent survey among electronic cigarette users in Poland, most used the product primarily to quit smoking or to reduce harm associated with smoking (both 41%). An online survey of more than 3500 e-cigarette smokers found that the vast majority of respondents were using e-cigarettes to quit smoking or reduce their smoking (92%) and a large proportion felt these products were less toxic than traditional tobacco products (84%). These studies are well in line with information received from the Electronic Cigarette Industry Trade Association (ECITA), namely that the “vast majority of consumers” use e-cigarettes as a harm reduction alternative to smoking/a substitute for FMC/for smoking cessation purposes and that many use them to get around smoke-free environments (including lorry drivers). Due to the high numbers of possible flavours, ECITA has also reported that e-cigarettes would be used as a “fun product” (annual “vape festivals”). In terms of age of the users, ECITA has stated that statistics show that the proportion of users under 20 is low but they cannot exclude that young people or minors might use the products, although they believe that the comparatively high start-up costs, together with a lack of peer pressure would make this unlikely. The previously mentioned Polish survey also found that one in five young people had tested electronic cigarettes. As explained in subsequent sections of this report, the regulatory framework for electronic cigarettes and other NCP is complex and varies between Member States and a number of health and safety concerns are associated with the products. NCP other than electronic cigarettes appear, at this stage, to be less common on the EU market.

**Herbal products for smoking**

Euromonitor data indicate that herbal cigarettes were sold in significant amounts in seven Member States over the period 2000-2010. The overall size of the market grew from around 40.6 million units in 2000 to 50 million units in 2010, an increase of around 23%. These products are often marketed as "healthier" and natural products with "no additives", which gives the impression that these products are not harmful.

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83 Products marketed as smoking cessation aid would fall under the pharmaceutical framework and would need a prior authorization before being put on the market. ECITA explicitly advises its members to refrain from claims such as "quitting smoking" or satisfying cravings" (MOCK Audit Report shared with DG SANCO).
87 Idem.
88 Idem.
90 Matrix 2012
1.1.9. Regulatory framework

2.1.1.1. Tobacco control in the EU

Since the 1980s, there have been legislative initiatives in the EU to ensure harmonised product regulation for tobacco whilst also ensuring a high level of health protection. Today, the TPD constitutes, together with the Tobacco Advertising Directive from 2003, the key legislation in the field of tobacco in the EU. The content of the current TPD is described in section 1.2. The Tobacco Advertising Directive bans all forms of cross-border advertising in printed media, information society services, radio and sponsorships of events.

These two main Directives are complemented by two non-binding Council Recommendations: one on the prevention of smoking and on initiatives to improve tobacco control from 2003 and one on smoke-free environments from 2009.

Awareness-raising activities are also important instruments in EU’s tobacco control policy. The current campaign (Ex-smokers are Unstoppable) was launched on 16 June 2011 to encourage young adults in the 25 to 34 age group to stop smoking. The previous campaign, "HELP – For a life without tobacco", which ran from 2005 to 2010, was focused on smoking prevention, cessation, and passive smoking, targeting young Europeans between 15 and 25 years of age.

Measures to regulate and control tobacco are not only initiatives by DG SANCO. Tobacco is a cross-cutting issue which affects numerous policy areas. High taxes on tobacco products are generally seen as a very effective means to reduce tobacco use, with a particularly big impact on young people and people with lower incomes. Council Directive 2011/64/EU amends the structure and rates of excise duty on manufactured tobacco in the EU and in this context more convergence and further increases of tobacco taxes can be expected in the EU. The European Anti-Fraud Office (OLAF) investigates cases of illicit tobacco trade, which deprive Governments of significant tax revenues and undercut the prices charged by legal traders. In the area of advertising, the Audiovisual Media Services Directive complements the EU legislation on tobacco advertising, by providing a ban on all forms of audiovisual commercial communications, including product placement as regards tobacco. Direct tobacco subsidies to growers were once an important but controversial part of the EU’s agricultural policy. They have now been phased out, but were partially replaced by other subsidies. The Commission (DG EMPL) is also considering measures addressing the risks of employees arising from exposure to environmental tobacco smoke at the workplace.

The EU also works with international partners to reduce the consumption of tobacco worldwide, including in the context of the WHO Framework Convention on Tobacco Control (FCTC) (see 2.1.4.2).

All the instruments described in this section are mutually reinforcing and play an important role in the comprehensive tobacco control policy.

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94 OJ L 95/1 of 15.4.2010
2.1.1.2. Obligations in the context of the WHO FCTC

The WHO Framework Convention on Tobacco Control (FCTC) was adopted by the World Health Assembly in May 2003 and is the first ever international treaty on public health developed in response to the globalisation of tobacco consumption. The FCTC includes both demand reduction provisions (such as price and tax measures, protection from exposure to tobacco smoke, content and disclosure of tobacco products, packaging and labelling, education and communication, advertising, promotion and sponsorship) and supply reduction provisions (such as illicit trade, sales to and by minors and support for economically viable alternative activities).

The FCTC is a legally binding instrument which needs to be implemented and enforced by all Parties having ratified the Convention (so far over 170 countries across the world), including the EU and its Member States. At the time of the final approval of the FCTC it was considered that both the EU and Member States have competence in certain areas covered by the Convention (mixed agreement) and need to work together on the uniform application.95 A legally binding Illicit Trade Protocol based on Article 15 of the FCTC was adopted by the Conference of the Parties to the FCTC in November 2012. The EU and the Member States are expected to become Parties to this new instrument (as is the case for the FCTC). The core provisions of the protocol relate to the control of the supply chain for tobacco products through notably licensing (or equivalent approval), due diligence, record keeping, control of

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95 See Council Decision concerning the conclusion of the WHO Framework Convention on Tobacco Control (2004/513/EC), recital 4. See also Annex II thereof referring to Community competence in areas covered by Community legislation, including TPD.
duty free sales, of internet sales and of free zones and a tracking and tracing system. Seven sets of guidelines have also been adopted (by consensus) to assist Parties in meeting their implementation obligations under the FCTC (figure 11). The guidelines, while not legally binding, give an indication of Parties' political commitments under the FCTC. In general, the guidelines are more far reaching than the FCTC and reflect scientific developments in tobacco control. For the purpose of this impact assessment, a distinction has been made between the binding “FCTC obligations” and the political “commitments” contained in the guidelines.

For the purpose of the revision of the TPD, it is relevant to distinguish between the following categories of obligations and commitments stemming from the FCTC:

1. **Harmonised areas at EU level: obligations at EU level stemming from the legally binding FCTC:**
   The EU (as Party to the FCTC) has an international law obligation to ensure that the TPD provisions comply with the FCTC. Member States are bound by the TPD and cannot take action at national level. The Illicit Trade Protocol will also be a legally binding Treaty once adopted and entered into force. However it is not further considered in the context of this Impact Assessment.

2. **Harmonised areas at EU level: commitments at EU level stemming from the non-binding FCTC guidelines:**
   The EU should ensure that TPD reflects new developments based on scientific facts and internationally agreed product standards (TPD Article 11). Member States are bound by the TPD (where it allows no discretion to the Member States) and cannot take action at national level.

3. **Non-harmonised areas at EU level: obligations stemming from the legally binding FCTC:**
   The EU and its Member States are legally bound by the FCTC obligations and shall work together to ensure uniform application/avoid fragmentation of the internal market.

4. **Non-harmonised areas at EU level: commitments stemming from the non-binding FCTC guidelines**
   The EU and the Member States have expressed their (political) support for the FCTC guidelines, but have no legal obligations to implement them. EU action is justified if divergent implementation in Member States leads to (or is likely to lead to) fragmentation of the internal market.
Figure 11: Obligations in the context of FCTC in harmonised and non-harmonised areas

<table>
<thead>
<tr>
<th>1. TPD harmonised areas: Obligations stemming from the FCTC:</th>
<th>2. TPD harmonised areas: Commitments stemming from FCTC guidelines (GL):</th>
</tr>
</thead>
<tbody>
<tr>
<td>FCTC Art 11: Health warnings should be 50% or more, but not less than 30%</td>
<td>GL Art 11: Health warnings more than 50% - Pictorial health warnings on both sides of the packets - Removal of TNCO display</td>
</tr>
<tr>
<td>FCTC Art 15: Full traceability</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>3. TPD non-harmonised areas: Obligations stemming from the FCTC:</th>
<th>4. TPD non-harmonised areas: Commitments stemming from the FCTC guidelines (GL):</th>
</tr>
</thead>
<tbody>
<tr>
<td>FCTC Art 9 and 10: Effective ingredients regulation and measures for disclosure</td>
<td>GL Art 9 and 10: Restriction/ban on &quot;attractive&quot; ingredients - GL Art 11: Cessation information on the packages</td>
</tr>
<tr>
<td>FCTC Art 11: No misleading package promotion</td>
<td></td>
</tr>
<tr>
<td>FCTC Art 16: No small packages</td>
<td></td>
</tr>
<tr>
<td>FCTC Art 16: TVM not accessible to minors</td>
<td></td>
</tr>
</tbody>
</table>

Annex 3 gives a more detailed description of developments and plans in Member States as well as the links between the TPD and FCTC (A.3.2).

2.2. PROBLEM IDENTIFICATION

**Market developments**, such as the appearance of new products (e.g. electronic cigarettes) and new market strategies (e.g. appealing packages and flavoured tobacco products) have taken place in recent years. The same is true for **scientific development** (e.g. SCENIHR opinions). In terms of **international obligations**, the FCTC was concluded after the adoption of the current TPD, which was later supplemented by guidelines. In particular the guidelines leave Member States with substantial discretion in terms of implementation. This could lead to heterogeneous development in Member States and have a negative effect on the internal market, in general, and, in particular, in Commission legislative proposal in the field of internal market.

The lack of EU action negatively affects EU **citizens** in terms of premature mortality, expensive health care treatment and inadequate consumer information. This is not in line with the overall aim of the EU to promote the wellbeing of its people (TEU Article 3) nor with TFEU Articles 168(1) and 114(3) which foresee a high level of health protection in all EU policies and activities.

**Economic stakeholders** have referred to some shortcomings with the current TPD (e.g. on the lack of a common ingredients reporting format) and the tobacco industry stressed that they...
are not against a revision of the TPD if it improves the functioning of the internal market.\textsuperscript{96} The industry has also drawn the attention of the Commission to the risks associated with increased illicit trade. However, in general, the industry does not support further harmonisation, possibly in the fear of stricter regulation ("high level of public health"), although one company interviewed in the context of the revision stated that a single regulation on ingredients across EU might be less expensive than the continuous adaptation to national provisions, as long as the regulation is proportionate and science based.\textsuperscript{97} The tobacco industry, in particular Swedish Match, has also asked for the lifting of the ban on oral tobacco. They strongly oppose measures such as plain packaging and display ban at point of sale (PoS). The health NGOs, on the other hand called for stricter measures in particular with respect to labelling and ingredients. They requested that the ban on oral tobacco (snus) is not lifted and that stricter measures for chewing and nasal tobacco are introduced. They also called for strict regulation of NCP and supported measures against illicit/non-compliant products. For further details, see Annex 1.

1.1.1. Problem 1: Smokeless tobacco and extension of the product scope

\textit{a) Smokeless tobacco products (STP)}

Within the category of STP, the placing on the marketing of oral tobacco was banned in the EU in 1992 and this ban was maintained in the TPD in 2001 to stop the expansion of a product considered harmful to health, attractive to young people and new on the markets of all Member States.\textsuperscript{98} Before the ban, Member States had started to take individual action, which in 1992 was replaced by action at the Community level. The producers of oral tobacco argue that the ban on oral tobacco is no longer appropriate. The health NGOs and most Member States are of a different view. This section also raises the question whether the current labelling requirements for all STP (Art 5(4) TPD) and ingredients regulation (reporting) are still adequate/sufficient, taking into account the high number of new products which are now marketed/attractive to young people.

- **Effectiveness of the current TPD**

General problems related to all STP

The current regulation of STP does not effectively address recent and foreseeable market developments. When the current regulation on STP (including the ban on oral tobacco) was introduced in 1992 and confirmed in 2001, chewing and nasal tobacco products were seen as traditional products on a declining market with virtually no market outside certain socio-professional groups (seafarers, miners and sectors of the army) and regions.\textsuperscript{99} The situation has somewhat changed since the current TPD was adopted. Recent years have seen an increase in the sale of chewing tobacco as well as a modest increase in sale of nasal tobacco on the main markets concerned.\textsuperscript{100} Chewing and nasal tobacco products are subject to

\textsuperscript{96} Minutes from the meeting with FMC manufacturers, 2 December 2011: http://ec.europa.eu/health/tobacco/events/index_en.htm#anchor4 (accessed 28 Nov 2012).

\textsuperscript{97} Matrix 2012


\textsuperscript{100} Euromonitor data indicates an increase in the sale of chewing tobacco from 4 tons to 14.2 tons between 2000 and 2010 in Denmark. Sales of nasal tobacco in Germany saw a small decrease in the beginning of the century,
cross-border trade, including via the internet, meaning that they can reach new user groups and new geographical markets.\(^{101}\) However, one chewing tobacco manufacturer has reported a greater potential for growth as more restrictions are put in place for smoking tobacco products as STP provides an alternative to be used where smoking is not allowed.\(^{102}\) The production method (partly done by hand) of traditional chewing tobacco and the relative expensiveness of the products is also an important factor in determining the market potential.

In addition, there has been an important **product development** in STP (both as regards chewing and nasal tobacco and as regards oral tobacco limited to the Swedish market). As indicated in the market description (section 2.1.1), the main manufacturer of oral tobacco (snus) increased its portfolio from 22 to 180 brands between 2002 and 2008.\(^{103}\) New market strategies target consumers outside the distinctive population groups who traditionally used these products, including young people. For example, there are STP available which are especially developed for modern taste or a younger generation. STP with characterising flavours (including chewing tobacco with tropical or bergamot flavours, nasal tobacco with peanut butter or cheese and bacon flavour and oral tobacco with elderflower and rhubarb taste) are put on the market and nasal tobacco has recently been promoted at youth parties throughout Germany. The packaging and labelling has also become more attractive masking the health risks associated with these products. In this light some Member States have taken action and banned STP all together leading to discrepancy between national markets (see Annex 3). The accession country Iceland is also reflecting about the introduction of a ban on chewing and nasal tobacco.\(^{104}\)

**The current labelling requirements for STP (oral, nasal and chewing tobacco) do not adequately inform consumers** about the adverse health effects of STP. In particular, the current warnings lack visibility as they are required on one side of the package only and are often placed on the bottom of the packet.\(^{105}\) One of the countries (Norway) responding to the public consultation has argued that the current **health warning** on STP (including oral tobacco) is outdated, insufficient and needs to be strengthened. However, Norway cannot take action at national level due to the current TPD requirements (TPD Article 5(4)) which applies to this country through the EEA Agreement.\(^{106}\)

**In terms of health, there are considerable differences between various types of STP**, but all STP contain nicotine and are addictive. While according to a relatively recent SCENIHR opinion (2008) STP are less harmful than for example FMC, the report also concluded that STP are not harmless. This is in line with the assessment of IARC, that has classified STP as carcinogenic to humans. These products contain carcinogenic substances, albeit at different levels, and are associated with a number of adverse health effects. The health effects, but the trend has reversed in recent years, from 168 tons 2008 to 170 tons in 2011. (Matrix 2012). The trend in chewing tobacco is fully in line with data reported by one manufacturer of chewing tobacco. Other stakeholders involved in chewing and nasal tobacco were not able to provide the Commission with specific data showing the development of the market from 2000 to 2010 despite several reminders.

\(^{101}\) There are no exact figures on cross-border trade in STP trade available, but indications that such trade exists are available on a number of websites.

\(^{102}\) SANCO meeting with Oliver Twist, 19 July 2012.

\(^{103}\) Swedish Match magazine Inside #2 from May 2008


\(^{106}\) Comment on the Consultation on the possible Revision of the Tobacco Products Directive 2001/37/EC by the Norwegian ministry of Health and Care Services, December 2010.
including findings from more recent studies, are further developed in section 5.2.1, assessing the impacts of lifting the ban on oral tobacco.

Oral tobacco (snus)

There are divergent views among stakeholders whether the current ban of placing on the market of oral tobacco is justified and whether the ban should be maintained, extended to other STP or lifted. Economic stakeholders, in particular those producing oral tobacco (snus), like Swedish Match, and citizens responding to the public consultation have argued that the ban of oral tobacco is no longer justified considering that oral tobacco is less harmful than FMC and also less harmful than other STP (chewing and nasal tobacco) which are not banned. These stakeholders also claim that oral tobacco can be used for smoking cessation. Other stakeholders, including most Member States and health NGOs, have defended the current ban and/or argued for an extension of the ban to all STP. The main arguments from these stakeholders relate to concerns about the harmful health effects of STP, the risk of STP as a gateway to FMC consumption and the risk of dual use (i.e. consumption of both cigarettes and snus). It was also argued that all STP need to be banned while these products still have relatively limited market shares as the supply of novel forms of STP is likely to increase. Respondents within this category also pointed to the fact that STP cannot be seen as an effective substitute for FMC.

The main producer of oral tobacco in the EU, has also claimed that the different treatment of oral tobacco and chewing tobacco is discriminatory. The producer has pointed to the huge marketing potential for oral tobacco (snus) outside Sweden and indicated that the current ban is equal to a hypothetical annual loss in export revenues to at least 3 bEUR and in the most optimistic scenario the market could even reach 9 bEUR per year.107 The marketing potential for oral tobacco has increased significantly in recent years as STP is a means to consume tobacco/nicotine in public places where smoking is forbidden. Manufacturing of some other STP requires manual labour which significantly limits the growth potential. Also, this type of STP is produced by a very limited number of SMEs who do not have the market power to export into new markets where the products have no traditional use.

The authorities of the autonomous Finnish island Åland have argued in their submission to the public consultation on the revision of the Tobacco Products Directive (TPD) that the current ban on oral tobacco distorts competition. Ships under Swedish flag would be allowed to sell oral tobacco (snus), whilst vessels under the flags of Finland or Åland serving the same routes would not be allowed to do so.108

Novel tobacco products

The STP market development is not limited to conventional oral, chewing and nasal tobacco products. There are also important developments in other novel tobacco products, often claimed to be less harmful than FMC and targeted to consumers who cannot or will not cease

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107 Swedish Match power point, 15 December 2011
108 Any differential treatment, however, is an effect of the exemption rather than the ban. See the Courts reasoning in Case C-434/02 Arnold André GmbH & Co. KG v Landrat des Kreises Herford [2004] ECR I-11825, p 41
The tobacco industry seems to see a very significant potential in this new product development. Like for other the STP, the question is whether these products can act as entry gates into tobacco consumption, in particular among young people, or whether these products prevent established smokers from quitting (dual use). The expected development in consumption of STP in the absence of further EU action is explored under section 2.3 (Baseline scenario).

- Enforcement of the current TPD

The current regulation of STP based on the mode of consumption of different STP categories is sometimes unclear and can facilitate circumvention of the current legislation. In particular, it may encourage STP manufacturers to market their products as chewing tobacco (allowed) instead of tobacco to be sucked (banned as oral tobacco). For example, the instruction on how to use a product marketed as a chewing tobacco found on an internet store refers to the placing “under your upper lip and keep in place for 5 minutes to a few hours according to your convenience”, which describes a sucking and not a chewing mode of use.

It has been reported that the current ban of placing on the market of oral tobacco has also led to some enforcement difficulties. A recent study concludes that many internet retailers of Swedish oral tobacco (snus) target non-Swedish customers and that ordering oral tobacco from EU Member States other than Sweden is quick and straightforward despite the current ban. Out of 43 test purchases in 10 Member States, 41 were successfully made. The availability of oral tobacco through internet throughout the EU was also pointed out in the Commission report on Sweden's implementation of measures necessary to ensure that oral tobacco is not placed on the market in other Member States. Further action might be warranted on this issue upon receipt of supplementary information.

b) Nicotine containing products (NCP)

Electronic cigarettes and other NCP for leisure purposes did not exist when the current TPD was adopted in 2001. This raises the question whether there is a need for action and if so how these products should be treated.

- Implementation of the current legal framework for NCP

About half of the Member States have reported that they consider electronic cigarettes and other NCP as medicinal products by function, even if they are not presented as smoking cessation aid but rather as alternative to cigarettes (leisure products). One third have said they have no specific regulation in place which means that the General Product Safety Directive applies and a minority of the Member States have chosen to ban these products or apply certain provisions that are used for tobacco products (for further details, see Annex 3, A.3.1.1.2.). So far, no electronic cigarette has been authorised in the EU under the pharmaceutical regulation, but at least one application has been submitted and others are

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110 See for ex. Thompson C. BAT invests in a smokeless future. Financial Times 2012 Sept 30, where it is reported that the market for tobacco alternatives (including non-combustible cigarettes and nicotine inhalers) could count for as much as 40% of BAT's revenues – which were £15bn in 2011 – in 20 years time.

111 Peeters S, Gilmore AB. How online sales and promotion of snus contravenes current European Union legislation. Tob Control 2012.

112 COM (2010) 399 final Report from the Commission to the Council on the implementation of the Kingdom of Sweden of the measures necessary to ensure that oral tobacco is not placed on the market in other Member States. 29.7.2010.

The increasing market volume (section 2.1.3) and cross-border trade together with the different legislations in Member States affects negatively the functioning of the internal market. It contributes to legal uncertainty for manufacturers and distributors and may also negatively affect consumer confidence in the internal market, in particular considering the safety concerns related to NCP (see figure 12). It also prevents NCP from moving freely across borders as manufacturers and distributors are required to comply with many different legal systems. Member States have expressed an urgent need for orientation from the Commission as to which legislation applies to electronic cigarettes.

Nicotine replacement products (NRTs) considered as medicinal products need to undergo strict marketing authorisation procedures and if other NCP can reach the market without such authorisation, it could lead to an unjustified advantage undermining a level playing field. The pharmaceutical industry has argued in favour of regulating NCP (some of them in the context of the pharmaceutical legislation and others by including them in the scope of the TPD), while Electronic Cigarettes Industry Trade Association (ECITA) representing primarily vendors of electronic cigarettes in the UK has argued that these products do not need to be further regulated (see Annex 1).

The current fragmented situation is a result of various efforts in Member States to respond to health concerns associated with NCP. The Commission has, so far, received fourteen notifications concerning (refill) liquids for electronic cigarettes via the RAPEX system, indicating serious health risks for consumers (17 December 2012). The serious health risks were due to the toxicity of nicotine and misleading presentation, for example labelling referring to fruit.

Nicotine is a toxic and addictive substance. Acute nicotine poisoning has occurred in children who accidentally ingest nicotine and the safety of heavy or long terms use are not (yet) known. Cartridges are sometimes sold in containers with minimal protection against tampering, opening by children etc. Ingestion of a single replacement cartridge is very likely to be lethal and users have reported leakage when replacing cartridges, suggesting that the quality of cartridges themselves is highly variable. A study of five different brands of electronic cigarette also found that most brands' cartridges were poorly constructed and leaked.

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114 Information about one application. It has also been reported in media that at least one of the big FMC manufacturers is preparing for an application: Thomas R. Cigarette giant to offer ‘safer alternative.’ England and Wales: Nicoventures; 2011. http://www.nicoventures.co.uk/Content/Downloads/august-2011/PM_Aug_p2.PDF (accessed 28 Nov 2012).
117 Nicotine is classified as a dangerous substance in Directive 67/548/EEC.
Studies of the nicotine content of cartridges have shown significant differences between labelled and true levels of nicotine cartridges and refill solutions. Analyses of electronic cigarette samples conducted by the US Food and Drug Administration (FDA) have shown detectable levels of known carcinogens and toxic chemicals; including diethylene glycol, tobacco-specific nitrosamines and tobacco specific impurities. A recent study has found immediate adverse physiologic effects (changes in the lung function) after short terms use which is similar to some of the effects associated with tobacco smoking. Another recent study concludes that ‘passive vaping’ must be expected from the consumption of e-cigarettes due to prominent components in the gas-phase, including 1,2-propanediol, 1,2,3-propanetriol, diacetin, flavorings, and traces of nicotine.

Some NCP also appear to be subject to vivid and innovative marketing, which could attract young people in particular. For example, e-cigarettes are available in a range of flavours, including coffee and cherry and a "smart pack" is being introduced that vibrates and flashes a blue light when a user is within 50 feet of someone with another "smart pack". The popularity of electronic cigarettes was also confirmed in a recent Polish survey which found that one in five young people had tested electronic cigarettes. ECITA on the other hand claims that consumers are mainly established smokers using the product for cessation and reduction purposes. At this stage the evidence on the effectiveness of electronic cigarettes in smoking cessation is inconclusive. As indicated, most users of electronic cigarettes seem to use them for cessation/reduction purposes, but the industry association, ECITA, explicitly advises its members to refrain from such claims as it could trigger application of medicinal products regulation (see section 2.1.3.). There are also some concerns that electronic cigarettes and other NCP can take advantage from national smoke-free environment policies, in particular as the products are often promoted as an alternative to smoking which allows smokers to keep up nicotine addiction in situations where smoking is prohibited. Electronic cigarettes can also become a starter/re-starter product attractive to young people or former smokers. However

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there are some studies being published that highlight the electronic cigarettes' potential as a smoking cessation aid.  

- **Enforcement of the current legal framework for NCP**

The current uncertainty in relation to the legal classification of NCP makes *enforcement difficult and contributes to circumvention. It also undermines reaching a level playing field* and leads to an unjustified differential treatment of different types of products containing nicotine. Today, there are two categories of products containing nicotine available on the market: NCP placed on the market without any prior control and NRT (Nicotine Replacement Therapies) which have been subject to a strict (and relatively costly) risk/benefit analysis and approved as medicinal products.

As illustrated under the market description (section 2.1.3), most electronic cigarettes are presented as alternatives to FMC rather than smoking cessation aids. It appears that the products are presented in this way in order to avoid the relatively burdensome authorisation procedure applicable to medicinal products.

Regardless of the presentation of the NCP, a product which may be used to modify physiological functions falls under the definition of medicinal products by function and needs to be authorised under this framework before being put on the market. The pharmacological effects of nicotine in NRTs are well documented. Nicotine attaches itself to receptors in the brain and has a long tradition of use in NRTs to reduce craving and help people stop smoking.

Moreover, electronic cigarettes without prior authorisation appear to be available also in Member States which have reported that they consider them as medicinal products or prohibit marketing. According to ECITA, regulatory agencies struggle to provide monitoring and enforcement in regulating the electronic cigarette industry.

c) **Herbal products for smoking**

- **Fragmentation of the internal market**

Herbal products for smoking, e.g. tobacco- and nicotine-free cigarettes and herbal fillings of water-pipes, fall outside the current scope of the TPD and are currently not subject to harmonised rules in the EU. This raises the question whether these products – considering their similarities with tobacco products – should be regulated in the revised TPD. Herbal products for smoking are subject to many different regulatory regimes in Member States. About half of the Member States have no specific regulation in place, while two Member States do not allow these products, one Member State has labelling requirements in place and

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130 For example, ECITA recommends to its members to stay out of medical claims as this could render the product illegal (material provided to DG SANCO).

131 Nicotine patches, sprays, inhalers and chewing gums have already been authorized as medicinal products.

132 See for example Minutes of SANCO's meeting with ECITA on 20 June 2012 reporting about two members in Greece despite the ban on such products in Greece (see Annex 3: http://ec.europa.eu/health/tobacco/docs/ev_20120703_mi_en.pdf

the remaining Member States have other rules (see Annex 3, A.3.1.1.1). This negatively affects the cross border trade in these products, which appears to be quite significant considering that many products are marketed and sold on-line. Manufacturers and retailers need to be familiar with and keep up to date with many different legal situations in all Member States. The free movement of goods and consumer confidence in the internal market cannot be secured in this case as the health requirements differ significantly from one Member State to another.

The divergent actions taken by Member States are a result of health concerns related to these products. Herbal products for smoking have not been extensively studied from a scientific/health perspective but it is generally acknowledged that inhalation of smoke of any kind can pose a health risk. This is also the reason for Member States' actions in this area. Evidence suggests that the combustion of these products produces a level of carbon monoxide similar to tobacco cigarettes as well as other toxic substances such as tar. On the other hand, these products do not contain nicotine and may therefore not be associated with the same risk of addictiveness. It is of particular concern that these products are often perceived as harmless or less harmful by consumers. This perception is reinforced by the current marketing of herbal products for smoking as "healthier" and "natural products without additives".

A particularly worrying trend in this context is the increase in water-pipe (sisha) smoking among young people. The content of the water-pipe filling is often unclear and sometimes herbal filling is used which does not contain tobacco. Water-pipe smoking as such is also often perceived as a less harmful activity than traditional tobacco smoking. As indicated, this is expected to trigger further actions at national level in the years to come.

1.1.2. Problem 2: Packaging and labelling

The type, size and location of health warnings as well as the display of TNCO levels are already subject to harmonised provisions in the current TPD (see figure 12). However, it might be necessary to update these rules to bring them in line with international, market and scientific development.

- Effectiveness of the current TPD

The current provisions on packaging and labelling are no longer in line with scientific evidence and commitments in the context of the FCTC. For example, the current obligation to print the TNCO yields on the package have been shown to be potentially misleading, as it makes people believe that some products are less risky to their health.

134 No exact figures are available, but herbal cigarettes are sold on line
136 However, addictiveness may also be associated with other factors, including behavioral factors.
Scientific evidence also suggests that bigger pictorial pictures on both sides are more effective than text-only warnings on a range of outcomes, including being a deterrent for new smokers\textsuperscript{141} and a means to increase cessation among current smokers.\textsuperscript{142} In particular, they increase smokers' and potential consumers' awareness of warnings, knowledge and credibility of health risks, depth of processing and also cessation behaviours such as forgoing FMC, quit intentions and actual quitting.\textsuperscript{143} Thus, the current TPD provisions in this area are not as effective as they could be.

Figure 12: Comparison between harmonised TPD provisions and FCTC guidelines in the area of packaging & labelling

<table>
<thead>
<tr>
<th>Current TPD Articles 5 and 7</th>
<th>Article 11 FCTC and Guidelines for implementing Articles 11 and 13 FCTC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Text warnings not less than 30% + 40% of both sides (Art 5(5)). For Member States having more than one official language, the warnings should be increased to 32-35% and 45-50%.</td>
<td>Health warnings shall be large, clear, visible and legible and should be 50% or more of the principal display areas but not less than 30% (FCTC Art 11b(iii) and (iv)) Health warnings should be more than 50% of the display areas (GL Art 11)</td>
</tr>
<tr>
<td>Member States may introduce pictorial warnings on one side of the package (Art 5(3))</td>
<td>Parties should consider pictorial health warnings on both principal display areas. (GL Art 11)</td>
</tr>
<tr>
<td>Mandatory display of TNCO levels on the package (Art 5(1))</td>
<td>Parties should prohibit the display of figures for emission yields such as TNCO. (GL Art 11)</td>
</tr>
<tr>
<td>Ban on misleading product descriptions (Art 7)</td>
<td>Packaging, individual cigarettes or other tobacco products should carry no advertising or promotion, including design features that make products attractive. (GL Art 13)</td>
</tr>
</tbody>
</table>

Some of the current packet shapes make it difficult to effectively display health warnings affecting negatively the visibility and legibility of the warning. This is particularly the case for very narrow (including “lip-stick” shaped) packets which distorts text and picture warnings.\textsuperscript{144}

Use of colours and other design features on packages and tobacco products undermines the effectiveness of the ban on misleading descriptors in Article 7 of the current TPD.


\textsuperscript{144} Tan YL, Foong K. How the Malaysian tobacco industry exploits loopholes in pictorial health warnings. Tob Control 2012;21(1):55-6.
Since the adoption of the current TPD, misleading descriptors such as “light,” “mild” and “ultra” has been replaced by the use of colours which can be misleading and give the impression that some products are less harmful than others (for example gold and white is used to indicate 'light' FMC). Recent studies have demonstrated that packages have the potential to mislead smokers and potential consumers and present them with an erroneous comfort about the risk of smoking. For instance, FMC packets featuring the descriptors 'slim' or 'extra-slim' were rated significantly more appealing than packets without those descriptors. In another study of young adults, so-called 'super-slim' 'parfume type' FMC packages were associated with femininity, elegance, slimmness and reduced harm.

Likewise, the shapes (e.g. slim) and colours (e.g. pink, black, denim blue) of individual FMC can mislead consumers by creating e.g. the impression of harmlessness. A study found that smokers of 'slim' FMC were more likely to believe that some FMC could be less harmful and that their own brand might be a little less harmful. A recent study in young Australian adults has shown that characteristics of the cigarette stick affect smokers' perceptions of the attributes of cigarettes. Some packages make different types of health claims by conveying the impression that a product has health benefits as it contains fruits, vitamins or is associated with energy. Other packages claim that FMC contain “no additives” or are “natural”, which can lead to misperceptions that certain products are less harmful.

- **Enforcement of the current TPD**

Article 5(5) of the current TPD allows Member States to impose stricter labelling measures for domestically produced products, but Member States can only impose stricter rules on imported products.\(^{152}\) This can lead to market distortion and the industry could circumvent strict domestic measures by shifting production to other Member States.

- **Currently not fully harmonised aspects of packaging and labelling**

Some of the not fully harmonised aspects of packaging and labelling are subject to important disparities between national regulations. For example, pictorial health warnings are already in use in eight Member States, two Member States will use them from 2013 onwards while the remaining seventeen Member States for the moment require textual warnings only (see Annex 3). In addition, Member States apply different rules on the package size/minimum number of FMC per package and different rules on display of cessation services on the packages. The choice of pictures used also varies from one Member State to another. At least one Member State (France) bans promotional elements on the package as part of its advertising ban, whilst technical discussions are taking place in a few Member States as regards promotional elements, including FMC and smoke appearance.\(^{153}\) As described under section 2.3, the disparities are expected to grow in coming years.

The heterogeneous development in Member States is closely linked to the legal obligations and political commitments in the context of the FCTC (see figure 13). The FCTC provides a broad margin of manoeuvre for parties in terms of implementation, both in terms of scope and time.

**Figure 13: Political commitments stemming from FCTC guidelines in areas not fully harmonised by the TPD**

<table>
<thead>
<tr>
<th>FCTC and its guidelines (GL)</th>
<th>Situation in Member States</th>
</tr>
</thead>
<tbody>
<tr>
<td>Each Party should endeavour to prohibit the sale of FMC individually or in small packets. (FCTC Article 16)</td>
<td>Fourteen Member States specify a minimum package size of twenty FMC: AT, CZ, DK, EE, FI, FR, EL, IE, LU, LT, PL, PT, RO, ES</td>
</tr>
<tr>
<td></td>
<td>Four Member States specify a minimum package of nineteen FMC: DE, HU, NL, SE</td>
</tr>
<tr>
<td></td>
<td>One Member State has a minimum package size of 10: UK</td>
</tr>
<tr>
<td></td>
<td>One Member States requires FMC to be sold in packages of 10 or 20: IT</td>
</tr>
<tr>
<td></td>
<td>Two Member States prohibit the sale of single FMC: LV, SI</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parties should consider pictorial health warnings on both principal display areas. (GL Article 11)</th>
<th>Pictorial health warnings (on one side) in use in eight Member States: BE, RO, UK, LT, FR, MT, ES and DK (in IE and HU starting in 2013)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Health warnings and messages should in addition to harmful effects address advice on cessation (GL Article 11)</td>
<td>Four Member States have mandatory references to cessation services displayed on the packages: BE, FR, NL, SI</td>
</tr>
<tr>
<td></td>
<td>Nine Member States have references to cessation services included in some warnings: AT, DK, DE, HU, IE, LV, PL, SE, UK</td>
</tr>
</tbody>
</table>

\(^{152}\) Of course, if a measure is outside the scope of the current TPD Member States are entitled to impose stricter rules (even restricting cross-border trade), if the measure can be justified under Art.36 TFEU.

\(^{153}\) ITC Project. ITC France National Report. Waterloo, CA and Paris: University of Waterloo, Institut national de prévention et d’éducation pour la santé (INPES), Institut national du cancer (INCa), and Observatoire français des drogues et des toxicomanies (OFDT); 2009.
Parties should consider adopting measures to restrict or prohibit the use of logos, colours, brand images or promotional information on, packaging other than brand names and product names displayed in standard colour and font size (plain packaging) (GL Article 11) - Public consultation in UK. (16 April -10 August 2012) 154  
- Statement of BE Health Minister in the parliament. 155  
- Proposal by FR MP in report to the FR Health Minister (February 2012). 156  
- Informal discussions in FI.

Parties should consider adopting plain packaging requirements to eliminate the effects of advertising and promotion on packaging. (GL Article 13) 

Parties should prohibit or restrict ingredients that have colouring properties in tobacco products. (GL Article 9 and 10) - At least BE bans an ingredient capable of colouring the smoke blue (see Annex 3).

Discrepancies between national legislations in product related areas are liable in themselves to constitute obstacles to the free movement of goods. 157 In this concrete case, the diversity between national legislations obliges the tobacco industry to be familiar with and adapt to multiple national legislations and possibly produce different labels and different packages for different markets.

The fact that the industry has not explicitly requested the harmonisation should not be misunderstood to suggest that the problem does not exist. It could rather be a sign that the industry might expect a harmonisation at a higher level ("high level of public health"). SME's could be more open to harmonisation as they have less resources to adapt to diverging legislation and harmonisation would thus facilitate their ambition to expand their activities beyond their home markets. 158 At least one of four big tobacco manufacturers has also indicated that it has no major concerns regarding the introduction of mandatory pictures, TNCO replacement and cessation services displayed on the packages if appropriate space is given for trademarks. 159 Other manufacturers have been less explicit.

For Member States, the current lack of harmonisation represents a lost opportunity to fully benefit from common solutions across the EU as Member States will have to come up with their own solutions to address scientific, market and international developments. The divergent labelling schemes in Member States also mean an unnecessary burden for economic stakeholder in terms of compliance costs (e.g. familiarisation, multiple adaptations to national measures and different product lines for different countries) (see section 5.1).

157 C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd.[2001] ECR I-11453, p 64  
158 See section 5.1 and the reference to the statement of one company that ingredients legislation across EU might be less expensive than if actions are taken at Member State level, Matrix 2012.  
1.1.3. **Problem 3: Ingredients**

The current TPD foresees reporting obligations in a non-harmonised format and invites the Commission to establish a list of ingredients in form of a positive/negative list, which was not accomplished.

**a) Reporting**

- **Effectiveness of the current TPD**

Despite its mandatory character, the ingredients reporting set out in Article 6 of the current TPD is not working effectively. This was emphasised by economic (and other) stakeholders in the context of the public consultation of the TPD (see Annex 1) and also highlighted in the Second Application Report. Despite the publication of the EU guide on harmonised reporting format\(^\text{160}\) and the development of an electronic tool for data submission (EMTOC\(^\text{161}\)), different reporting formats are still used for reporting of ingredients. **This makes it difficult for manufacturers and importers to provide requested information and for Member States to fulfil the reporting obligations** set out in the current TPD. It also makes it difficult for the Commission to analyse the reported data.

Manufacturers also have **concerns about confidential business information** and there is no uniform and reliable basis to inform consumers on the content of tobacco products. Finally, disparities in the current reporting system makes it burdensome for Member States and the Commission to compare, analyse and draw conclusions from the data received.

**b) Ingredients regulation**

- **Currently non-harmonised aspects of ingredients**

In the absence of a common ingredients regulation, **Member States have adopted different approaches.** Four Member States have introduced positive lists indicating additives allowed to be used in tobacco products, one has introduced a negative list that restricts specific additives from being included in tobacco products and five have a combination. One Member State has maintained a voluntary agreement with the manufacturers (see Annex 3, 3.1.2.2). The EU has no regulatory power within the current TPD to harmonise these national lists and thus prevent obstacles to the internal market\(^\text{162}\). Most of these national regulations appear to be rather old and based on food legislation. However, there is a growing tendency in Member States to regulate ingredients, in particular as regards additives attractive to young people. One key example is the French regulation, fixing maximum levels for additives that impart a sweet or fruity/acid taste to FMC (e.g. vanilla). This has led to the reformulation or removal of certain products on the French market. In the aftermath, the same products were also altered in the other EU markets. Lithuania bans clove, vanilla plant and a number of other additives (see Annex 3.1.3.2). Another example is the different approaches as regards to additives added to the filter of FMC (e.g. menthol burst-capsule which allows a smoker to activate the menthol flavour whenever he/she wishes to do so). These new products are

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\(^{161}\) EMTOC (Electronic Model Tobacco Control) is a European web application which enables safe submission of the lists of tobacco ingredients to the concerned authorities. It was funded from the Health Programme by the European Executive Agency for Health and Consumers (EAHC) (60%) and a consortium of 13 Member States (40%).

\(^{162}\) Article 12 of the current TPD invites the Commission to propose a common list taking into account inter alia addictiveness.
allowed in a number of Member States, while authorisation has been refused in other Member States (Germany, Belgium) (see Annex 3, A.3.1.3). The lack of a harmonised approach on additives affects the functioning of the internal market and prevents free movement of products across the EU. Manufacturers have to produce different products for different markets.

As outlined under the baseline scenario (section 2.3), the heterogeneous development on the internal market is expected to increase further in coming years in the absence of a common EU approach. The expected development in Member States is not only based on the FCTC guidelines, but also on scientific progress and development of the market.

A significant number of scientific studies show that certain tobacco additives make FMC more appealing. The WHO Study Group on Tobacco Product Regulation summarises the international public health knowledge about flavourings added to FMC and other tobacco products and their attractiveness to young and older smokers. The Commission’s independent Scientific Committee on Emerging and Newly Identified Health Risks (SCENIHR) concludes, in its Opinion of 2010, that the use of fruit and candy flavourings in high amounts seems to favour smoking initiation by young people. It is also suggested that some additives decrease the harshness and increase the smoothness of the smoke. This also applies to menthol FMC, which show increased use in some Member States, e.g. Germany where the market share of menthol FMC more than doubled in the past ten years, from 1.3 to 3%.

In terms of market development, additives are added to tobacco products which will make them more palatable and capable of misleading consumers to believe that they are less harmful or have some beneficial effects. For example, spices and herbs can also be used to improve the palatability. As mentioned above, another recent development is FMC containing a burst capsule that enhances menthol (or other) flavour when squeezed and turn a FMC from non-menthol to a menthol FMC. Colouring agents are another way of enticing young people. Both pink and black coloured FMC are available on the market and a tobacco company recently wanted to use an additive which would colour the smoke blue. Various additives have been used in tobacco products to help create the misleading impression that such products have health benefits, present reduced health hazards or increase mental alertness and physical performance. For example, an oral tobacco product containing caffeine, taurine and guarana as well as baking soda claimed to help to keep the teeth white are available. 19% of EU citizens (smokers and non-smokers) believe that some FMC, e.g. those with menthol or other taste/flavour (spice, sweet, fruit) or those without additives or labelled as “organic” or “natural” are less harmful than others which is not the case. There is also a trend in several countries to use products labelled “without additives”.

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164 SCENIHR 2010
166 WHO FCTC guidelines for the implementation of Article 9 and 10.
168 Annex 3 (3.1.3). See also: De Standaard. Belgium. 5 December 2011.
169 WHO FCTC guidelines for the implementation of Article 9 and 10
170 Eurobarometer 2012
171 SCENIHR 2010
industry has expressed doubts whether a ban on certain fruit and candy flavoured products, such as pina colada and strawberry, would affect smoking behaviour, but showed some understanding for the need of further regulation.\textsuperscript{172}

Finally, it should be noted that tobacco additives also transform tobacco smoke into an even more complex chemical mixture and thereby may further increase the \textbf{carcinogenic and harmful effects of tobacco}. Additives that facilitate deeper inhalation (e.g. menthol) or inhibit the metabolism of nicotine may enhance the \textit{addictiveness} of nicotine indirectly.\textsuperscript{173} A review made of nearly 600 additives to FMC suggests that more than 100 are known to “have pharmacological actions that camouflage the odour of environmental tobacco smoke emitted from FMC, enhance or maintain nicotine delivery, could increase the addictiveness of FMC, and mask symptoms and illnesses associated with smoking behaviours.”\textsuperscript{174} The less aversive, cooler and milder smoke seems to improve the experience of smoking and facilitate smoking initiation. On the basis of an extensive review of all available information, the US FDA Tobacco Products Scientific Advisory Committee confirmed in 2011 that the evidence was sufficient to conclude that it is more likely than not that the availability of menthol cigarettes increases the likelihood of experimentation and regular smoking beyond.\textsuperscript{175} However, as mentioned previously (section 1.2) international work on ingredients has, so far, focused on attractiveness and the FCTC guidelines adopted in November 2010 do not address toxicity and addictiveness. However, work has been initiated in the areas of addictiveness and toxicity and it is expected that guidelines will be developed for these issues in coming years.

In the absence of a harmonised approach under the current situation, Member States are faced with \textit{unnecessary administrative burden} in terms of finding their own solutions on how to implement the FCTC guidelines and adapt national legislations to the above mentioned market and scientific developments. They cannot fully benefit from the economy of scale associated with one common solution across the EU and enforcement of national rules is becoming more burdensome. In addition, \textit{economic stakeholders are/would be faced with unnecessary compliance costs} in terms of country specific familiarisation, reformulations of tobacco products and different production lines for different Member States (see section 5.1).\textsuperscript{176}

1.1.4. \textbf{Problem 4: Cross-border distance sales}

Cross-border distance sales of tobacco products fall outside the scope of the current TPD. According to Council Recommendation 2003/54, Member States are recommended to restrict tobacco distance sales for general retail. The FCTC guidelines for implementing Article 13 recommend Parties to ban internet sales. The Protocol on Illicit Trade recommends regulating cross-border internet sales.

\textbf{Cross-border distance sales of tobacco implies a risk of circumvention of the safeguards of the TPD}. Typically, tobacco products sold on the internet do not comply with the provisions of TPD, which means that consumers do not benefit from safeguards of the TPD (e.g. health warnings and ingredients). Legal cross-border internet sale of tobacco products

\begin{footnotesize}\begin{enumerate}
\item SCENIHR 2010
\item Matrix 2012. Summary of industry responses to questionnaire.
\end{enumerate}\end{footnotesize}
also makes very little sense as taxes have to be paid, by the vendor, in the country of the buyer (i.e. no tax savings can be made by the buyer). Given the characteristics of the tobacco market, most tobacco products are also available on the domestic market. If the taxes in the country of destination are not paid consumers gets access to this product below the price level considered appropriate by the country of destination.

- Circumvention of the TPD and problems related to enforcement

The lack of effective common rules on cross-border distance sales of tobacco products increases the volume of **tobacco products circulating on the internal market without complying with the TPD provisions** (e.g. on health warnings, ingredients and ban on the placing on the market of oral tobacco). This is particularly the case under the current situation where internet sale of tobacco products is expected to increase in coming years (see section 2.1.1) and also taking into consideration that legal cross-border internet sales of tobacco products make very little sense as described above.

The **difficulties to enforce the ban of placing on the market of oral tobacco** are further described in section 2.2.1.

In a report from 2009 on the implementation of the Recommendation 2003/54, the Commission also concluded that, despite the overall satisfactory transposition, the area of **distance sale of tobacco represent particular challenges and problems** as far as enforcement is concerned.178

In addition to the non-compliance with the TPD provisions, many of the sites offering tobacco products on-line do not have information as regards **sales to minors** and have no system in place to verify the age of the purchaser.179 In this context, it should be noted that age limits, between 16 and 18 years, for purchasing of tobacco are in place in all Member States (see Annex 3, A.3.1.4).

Access to tobacco on the internet is therefore easier and cheaper than from other sources. Internet sales of tobacco also often imply illegal advertising.180

As illustrated in Annex 3 (3.1.4), nine Member States do not allow internet sale of tobacco while some others have no restrictions in place.181 Among those Member States, some (including France and Lithuania) have introduced outright ban on this type of sale. Others (including Austria, Bulgaria, Hungary, Latvia and Spain) only grant licences or permission to sell tobacco through other sales channels. Several other Member States have different restrictions in place as far as internet sale of tobacco is concerned. The restrictions range from licensing of the internet retailer, to age limits and advertising bans. The different regulatory schemes in Member States and the lack of any common rules make it difficult for Member

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177 Article 36 of Directive 2008/118 on excise duty of tobacco indicates that in the case of cross border sale, the excise duty have to be paid for in the country of destination. However, from a perspective of a consumer, cross-border purchase makes primarily sense when the consumer avoids the higher excise duties in the country of destination.


180 Peeters S, Gilmore AB. How online sales and promotion of snus contravenes current European Union legislation. Tob Control 2012.

181 AT, BG, ES, FR, HU, IT, LT, LV and SK do not allow internet sale of tobacco.
States to enforce their national legislation. This is particularly the case given the cross-border dimension of this type of sales and the fact that the economic operators (retailers) compete on the same virtual market and offer their products to consumers regardless of their locations.

- Impact on the internal market

As described above, illegal cross-border distance sales of tobacco undermine the application of the provisions contained in the TPD, e.g. on health warnings and ingredients.

The steady increase in (illegal) cross-border internet sales also reduce the legal movement of goods both within and between Member States and makes it more difficult for legal business (traditional retailers) to compete on the market.

1.1.5. Problem 5: Traceability and security features

- Implementation of the current TPD

Illicit trade in FMC currently accounts for 8.25 % of total trade in the EU and is estimated to increase by 1 % per year in the next five years.\(^\text{182}\) The negative impact of illicit trade is manifold for the legal supply chain, consumers and Member States. From the perspective of TPD, the main concern is that the safeguards of the TPD are not respected by illicit products (e.g. health warnings in the correct language). For Member States, the loss in tax revenues (10 bEUR per year according to OLAF)\(^\text{183}\) is certainly also a major concern. The main beneficiaries are criminals and the organised crime. The main categories of illicit products are contraband (i.e. products which have been diverted into illicit trade, not respecting the legal requirements in the jurisdiction of destination), counterfeit (i.e. brand protected products which have been falsified without consent of the brand owner and are not respecting the legal requirements in the jurisdiction of destination) and illicit/“cheap” whites (i.e. products produced (often legitimately) in their country of origin at very low cost, destined to be smuggled into other jurisdictions and not respecting requirements in the jurisdiction of the destination). According to OLAF indications, the ratio between these categories is 30 % contraband, 50 % counterfeit and 20 % illicit whites.

The current TPD contains, as part of its Article 5(9) the possibility to introduce provisions on traceability and product identification, but so far no effective implementation has taken place. In particular, the Commission has not used the power in Article 5(9) to adopt technical measures related to traceability and identification. A major reason for this delay was that since 2000 the concepts of traceability have been discussed internationally in the development of Article 15 FCTC and later on in the negotiations of the Illicit Trade Protocol based on Article 15 FCTC. It would therefore not appear useful if the Commission made use of the powers under Article 5(9) TPD at this stage, as the current provision does not provide for a fully-fledged traceability concept and is thus no longer state of the art. The currently foreseen batch marking is only one element for achieving traceability and control of the supply chain. Already the FCTC goes further and stipulates that each unit package has to be marked in order to determine the origin and the point of diversion and to monitor, document and control

\(^{182}\) Euromonitor data presented in Matrix 2012. According to OLAF indications, the ratio between categories of illicit trade is 30 % contraband, 50 % counterfeit and 20 % illicit whites (for definitions, see glossary).

the movement of tobacco products and their legal status. As regards Article 5 (9) TPD, in the absence of the adoption of an implementing measure, Member States are, in principle, unable to take actions unilaterally, while being obliged by the FCTC. Taking into account that a very significant part of FMC and RYO are traded across EU borders (2.1.1), an EU wide system would clearly provide added value.

- **Enforcement of the TPD**

  Typically, illicit trade products (both contraband and counterfeit) do not comply with the TPD. This undermines the enforcement of the Directive and thus the effectiveness of the internal market in legal tobacco products. Sales of products not complying with TPD mean that the consumers do not benefit from the safeguards in form of the appropriate labelling (e.g. text warnings in a different language), or ingredients control.

  In terms of traceability, there is currently no level playing field for economic operators on the internal market. The four largest tobacco manufacturers have concluded legally binding agreements with the EU and the participating Member States (‘the agreements’), including provisions on tracking and tracing, but other manufacturers and importers do not have similar legal obligations to control the supply chain through tracking and tracing.

  It is highly likely that Member States will also consider additional measures in terms of security features to act against counterfeit and empower consumers to verify the authenticity of the product concerned. Article 15 FCTC contains obligations in this regard, but without a common EU approach Member States are expected to take different approaches. **Such heterogeneous development will have a negative impact on the internal market and prevent free movement of products.**

**2.3. BASELINE SCENARIO**

This section describes how the tobacco market is expected to evolve in the coming years if no changes are made to the TPD. In this respect it is first important to underline that the current difficulties in terms of implementation, application and enforcement of the current TPD (see section 1.2) would persist under the baseline scenario.

**1.1.6. Development of the market**

As mentioned in the market description (section 2.1) significant development and diversification have taken place on the tobacco market in recent years. This development towards further fragmentation of the internal market is expected to continue under the baseline scenario.

The rapid development of the traditional **STP (oral, chewing and nasal) market** is likely to continue under the baseline scenario (new brands, new flavours, new attractive packaging). The development of novel tobacco products is also a priority for FMC manufacturers and this market is expected to increase significantly. In addition, the development of tobacco-free

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products, such as NCP (notably electronic cigarettes) described in the market description section, 2.1.3) is expected to continue and even intensify. One of the main drivers for this trend is the implementation of smoke-free environments and tobacco control measures in the Member States.

Also the new marketing strategies for tobacco for smoking are expected to continue. This applies in particular to innovative and appealing tobacco packaging (e.g. new "lip stick" packets with slim FMC). The development is even expected to aggravate under the baseline scenario, especially following stricter advertising regimes in Member States and taking into consideration that packaging constitutes an important factor in choosing a FMC brand (23%).

Moreover, further market development as regards ingredients for combusted tobacco is expected to continue in coming years, in particular in light of the recent development of distinctive flavoured tobacco products (see section 2.1.1). The various patents required by tobacco industry to embed flavourings in the filter are also an indication of further development in this regard. Increased sale of menthol FMC can also be expected under the baseline scenario and these products are likely to be most popular among young people. The market share of menthol FMC has more than doubled in Germany in the past ten years, from 1.3 to 3%. In the United States, menthol FMC sales have remained stable as cigarette sales have decreased and the share of menthol brands is higher in new adolescent smokers, who have been smoking for less than one year, than for those who have smoked for more.

1.1.7. Development in Member States

2.3.1.1. Already harmonised areas

In areas already covered by harmonisation, Member States would be prevented under the baseline scenario from taking actions to maintain or strengthen a high level of health protection.

2.3.1.2. Non-harmonised areas

In reaction to the market developments outlined above, some Member States have already taken actions and more are likely to follow (see sub-sections under 2.2 and Annex 3). Member States will also adapt to scientific progress and international development, e.g. implement the FCTC and its guidelines.

Although the FCTC provides good orientations, it leaves significant discretion to Member States on how to implement the Convention and guidelines in terms of content and time. For example, the Convention refers to health warnings of a certain size (50% or more but not less than 30%) and effective ingredients regulation. However, the Convention itself does not

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187 Eurobarometer 2012


specify how and where to place the warnings on the packets, nor which ingredients should be addressed by the regulation. The guidelines are a bit more specific and more far reaching, but also they give some discretion to Member States and they are not legally binding. Whilst some Member States are expected to take actions to implement fully also the non-binding guidelines, others are likely to be less ambitious in this respect. The legal obligations stemming from FCTC are also different from obligations based on an EU Directive in the sense that the enforcement mechanisms are different. Whilst a failure to implement an EU Directive may lead to infringements procedures, settlement of the disputes in the context of FCTC leaves more flexibility to Parties (FCTC Article 27 refers to negotiations or any other peaceful means of their choice). Also this can affect the implementation of the FCTC. In light of this, further discrepancies can be expected under the baseline scenario.

More specifically, in the areas of **STP, NCP and herbal products for smoking**, it can be expected that Member States continue to take unilateral actions to address the market development described above. For example, some Member States can be expected to react to the development of flavoured STP and implement the FCTC guidelines on Articles 9 and 10 which recommend banning additives capable of making these products more attractive or of misleading consumers to believe they are less harmful or have health benefits. Some Member States are also likely act to ensure maximum effect of their smoke-free environment policies. Although it is difficult to predict the exact shape and timing of these actions, it appears certain that they will consist of divergent approaches reinforcing the already existing discrepancies.

Also in the area of **packaging and labelling**, the disparities are expected to grow in coming years as Member States continue to take further measures, e.g. to adopt pictorial health warnings, introduce cessation information and/or further standardise tobacco packaging in line with the guidelines for implementing Articles 11 and 13 of the FCTC. In the context of the public consultation on the TPD, five Member States who have not yet adopted pictorial health warnings nevertheless supported such a proposal at the EU level. It is likely that at least some of them will go forward at national level in the absence of a common EU approach. The UK Government launched a public consultation on plain packaging on 16 April 2012 and appears to be the most advanced in terms of considering the full standardisation of the package, but reflections on plain packaging are also on-going at various levels in some other Member States, including Belgium, Finland and France (see Annex 3, A.3.1.2.2). Member States' interest in plain packaging is also reflected in a Council Recommendation from 2009 inviting the Commission to analyse the legal issues and the evidence base for the impact of plain packaging, including its effect on the functioning of the internal market.

In order to comply with their obligations under FCTC (see Annex 3, A.3.1.3.1), Member States have already different measures in place and are likely to adopt rules on **ingredients** unilaterally in the years to come, in particular to address the attractiveness of tobacco products and considering the market development in recent years with more flavoured tobacco products being placed on the market. At least one Member State has declared that it will discuss a national ban on “candy-cigarettes” if no action is taken at EU level. A majority of

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191 AT, FI, LV, PL and SI.
194 DK political agreement between the Government and the Alliance of 21 April 2012
Member States also supports some kind of regulation in the context of the public consultation (see Annex 1, section A1.2.2). It is also likely that Member States will seek inspiration from other jurisdictions. A number countries around the world have taken measures (e.g. Canada and Brazil banning in principle all additives and US banning characterising flavours).  

Finally, Member States are expected to further develop their legislation in order to address easy access to and visibility of tobacco, including through the internet. In this case the FCTC plays an important role as well as the enforcement difficulties faced by Member States under the current situation (e.g. age limits for purchasing of tobacco).

1.1.8. The demand side

The overall smoking prevalence has decreased significantly in the EU in recent decades. Whilst Euromonitor predicts also some moderate decline in consumption (decrease in volume sales) in the years to come it cannot be assumed that this predicted decrease will in reality continue in the absence of new adjusted tobacco control measures. Firstly, the decline was the effect of a concerted and comprehensive tobacco control policy consisting of a broad range of measures used in a complementary manner and where new elements have been constantly introduced. This includes the adoption of the FCTC, increases in tobacco taxation, bans on tobacco advertising, restrictions on sales to minors, comprehensive laws on smoke-free environments, health warnings on tobacco packages, awareness raising campaigns as well as measures adopted in other policy areas (see section 2.1.4). In the absence of further tobacco control measures at EU level, it is likely that the trend in prevalence would revert, at least in those Member States not taking actions under the baseline scenario. In any case, even if some Member States take stronger actions on their own initiative, this would create a patchwork of legislations interfering with the internal market.

Secondly, market developments in terms of packaging, ingredients, new products and sales strategies have the potential of misleading consumers, undermining awareness of health risks and encouraging (in particular) young people to take up smoking. This has been demonstrated in the problem identification above (section 2.2.1-2.2.5) for each of the relevant policy areas, including references to scientific evidence. Over time, this could encourage young people to take up smoking and prevent current smokers from quitting. Whilst the latest Eurobarometer showed a welcome decline in smoking prevalence amongst young people, studies from the WHO showed that several Member States have seen an increase in smoking prevalence among youth since 2005.

Based on this it is assumed that the overall smoking prevalence will remain at the current level if no EU action is taken. The precise prediction of the baseline is, however, of limited relevance for this impact assessment as the impact of all measures is expressed in relative terms, i.e. if the consumption/prevalence were to decrease as predicted by Euromonitor, the

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195 Canada has an exemption for menthol and Brazil accepts sugar up to a certain level. US bans cigarettes with characterising flavours other than tobacco or menthol, including strawberry, grape, orange, clove, cinnamon, pineapple, vanilla, coconut, licorice, cocoa, chocolate, cherry or coffee.
196 According to 2012 Eurobarometer survey, smoking prevalence was 28% in the EU-25 (excluding RO and BU) and 27% in EU-15. This is considerably less than 32% in 2006 (EU-25), respectively 39% in 2002 (EU-15). The volume of the EU-27 cigarettes market in 2010 was 608.8 bn sticks, what represents a decline of 23.3% in comparison to 2000, when 793.7 bn sticks were sold across the EU.
197 Matrix 2012.
proposed measures would accelerate the decrease. If, on the other hand, the consumption were to remain stable, it could decrease thanks to the envisaged measures.

**Consumption of STP is expected to increase under the baseline scenario** in light of new product developments and the introduction of smoke-free environments in more and more Member States.¹⁹⁹ This prediction is also shared by one of the chewing tobacco manufacturers who sent information to the Commission in the context of the revision of the TPD. The same manufacturer also confirmed that multinational tobacco companies' engagement in the STP segment is likely to lead to greater awareness of STP in general. Moreover, reports from other jurisdictions also suggest an upwards trend in STP worldwide. Use of nasal tobacco in Iceland has increased significantly from 11.7 tons in 2003 to 30.2 tons in 2011.²⁰⁰ New STP products have also been introduced on the US market, including dissolvable STP resembling candies.

There is not conclusive evidence as regards the substitution between STP (including novel non-combusted tobacco products) and smoking products and it is therefore not possible to draw any firm conclusion whether the expected increase in STP use will have an impact on the smoking prevalence. STP can, indeed, be used by current smokers, e.g. to reduce consumption or in any effort to quit, but at the same time and as illustrated above, STP can attract young people who do not yet smoke and who do not intend to take up smoking (entry gate). STP can also prevent people from effectively quitting (dual use). The issue of substitution will be further explored in section 5.1.1.

**The lack of visible health warnings** on STP under the current TPD (see 2.2.1) could also undermine consumers' and potential consumers' awareness of the adverse health effects of STP and thus lead to higher uptake of these products.

### 2.4. EU BASIS TO ACT

This section includes a general assessment of the relevant legal basis, subsidiarity and proportionality. The proportionality is further examined as part of the assessment and comparison of policy options (section 5).

Compliance with fundamental right has also been ensured throughout the document. The proposal affects the freedom of economic operators to conduct business (Article 16) as well as their freedom of expression and information (Article 11) and right to property (Article 17) but the obligations imposed on manufacturers, importers and distributors of tobacco products are necessary to improve the functioning of the internal market while ensuring a high level of health protection.

### 1.1.9. Legal basis

The current TPD is based on Article 95 TEC (now Article 114 TFEU).²⁰¹ The choice of the legal base has been confirmed by the Court of Justice of the European Union (hereinafter: "the Court").²⁰² The same legal basis is appropriate for revising the TPD. Article 114(1)

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¹⁹⁹ Despite repeated efforts to gather reliable statistics from the industry (European Associations and larger producers such as Pöchl) no information could be obtained.

²⁰⁰ Information received from the Icelandic Ministry of Welfare, July 2012.

²⁰¹ In addition to Article 95 TEC the TPD was also adopted on the basis of Article 133 TEC. Case: C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd. [2001] ECR I-11453, the Court found, however, that Article 95 TEC was the only appropriate legal base, but that the addition of Article 133 TEC as a legal base was not a reason for declaring the Directive invalid.

²⁰² C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd.[2001] ECR I-11453
TFEU empowers the European Parliament and the Council to adopt measures for the approximation of the provisions laid down by law, regulation or administrative action in Member States which have as their object the establishment and functioning of the internal market. According to Article 114(3) TFEU, the Commission should aim at ensuring a high level of health protection in its proposal envisaged in paragraph 1 of Article 114.

It follows from the case law that measures adopted on the basis of Article 114 TFEU must genuinely have as their object the improvement of the conditions for the establishment and functioning of the internal market. A mere finding of disparities between national rules is not sufficient to justify having recourse to Article 114 TFEU. At the same time it is settled case-law that recourse to Article 114 TFEU as a legal basis is possible if the aim is to prevent the emergence of future obstacles to trade resulting from divergent development of national laws. The emergence of such obstacles must be likely and the measures in question must be designed to prevent the discrepancies. The (existing or likely) disparity must have (actually or probably) the effect of creating an obstacle to trade, by preventing a product or service from moving freely within the Union, or by appreciably distorting competition on the internal market.

The Court has held that, provided that the conditions for recourse to Article 114 TFEU are fulfilled, the legislature cannot be prevented from relying on that legal basis on the ground that public health protection is a decisive factor in the choices to be made. On the contrary, priority must be given to measures ensuring a high level of health protection.

In line with the jurisprudence of the Court, the following situations of relevance for this impact assessment can be distinguished, under which harmonisation based on Article 114 is justified:

1. There is an **existing level of harmonisation** which needs an update in light of scientific and international developments. This is the case for display of TNCO levels, the size of the warnings and certain aspects of traceability features. The current legislation needs to be updated to take into account new evidence.

2. For **product specific measures** the decisive question is whether there are discrepancies between national legislations impacting on the marketing of products across borders or whether there is a sufficient likelihood that such discrepancies will appear in the future. Measures related to scope, labelling and ingredients typically fall within this category. It follows from jurisprudence that these types of product

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206 C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd.[2001] ECR I-11453, para 62

207 Idem, para 77-79
requirements are in themselves liable, in the absence of harmonisation at EU level, to constitute obstacles to the free movement of goods. 208

3. Even a measure not directly aimed at improving the conditions for the functioning of the internal market can be adopted under Article 114 if its purpose is to ensure that certain provisions concerning the internal market are not circumvented. 209 This is particularly relevant for the areas on cross-border distance sales and tracking, tracing and security features in this impact assessment.

4. A ban of a product can also, under specific circumstances, be adopted on the basis of Article 114. 210 In the case of oral tobacco, the Court confirmed the validity of the ban taking into consideration that national legislations were divergent and were therefore such as to constitute obstacles to the free movement of goods, the product (oral tobacco) was new on the market and could be attractive to young people. The Court also found the ban on oral tobacco proportionate due to the harmful effects, the uncertainty of oral tobacco as a substitute for cigarettes, the addictive and toxic characters of nicotine, oral tobacco's danger to young people and the novelty of the product. 211

1.1.10. Subsidiarity
In accordance with the principle of subsidiarity, the Union shall act only if and in so far as the objectives of the proposed action cannot be sufficiently achieved by the Member States, either at central level or at regional or local level, but can rather be better achieved at Union level (Article 5(3) TEU).

As indicated above, some of the areas included in this impact assessment are already covered by the current TPD, but need to be updated in accordance with market, scientific and international developments. Due to the harmonisation which already exists, Member States are prevented / limited from acting unilaterally. For example, Member States are not allowed to remove the display of tar, nicotine and carbon monoxide (TNCO) levels or choose to put the pictorial health warning on the most visible side of the package and they have limited competence in terms of traceability. Only a common approach is possible to update these provisions.

Other areas relevant for this impact assessment are subject to different approaches in Member States which have led to obstacles to the functioning of the internal market. For example, in policy areas labelling and ingredients, the heterogeneous situations in Member States have resulted in a situation where the industry has to produce different product lines for different markets. Even more disparity is expected in coming years when Member States implement the FCTC obligations and commitments. Only a harmonised approach at EU-level can remove obstacles to cross-border trade. Only a harmonised approach would ensure that industry is not obliged to adapt at different times to 27 national regimes.

Finally, in some areas it would be very difficult for a Member State to act unilaterally due to the difficulties to enforce such an action when other Member States have different rules. For example, it appears almost impossible for a Member State to enforce restrictions on

208 Idem, para 64
209 Case C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd.[2001] ECR I-11453, p 82-83
210 For example Case C-434/02 Arnold André GmbH & Co. KG v Landrat des Kreises Herford [2004] ECR I-11825
211 Case C-434/02 Arnold André GmbH & Co. KG v Landrat des Kreises Herford [2004] ECR I-11825
tobacco internet sales if such sales are unregulated in other Member States. The current Council Recommendation 2003/54/EC does not indicate how tobacco distance sale should be regulated and the challenge related to enforcement was outlined in the Commission’s Staff Working Document from 2009 on the implementation of the Recommendation. A legally binding and EU wide measure in the framework of the revision of the TPD would therefore produce clear benefits. The same holds true for the EU system for tracking and tracing, when tobacco products regularly move across borders.

EU action would also contribute to greater consistency, both between and within Member States, and a higher level of legal certainty, for example in the area of NCP where the legal situation is complex and unclear, which undermines the level playing field.

1.1.11. Proportionality

Under the principle of proportionality, the content and form of the Union action shall not exceed what is necessary to achieve the objectives of the Treaty (Article 5(4) TEU). In the area of health, the Court has held that the EU legislature must be allowed broad discretion and the legality of a measure can be affected only if the measure is manifestly inappropriate having regard to the objective which the competent institution is seeking to pursue. The revision shall aim at providing an appropriate level of margin for implementation by the Member States and it must fully respects responsibilities of the Member States to organise, finance and deliver health services and medical care.

3. POLICY OBJECTIVES

Overall objectives

The overall objective of the revision is to improve the functioning of the internal market, while ensuring a high level of health protection. In terms of internal market, the proposal aims to:

1. Update already harmonised areas to overcome Member States' obstacles to bring their national legislations in line with new market, scientific and international developments.

2. Address product related measures not yet covered by the TPD insofar as heterogeneous development in Member States has led to, or is likely to lead to, fragmentation of the internal market.

3. Ensure that certain provisions of the TPD are not circumvented by placing on the market of products not compliant with the TPD.

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212 See C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd.[2001] ECR I-11453, p 123
213 Without an update, Member States cannot, for example, increase the size of the health warnings, change their location of the package or replace the display of tar, nicotine and carbon monoxide levels. For example, at this stage, eight Member States have adopted pictorial health warnings and the regulations of ingredients differ between Member States.
214 For example, at this stage, eight Member States have adopted pictorial health warnings and the regulations of ingredients differ between Member States. For example, measures on cross-border distance sales and traceability will facilitate legal activity and thus prevent sale of tobacco products not complying with the TPD (e.g. health warnings and ingredients).
It is also important to ensure a **harmonised implementation of international obligations** following from the WHO Framework Convention on Tobacco Control (FCTC), which is binding for the EU and all Member States, and a consistent approach to non-binding FCTC commitments, if there is a risk of diverging national transposition.

The revision will contribute to the overall aim of the EU to promote the well-being of its people (TEU Article 3). The proposal should contribute to the Europe 2020 strategy in so far as keeping people healthy and active longer will have a positive impact on productivity and competitiveness. The revision should also fully respect the EU Charter on Fundamental Rights.

**Specific and operational objectives**

<table>
<thead>
<tr>
<th>Specific objectives</th>
<th>Operational objectives (PA=Policy Area)</th>
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<tbody>
<tr>
<td><strong>A</strong></td>
<td><strong>To remove obstacles to cross-border trade and ensure a level playing field for manufacturers and other economic operators</strong></td>
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<tr>
<td>1. remove unjustified differential treatment between products (PA1a, b, 3)</td>
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2. facilitate a level playing field for economic actors in the field of STP, NCP and herbal products for smoking (PA1), for retailers involved in cross-border distance sale (PA4) and traceability and security features (PA5) |
| 3. remove national disparities and ensure a harmonised approach in packaging & labelling and ingredients (PA1, 2,3) and traceability and security features (PA5) |  
| **B** | **To reduce the administrative burden for economic actors and public authorities due to the complexity of the current TPD and remaining disparities in legislations** |
| 1. unify the rules on labelling and ingredients and establish one single format for ingredients reporting (PA1, PA2, PA3) |  
2. facilitate market surveillance of Member States and improve the overall enforceability, including by reducing the number of products on the market which do not comply with TPD specifications (PA 1a, 3, 4, 5) |

**Choosing between options**

A high level of **health protection** has been considered when choosing between different policy options. In this context, the revision seeks to regulate tobacco products in a way that reflects their characteristics and negative impact on health, e.g. by ensuring that the ingredients and packaging of the products do not encourage or facilitate initiation by young people. 216 The focus on young people is also reflected in the selection of the policy options and the products primarily targeted (FMC, RYO and STP). In addition, the revision should create conditions which allow all citizens across the EU to take informed decisions about the products, based on accurate information on the health consequences of consuming tobacco products. Finally, all smokers should benefit from measures contained in the TPD (e.g. health warnings and ingredients regulation). 217

<table>
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<tr>
<th>Specific considerations</th>
<th>Operational considerations (PA=Policy Area)</th>
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215 For example, measures on cross-border distance sales and traceability will facilitate legal activity and thus prevent sale of tobacco products not complying with the TPD (e.g. health warnings and ingredients).


217 An unintended, but welcome side effect of the measures against illicit trade is that it protects tax revenues of Member States.
To provide a high level of protection to citizens throughout the EU

1. regulate the placing on the market of hazardous and potentially hazardous products (PA1, 3)
2. remove from the market products which are particularly attractive, in particular to young people, because of their appearance or taste/smell (PA1, 2, 3)
3. assist consumers to verify the authenticity of tobacco products and protect them against non-complying supply (PA5)
4. inform the consumer, through the labelling, about the harmful effects of tobacco and related products and remove misleading information (PA1, 2)
5. reduce easy availability and access of tobacco and related products in the interest of protecting vulnerable groups, in particularly young people (PA1a, c, 4, 5)
6. ensure that consumers across the EU benefit from a minimum level of protection when purchasing tobacco products (e.g. health warnings and ingredients control) and reduce the appeal of cheaply available illicit tobacco products to protect vulnerable groups. (PA4, 5)

4. **POLICY OPTIONS**

A number of factors have been taken into consideration when identifying the policy options, most prominently the benefits for the internal market and its main stakeholders (industry, suppliers, distributors, consumers, NGOs and Governments), international obligations and commitments (notably FCTC and its guidelines), and the EU Charter on Fundamental Rights. The views put forward by stakeholders have been carefully considered. Reference is also made to the criteria contained in Annex 4. Some general comments are necessary upfront.

**The role of public health**

While all options identified seek to improve the functioning of the internal market, the protection of public health has played a key role in designing/shaping the policy options. This follows from the Commission’s duty to ensure a high level of health protection in its proposals and is logical considering that the national legislations, which the revised TPD seeks to harmonise, have been adopted in order to protect public health.

**The relation to FCTC**

The table in section 3.2 of Annex 3 sets out the relation between the current TPD obligations, obligations stemming from FCTC, commitments in the context of FCTC guidelines as well as the preferred policy options.

**The products primarily targeted**

The policy options identified in this impact assessment cover in principle all tobacco products. However, in the policy areas on packaging & labelling (PA2), ingredients (PA3), and traceability & security features (PA5), stricter rules will - in a first stage - apply only to FMC, RYO and STP and other products will be exempted. The products initially exempted will continue to be subject to current TPD requirements. This means that the identified options primarily target large manufacturers of FMC and RYO and exempt the SMEs and micro-enterprises involved in the manufacturing of pipe tobacco and cigars. 218 This reasoning is based on a proportionality assessment, taking into account the interest of these economic stakeholders, their marketing mainly towards adult consumers and the focus of this report on regulating tobacco products in such a way that they do not appeal to underage citizens.

218 It should be noted, however, that also in the FMC, RYO and STP segment, some smaller manufacturers will be affected by the proposal.
However, for policy area 5 (traceability and security features) the exemption is only of a temporary nature (longer transitional period) and for the other concerned policy areas the exemption will be removed if the consumption trend changes and consumption in young people increases significantly.

The level of harmonisation

In general, the policy options foresee full harmonisation, but obviously Member States are allowed to maintain or introduce stricter national measures in areas not covered by the harmonisation provided that national rules are compatible with the Treaty, in particular with the principle of free movement of goods. The Directive also foresees the possibility for Member States to maintain or introduce stricter measures in areas falling within the scope of the Directive provided that the measures are notified to and approved by the Commission which will assess if they are necessary, proportionate and non-discriminatory taking into account the high level of harmonisation achieved by the Directive.

Delegated and implementing powers

Delegated and implementing powers are foreseen to allow for an appropriate reaction to market, scientific and international developments, to amend or supplement the basic act and to give effect or shape the rules laid down in the basic act. The exact scope of these two categories of acts is further defined in the legislative proposal and the nature of the power is clearly defined in each case and linked to strict conditions in order to ensure legal certainty and respect the institutional division of tasks. Delegated/implemented powers are foreseen in the following situations:

- Establish rules for the use of health warnings, unique identifiers and security features and adapt health warnings to technical and scientific developments.
- Establish further rules for the shape and size of unit packets.
- Setting/adapting maximum yields for emissions and their measurement methods.
- Establish the format of ingredients reporting.
- Remove from the market of tobacco products with characterising flavours and products with increased toxicity or addictiveness levels.
- To remove the exemption for products other than FMC and RYO as regards the labelling and other than FMC, RYO and STP as regards ingredients.

Subsequent impact assessments for implementing measures and delegated acts will be carried out as appropriate and in line with the Impact Assessment guidelines.219

4.1. DISCARDED POLICY AREAS

A number of additional policy areas were considered at an earlier stage of the impact assessment, but these were subsequently rejected.

Firstly, regulation of toys and sweets resembling tobacco was considered at one stage. This suggestion has not been retained in the impact assessment as it was concluded in an early stage that the TPD is not the most adequate instrument to deal with these products.

Secondly, the introduction of manufacturer liability for the financing of all health costs arising from tobacco consumption was proposed by the European Parliament in its

resolution from 2007. Full implementation of the "polluter pays principle" was assessed in the external study produced by RAND Europe in September 2010.\textsuperscript{220} An external study on liability and the health costs of smoking was also commissioned to examine this proposal in more detail.\textsuperscript{221} However, the proposal has not been assessed further in the context of this impact assessment in the light of subsidiarity and legal considerations.

Thirdly, a **total ban of all tobacco products** was discarded without further analysis. Despite the harmful effects of tobacco consumption, this option was not considered feasible. A total ban on tobacco products would lead to unreasonable compliance costs and not have the desired effect of stopping the use of tobacco in the EU. An illegal market would most likely appear, in particular considering that 28% of EU citizens currently smoke.

Fourthly, the approximation of Member States' legislations on **Tobacco Vending Machines (TVM)** was considered in the impact assessment process and public consultation. This policy area was discarded given subsidiarity concerns and taking into account the already good progress in this area following the Council Recommendation 2003/54 and the FCTC provisions and guidelines. Thirteen Member States have banned TVM, while all remaining Member States have put in place some kind of restrictions to limit access to young people under the legal age for purchasing.\textsuperscript{222} For more information about FCTC obligations and commitments as well as national legislations see Annex 3 (A.3.1.4.2).

Fifthly, the approximation of Member States' legislations in the area of **tobacco display at point of sale (PoS)** was considered in the impact assessment process and the public consultation. Also this policy area was discarded due to subsidiarity concerns as well as the limited support from Member States at this point in time. For further information on international commitments and regulations in Member States, see Annex 3 (A.3.1.4.2).

### 4.2. PROBLEM 1A – SMOKELESS TOBACCO PRODUCTS (STP)

For STP, the main issues are how to effectively regulate the products (including oral tobacco, chewing and nasal tobacco as well as novel non-combustible tobacco products), which develop rapidly (labelling and ingredients) and which are addictive products with adverse health effects (although less harmful than FMC), taking into account the growth potential (smoke-free environments)) and the current differential treatment between oral tobacco (snus) and other STP. The following options were considered:

- **Option 0: No change**

Status quo means:

a) Oral tobacco is banned (except for Sweden which has an exemption but needs to respect labelling requirements, i.e. health warnings, and ingredients reporting).

b) Placing on the market of chewing and nasal tobacco is allowed subject to labelling requirements (health warning) and ingredients reporting.

c) Placing on the market of novel forms of STP is allowed subject to labelling requirements (health warnings) and ingredients reporting.

- **Option 1: Lift the ban on oral tobacco and subject all STP to stricter labelling and ingredients regulation**

\textsuperscript{220} Rand 2010
\textsuperscript{221} GHK 2012
\textsuperscript{222} Some of the national systems might be confronted with enforcement challenges.
a) The current ban on oral tobacco is lifted and these products can be placed on the market subject to stricter labelling and ingredients requirements (e.g. health warnings on both sides of the package and a ban of products with characterising flavour (glossary) and increased toxicity or addictiveness).223

b) The placing on the market of chewing and nasal tobacco continues to be allowed, but subject to the same rules on labelling and ingredients as set out under a.

c) Placing on the market of novel tobacco products (glossary) continues to be allowed subject to the same rules on labelling and ingredients as set out under a.

Delegated/implementing power to adapt health warnings, act on products with characterising flavours on products with increased toxicity or addictiveness and to regulate additives that cause a characterising flavour.

- **Option 2: Maintain the ban on oral tobacco, subject all novel tobacco products to a notification obligation and all STP to stricter labelling and ingredients regulation**

a) The current ban on oral tobacco is maintained (except for Sweden which has an exemption), but oral tobacco needs to comply with stricter labelling and ingredients regulation (e.g. health warnings on both sides of the package and a ban of products with characterising flavours and increased toxicity or addictiveness).

b) The placing on the market of chewing and nasal tobacco continues to be allowed, subject to the same rules as set out under a. A clearer definition of chewing tobacco is inserted in the TPD.

c) A notification obligation is introduced for novel tobacco products (glossary) and a report on the market development in these products will be issued by the Commission five years after the transposition of the TPD. Novel tobacco products placed on the market must respect the rules on labelling (health warnings on both sides) and ingredients regulation (ban on products with characterising flavours).

Delegated/implementing power as in option 1.

- **Option 3: Maintain the ban on oral tobacco, restrict the sale of other STP to areas of traditional use only and subject all STP to stricter labelling and ingredients rules**

a) The current ban on oral tobacco is maintained (except for Sweden which has an exemption), but oral tobacco needs to comply with stricter labelling and ingredients regulation (e.g. health warnings on both sides of the package and a ban on products with characterising flavours and increased toxicity or addictiveness).

b) The placing on the market of chewing and nasal tobacco is banned unless traditionally used (glossary) in Member States. The placing on the market of traditionally used STP is limited to the relevant territory/Member State and must comply with the same labelling and ingredients requirements as set out under a. Member States will have to prove traditional use and notify to the Commission.

c) The placing on the market of novel tobacco products is banned.

Delegated/implementing power as in option 1.

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223 The product regulation developed under this option could be inspired by the voluntary industry standards already used for snus, the Gothia Tek standards. Rutqvist LE, Curvall M, Hassler T, Ringberger T, Wahlberg I. Swedish snus and the GothiaTek standard. Harm Reduct J 2011; 8:11.
• **Option 4: Ban all STP with the exception of oral tobacco in Sweden which would be subject to stricter labelling and ingredients rules**

a) The current ban on oral tobacco is maintained (except for Sweden which has an exemption)

b) The placing on the market of chewing and nasal tobacco is banned given their circumvention potential, in particular as regards chewing tobacco.

c) The placing on the market of novel tobacco products is banned.

• **Discarded options:**

An alternative option would have been to lift the ban of placing on the market of oral tobacco without suggesting any regulation. However, this option was not considered viable as it would mean lifting a ban for a product with adverse health properties without adequate control/limits. This option was not suggested by any of the stakeholders.

Another option which was discarded was to introduce a new authorisation regime for novel tobacco products. Apart from the very significant burden for the evaluating and authorising bodies (either at national or EU level) and possibly new structures to be set up, more information is needed about these products as well as expected overall health effects, including on non-smokers, young people and people who would otherwise had quit tobacco consumption altogether.

4.3. **PROBLEM 1B – NICOTINE CONTAINING PRODUCTS (NCP)**

The main issue is how to effectively regulate NCP considering the heterogeneous development in Member States. Also, the addictive nature of NCP, health and safety concerns and uncertainty, growth potential (smoke-free environments) and market development (labelling and ingredients) need to be considered.

• **Option 0: No change**

Status quo means: The TPD remains limited to tobacco products. NCP remain subject to the General Product Safety Directive,\(^{224}\) or other more specific legislations.

• **Option 1: Subject NCP to labelling and ingredients requirement under TPD**

NCP placed on the market are subject to adapted health warnings, ingredients reporting and a prohibition to place on the market NCP with characterising flavours (glossary). Delegated/implementing power to adapt health warnings and act on products with characterising flavours, on products with increased toxicity or addictiveness and to regulate additives that cause characterising flavours.

• **Option 2: Establish a new authorisation scheme for NCP**

Only NCP that have been authorised under a new authorisation procedure (risk/benefit analysis) set up under TPD are allowed to be placed on the market. Otherwise, placing on the market of NCP is prohibited. The authorisation procedure would also cover labelling and additives control.

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• **Option 3: Subject NCP over a certain nicotine threshold to the medicinal products' legislation and the remaining NCP to labelling requirements**

NCP with a nicotine level over a certain threshold may only be placed on the market if they have been authorised as medicinal products on the basis of their quality, safety and efficacy, and with a positive risk/benefit balance under the medicinal products legislation. NCP with nicotine levels below this threshold will be subject to an adapted health warning. The nicotine threshold identified under this policy option should be established by considering the nicotine content of medicinal products (NRTs) for smoking cessation which have already received a market authorisation under the medicinal products' legislation.

Delegated/implementing power to adapt the health warning and the identified nicotine threshold taking into account scientific and technical developments.

• **Option 4: Subject all NCP to the medicinal products' legislation**

Only NCP that are authorised as medicinal products on the basis of their quality, safety and efficacy, and with a positive risk/benefit balance are allowed to be placed on the market. Otherwise, the placing on the market of NCP is prohibited. The authorisation procedure is described under option 3.

• **Discarded option**

Development of minimum safety standards for NCP under the General Product Safety Directive was not considered to address the health problems identified. Safety standards developed under this Directive and referenced in the Official Journal provide that products complying with the standards are presumed to be safe. However, since nicotine is toxic and addictive, it is impossible to set safety standards for products releasing nicotine. A safety standard under the General Product Safety Directive could therefore not fulfil its objective.

Another option would have been to update the orientation note on electronic cigarettes from 2008, but considering the non-binding character of such a document and the need for a clear legal framework, this option has been discarded.

4.4. **PROBLEM 1C – HERBAL PRODUCTS FOR SMOKING**

The main concern related to herbal products for smoking is the different regulatory approaches in Member States. Also, the misperception of adverse health effects needs to be taken into account.

• **Option 0: No change**

Status quo means: TPD remains limited to tobacco products. Herbal products for smoking remain unregulated or subject to different national regulations.

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• **Option 1: Subject all herbal products for smoking to labelling requirements under TPD**
  Adapted health warnings are required for herbal products for smoking. Delegated/implementing power to adapt the warnings.

• **Option 2: Phase out the placing on the market of herbal products for smoking**
  Placing on the market of herbal products for smoking is phased out.

• **Discarded option**
  An alternative option would have been to regulate the content of these products. However, this option was discarded as not relevant for the main problem identified.

4.5. **PROBLEM 2 – PACKAGING AND LABELLING**

The main issues are that some of the current provisions are outdated (e.g. size of the warnings, display of quantitative TNCO-values) and that there is heterogeneous development in Member States (e.g. pictorial warnings). There is also a need to implement FCTC obligations and commitments, to address the potential of packaging and labelling to mislead consumers and encourage people to start or maintain smoking.

• **Option 0: No change**
  Status quo means: Current labelling rules are maintained, i.e. a general text warning of not less than 30% and an additional text warning of not less than 40%. Member States can choose to use a combined warning (picture and additional text warning) instead of the additional text warning (40%). Delegated/implementing power to adapt the additional health warnings.

• **Option 1: Mandatory enlarged picture warnings**
  Combined warnings (picture plus text) of 75% displayed on both sides of the packages of tobacco products, presented in rotation. TNCO levels on the packages are replaced with descriptive information on content, emissions and risks. Display of cessation information (e.g. quit-lines, websites) is added to the packages.

  Tobacco products other than FMC and RYO are exempted (current TPD rules apply). Delegated/implementing power to remove/extend the exemption for these products if there is a change of circumstances and to adapt the health warnings.

• **Option 2: Option 1 plus harmonise certain aspects of packets and prohibit promotional and misleading elements**
  Option 1 plus:
  1) The tobacco labelling and packaging and the tobacco product itself shall not include any promotional and misleading elements (e.g. misleading colours, symbols, slim FMC),
  2) setting certain requirements for packages (e.g. cuboid shape, minimum number of and FMC per package) as well as for the size of the warnings

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228 For STP, NCP and herbal products for smoking, see problem 1a, 1b, and 1c
229 Required increase of the size if more than one official language (32-35% and 45-50%)
230 Procedures foreseen in current TPD
231 The size of the health warning proposed is based on scientific evidence and international developments.
232 For definition, see glossary
Member States are allowed to regulate the area not regulated by the TPD or other Union legislation, including adopting provisions providing full standardisation of packaging of tobacco products (i.e. plain packaging) as far as these provisions are compatible with the Treaty. The Commission will report on experiences gained with respect to surfaces not governed by the TPD five years after the transposition of the TPD.

For Point 2) Tobacco products other than FMC and RYO are exempted (current TPD rules apply). Delegated/implementing power to remove the exemption for these products if there is a change in circumstances.

- **Option 3: Option 2 plus full plain packaging**
  Option 2 plus: standardised colour, font, size and position of brand name and brand variant on packages (plain packaging) and a readable health warning on each FMC stick.

Other tobacco products than FMC and RYO will be exempted from the requirements (current TPD rules apply). Delegated/implementing power to remove the exemption if there is a change in circumstances.

- **Discarded policy option**
  An additional option which was considered in an earlier stage of the impact assessment was to introduce health warnings on pipes and water-pipes as such. This option was not considered appropriate within this revision.

### 4.6. **Problem 3 – Ingredients**

The main problems are the lack of a harmonised reporting format resulting in additional burden for all stakeholders involved, the heterogeneous development in Member States in terms of ingredients regulation (toxicity, addictiveness, attractiveness) and the need to implement FCTC obligations and commitments. Also, the potential of certain tobacco products to mislead consumers and encourage people to take up or maintain smoking should be considered.

- **Option 0: No change**
  Status quo means: Current mandatory reporting without common format. No common ingredients regulation beyond toxicity and addictiveness reporting.

- **Option 1: Common reporting format on a voluntary basis. Prohibit toxic, addictive and attractive additives in tobacco products.**

  Member States are free to decide whether they oblige manufacturers to report additives in tobacco products. If Member States decide to make the reporting obligatory, the common reporting format must be used. Member States shall prohibit additives based on the general criteria toxicity, addictiveness and attractiveness.

- **Option 2: Mandatory reporting in harmonised format. Prohibit tobacco products with characterising flavours and products with increased toxicity or addictiveness.**

  Manufacturers are obliged to electronically report ingredients (glossary) of tobacco products in accordance with a common format and provide supporting data (e.g. marketing reports). Fees charged by Member States for handling the information submitted to them shall not exceed the cost attributable to those activities. Placing on the market of new or modified

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233 For STP, NCP and herbal products for smoking, see problem 1a, 1b and 1c.
tobacco products must not take place before the submission of ingredients data. Reported data, excluding confidential information, is published. Delegated/implementing power to specify the reporting format.

Tobacco products with characterising flavours (glossary) are prohibited (this is similar to the US model). Test panels assist in the decision making process. Additives associated with energy and vitality (e.g. caffeine and taurine) or creating the impression that products have health benefits (e.g. vitamins) are prohibited. No flavourings are allowed in filters, papers or packages. Tobacco products with increased toxicity or addictiveness shall not be placed on the market. Member States shall remove from the market tobacco products that include ingredients not complying with REACH. Delegated/implementing power to set limits for additives imparting a characterising flavour, toxicity and addictiveness.

Tobacco products other than FMC, RYO and STP (i.e. cigars, cigarillos and pipes) are exempted from the prohibition of products with characterising flavour and the prohibition of additives associated with energy and vitality or health benefits. Delegated/implementing powers to remove the exemption for these products are foreseen if there is a substantial change of circumstances.

This approach is similar to the US approach.

- **Option 3: Mandatory reporting in harmonised format. Prohibit all additives not essential for manufacturing.**

Ingredients reporting as in option 2. In terms of ingredients regulation, all additives in tobacco products, except those essential for manufacturing are prohibited. Maximum limits are set for sugar and sweeteners.

Other tobacco products than FMC, RYO and STP are exempted from the ingredients regulation. Delegated/implementing power to remove the exemption for these products if there is a change of circumstances.

This option is similar to legislations in Canada and Brazil.

- **Discarded options**

An alternative approach would have been to develop a common list of ingredients based on addictiveness and toxicity as foreseen in the current TPD (Articles 6 and 12). This would have responded to the concerns expressed by the European Parliament in its Resolution from October 2007. However, as explained in previous sections, at this stage scientific and international progress has primarily been achieved in terms of attractiveness rather than addictiveness and toxicity. Therefore, this approach was not considered, but should rather be seen as a measure to be taken at a later stage/over time.

### 4.7. PROBLEM 4 – CROSS-BORDER DISTANCE SALES OF TOBACCO PRODUCTS

Cross-border distance sales of tobacco products undermine the safeguards/full effect of the TPD provisions (labelling, ingredients, ban on oral tobacco) and encourages illicit cross-border trade (legal purchasing via the internet makes little sense). There is also a need to implement FCTC obligations and commitments. Finally, it facilitates access for young people.

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• **Option 0: No change**

Status quo means: Regulation is left to Member States. Council Recommendation 2003/54 applies, i.e. recommendation to restrict tobacco distance sales for general retail.

• **Option 1: Notification and age verification system**

Retailers of tobacco products intending to engage in cross-border distance sales shall notify their cross-border activities to the Member States where the company has its seat and where it intends to sell. Member States may require the retailer to appoint a natural person, who ensures compliance with the TPD of products delivered to customers in Member States concerned. Mandatory age verification mechanism is foreseen.

• **Option 2: Prohibit cross-border distance sales of tobacco products**

Cross-border distance sales of tobacco products are prohibited in the EU.

4.8. **Problem 5 – Traceability and security features**

The main problem in this area is that illicit products (estimated 8.25% of the market with increasing tendency\(^{236}\)) undermine the safeguards/full effect of the TPD provisions (labelling, ingredients). The current TPD provisions are incomplete, do not provide for a fully-fledged traceability system, and consumers have difficulties to verify the authenticity of tobacco products. There are also obligations as regards the fight against illicit products stemming from the FCTC (Article 15).

• **Option 0: No change**

The legally binding Agreements between the four biggest FMC manufacturers and the EU and Member States would continue to apply (until potential expiry as of 2016).\(^{237}\) Equivalent measures would not be in place for tobacco manufacturers without Agreement. The EU could adopt measures only on batch numbering (TPD Article 5(9)).

• **Option 1: EU tracking and tracing system**

An EU tracking and tracing system at packet level for tobacco products throughout the supply chain (excluding retail) is introduced. Tobacco manufacturers shall conclude contracts with independent third parties that provide data storage capacities for such system ensuring full transparency and accessibility by Member States at all times. Tobacco products other than FMC and RYO are granted a transitional period of five years.

Delegated/implementing power to adopt technical specification to ensure compatibility between the systems used.

• **Option 2: Tracking and tracing system, complemented by security features**

Option 1 plus: security features against counterfeiting and against illicit/cheap whites (glossary) on all tobacco products (e.g. holograms). Tobacco products other than FMC and RYO are granted a transitional period for five years. Delegated/implementing power to adopt technical specifications for the security features.

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\(^{236}\) Euromonitor as presented by Matrix 2012.

\(^{237}\) The European Union and all Member States have signed legally binding and enforceable agreements with PMI (2004) and JTI (2007) and the European Union and 26 Member States have signed agreements with BAT and ITL (2010)
5. ANALYSIS OF IMPACTS

In accordance with the Commission's impact assessment guidelines, this impact assessment analyses the likely impacts of identified policy options. As the overall objective of revising the TPD is to ensure a good functioning of the internal market while ensuring a high level of health protection, the **focus of the analysis is on economic-, social and health impacts**. Environmental impacts have been considered to be less significant and are therefore not specifically referred to in this section. The assessment criteria (listed in Annex 4) have been identified based on the specific objectives and health considerations set in chapter 3. Some additional comments are warranted upfront.

**Effectiveness, efficiency and coherence**

The comparison of policy options and the identification of preferred options are based on the general evaluation criteria **effectiveness** (the extent to which options achieve the objectives of the proposal), **efficiency** (the extent to which objectives can be achieved for a given level of resources/at least cost) and **coherence** (the extent to which options are coherent with the overarching objectives of EU policy).

**Data collection**

Collecting data and presenting it in a coherent form was a challenge when preparing this report. Data has been collected/received a.o. from NGOs, industry, external consultants, scientific studies, Member States, other Commission services and publicly accessible information. When, despite these efforts, information was considered inadequate, information from other comparable sectors were used. The analysis was complemented by qualitative assessments. Some stakeholders did not provide the information as requested or not in the format requested. Sometimes it was also difficult to reconcile publicly available data (e.g. Eurostat) with information received from the economic stakeholders. The data sets received from industry were also not always fully coherent (here an effort was made to reconcile the data to the extent possible). In order to ensure overall quality some of the key data was verified with the associated services (e.g. on tax revenues) and the industry (e.g. on turnover generated with tobacco products and the value chain by various stakeholders (farmers, upstream suppliers, tobacco industry, downstream distributors). Finally, information on the illicit part of the market is by definition difficult to establish in a robust form taking into account the nature of these activities.

**Direct impacts**

This impact assessment distinguish between direct impacts associated with the implantation of the options for the stakeholders' one-off and on-going costs and indirect impacts associated, in particular, with their effect on revenues. As for the **direct costs** for the economic stakeholders, five main cost categories have been identified and considered throughout the impact assessment: 1) familiarisation with the requirement of the new regulation, 2) redesign/reformulation of the package/product, 3) acquisition of new equipment (e.g. printing equipment), 4) costs for disposal of old stocks and equipment (e.g. printing equipment) and 5) variable/on-going costs (e.g. changes in types and quantities of material to be used). The estimation of the direct cost factor 1-4 has also been used as an indicator for the functioning

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239 For ex., see: Matrix 2012: "Industry has been able to provide limited data to demonstrate how particular regulatory change has impacted across the value chain in terms of costs or sales. The majority of evidence provided was indicative or too general."
of the internal market, i.e. the preferred option should normally lead to reduced compliance costs compared to the status quo, as economic stakeholders can adapt to EU provisions (as nationally implemented) in one go compared to consecutive changes at national level leading to diverging set of rules. For Governments, direct impacts are also estimated, e.g. in terms of administrative burden associated with the options.

**Indirect impacts**

The indirect impacts affect both the economic stakeholders (industry concerned, upstream/downstream actors, others) and Governments. Economic stakeholders are impacted in terms of revenue/profit and in terms of employment. In particular as regards employment, an input/output model was used which describes that money not spent on tobacco would be spent on other sectors. The expected main gain for Governments/society is the improvement of public health (gained life years). This impact has been monetised in line with the Commission’s impact assessment guidelines.\(^{240}\) In addition, the impacts on health care costs, reduced absenteeism and tax revenues are addressed. The calculations are based on an expected consumption drop of 2% within five years after the transposition.\(^{241}\) The indirect impacts are further described in section 5.7 and Annex 5.

**SMEs**

When analysing the impacts, specific considerations have been given to SMEs and micro-enterprises both in identifying the policy options and assessing their impacts. A summary of the impact on SMEs is set out in section 6.2.2.

**Stakeholders’** comments and concerns have also been carefully considered throughout the document, as illustrated by separate sections under each policy area describing the views expressed by key stakeholders, integration of key comments or essential points, and a detailed description in Annex 1.

The text is divided in five subsections following the five identified policy areas (STP and extension of the product scope, packaging & labelling, ingredients, cross-border distance sales and traceability & security features). Assessment of impacts, comparison of policy options and presentation of a preferred option will thereafter be undertaken separately for each of the policy areas. Scoring tables for the different policy areas can be found in Annex 4. A general conclusion presenting the overall combined preferred options can be found in section 6.1.

### 5.1. ASSESSMENT OF POLICY OPTION 0: NO CHANGE

#### 5.1.1. Economic impacts

**Internal market**

The increasing divergence between Member States' legislations under the baseline scenario leads to further fragmentation of the internal market. Given the significant amount of cross-border trade in tobacco and related products (section 2.1), this will negatively affect cross-border trade and undermine the level playing field for the economic actors involved.\(^{242}\)

**Direct impacts**

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\(^{240}\) For each life year, a value of 52,000 EUR is assumed.

\(^{241}\) For further information on how this drop was estimated, see Annex 5.

\(^{242}\) In certain areas Member States cannot adapt their legislation due to existing level of harmonisation. Also, there is a risk that the safeguards of the Directive are undermined by non-compliant products.
More fragmentation of the internal market leads to higher costs for economic stakeholders to comply with different national measures. In general, a qualitative description of the various cost categories suggests that the compliance costs under option 0 (consecutive national changes) are expected to be higher than if a harmonised approach across the EU was taken.

Following the cost categories identified above, the direct economic impact (compliance costs) can be described according to the table below. Categories 1-4 refer to one-off costs, while category 5 is a variable/on-going cost.

<table>
<thead>
<tr>
<th>Cost category (compliance cost)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Familiarisation with the requirements of the new regulation.</td>
<td>Economic stakeholders need to monitor 27 different legal systems and get familiar with all unilateral changes adopted by Member States. This can also include seeking legal advice.</td>
</tr>
<tr>
<td>2. Redesign/reformulation of the package/product</td>
<td>Economic stakeholders need to adapt to unilateral changes adopted by Member States. This also includes testing and legal/marketing/scientific advice. Stakeholders also have to adapt production lines to comply with different national legislations.</td>
</tr>
<tr>
<td>3. Acquisition of new equipment</td>
<td>Economic stakeholders may need to buy new equipment (e.g. printing equipment) to comply with national measures.</td>
</tr>
<tr>
<td>4. Costs for disposal of old stocks and equipment</td>
<td>Cost for the disposal of stocks can normally be disregarded provided sufficient transposition periods are granted.</td>
</tr>
<tr>
<td>5. Variable/on-going costs</td>
<td>Economic stakeholders might have to change the types and quantities of material to be used in the manufacturing process. This could be either a cost (e.g. more expensive ink) or a saving (e.g. less ingredients to be used) and is proportionate to the volume of sale.</td>
</tr>
</tbody>
</table>

The approach can be exemplified for the area of labelling, ingredients and traceability and security features. For estimations of costs, see impact assessment sections 5.2-5.6.

**Indirect impacts**

As far as indirect impacts (linked to consumption) are concerned, the impact on companies' revenues under the baseline scenario is expected to be neutral. This is based on the estimated stagnation of smoking consumption/prevalence (see above under baseline scenario, 2.3.3).

Also Governments' monetised value of public health, the tax revenues, health care costs and productivity losses are linked to the consumption/prevalence forecast. As mentioned in section 2.1.2, approximately 25 bEUR are currently spent every year to treat smoking related diseases and 8.3 bEUR are lost annually on productivity losses. From an economic point of view, the baseline scenario constitutes a missed opportunity for Governments to further reduce costs. From a macro-economic point of view, taxes are not a cost (taxes just reallocate revenues from economic stakeholders to the Government), but obviously reduced taxes have an impact on Government revenues.

**Consumers**

Under option 0, industry would have an incentive to present consumers with a larger choice of products. Consumers would also be subject to new innovative marketing, likely to target

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244 In the light of existing entry barriers it might not be easy, in particular for SMEs to enter the market in a sustainable manner with new brands.

243 It could be noted that a UK administrative burden exercise estimated the costs attributed to familiarisation and understanding of the food labelling regulation as 13% of all administrative costs across food legislation, see FSA (2006), "Food Standards Agency: Administrative Burdens Measurements Exercise: Final Report", June 2006
young consumers in particular and there would be a lack of protective measures.\textsuperscript{245} No particular impact is foreseen as far as the price is concerned. Consumers would also be exposed to serious health risks due to a lack of harmonised standards for some of the products, including STP, NCP and herbal products for smoking. The supply of illicit tobacco products would continue to be of (increasing) safety concern and to undermine tobacco control measures intended to protect consumers. In addition, as indicated in the problem identification, certain aspects of packaging & labelling, as well as certain ingredients may have the capacity to mislead consumers.

\textit{International}

In terms of \textit{international} impact, under option 0, the EU does not fully comply with its international obligations/non-binding commitments (FCTC)\textsuperscript{246} and lags behind other jurisdictions (including Canada, Australia and the US) in many of the areas covered by this impact assessment, including labelling and ingredients.

5.1.2. \textbf{Social impacts}

The specific impact on \textit{employment} is estimated under option 0 following the consumption forecast foreseen under the baseline scenario (section 2.3.3) (including redistribution effects). This is without prejudice to further consolidation and automatisation of production methods which can be expected and that would potentially entail employment cuts. In terms of \textit{equality}, option 0 has negative consequences on young people in particular. As illustrated in the problem definition, many of the products on the market currently target young people. This is true both for STP, NCP and herbal products for smoking, but also for FMC marketed in appealing packages or with flavours particularly attractive to young people. Due to the price structure RYO are also attractive for young people. The targeting of young people is expected to continue in coming years.

5.1.3. \textbf{Health impacts}

As outlined above, under the baseline scenario (section 2.3.2), Member States would be prevented from taking unilateral actions in \textit{areas already harmonised in the TPD}, notably as regards the size and placement of the health warnings, the removal of misleading TNCO levels and in the area of traceability. This is unsatisfactory from a public health point of view and is not in line with the obligation to ensure a high level of health protection in all policies and activities (TFEU Articles 168 and 114). The high level of premature mortality and morbidity would prevail (see section 2.1.2).

5.2. \textbf{SMOKELESS TOBACCO PRODUCTS AND EXTENSION OF THE PRODUCT SCOPE}

1.1.12. \textbf{Smokeless tobacco products (STP)}

5.2.1.1. \textit{PO1: Lift the ban on oral tobacco and subject all STP to stricter labelling and ingredients regulation}

a) The current ban on oral tobacco is lifted and these products can be placed on the market subject to stricter labelling and ingredients requirements (e.g. health warnings on both sides of


\textsuperscript{246} For ex. FCCT refers to effective legislation regulating the content of tobacco products, marking of all tobacco packages in order to determine the origin and health warnings of 50% or more (but no less than 30%).
the package and a ban of products with characterising flavour (glossary) and increased toxicity or addictiveness).

b) The placing on the market of chewing and nasal tobacco continues to be allowed, but subject to the same rules on labelling and ingredients as set out under a.

c) Placing on the market of novel tobacco products (glossary) continues to be allowed subject to the same rules on labelling and ingredients as for oral tobacco.

Delegated/implementing power to adapt health warnings, act on products with characterising flavours on products with increased toxicity or addictiveness and to regulate additives that cause a characterising flavour.

**Economic impacts**

Option 1 would allow the placing on the market of oral tobacco (snus) in the EU and remove the current differential treatment between different types of STP. However, when the ban on oral tobacco was introduced in 1992, three Member States had already adopted national bans. It is likely that, at the very least, these Member States will wish to maintain their approach, in particular when considering that most of the Member States responding to the public consultation explicitly supported keeping the ban on oral tobacco (Annex 1). This could negatively affect the functioning of the internal market if it reintroduces the fragmentation of the market that existed before 1992. Moreover, it is difficult to draw any firm conclusion on the effectiveness on STP in smoking cessation (see below under health impacts). In **regional** terms, option 1 would have a positive impact on the Finnish island Åland which claims being subject to unequal treatment under the current situation (section 2.2.1).

The most important impact for the **economic stakeholders** is that of increased sales of oral tobacco following the lifting of the ban, in particular as this product is new to the EU market (except for Sweden) and is potentially attractive to new customers, including young people. Smoke-free environments are a key driver for this expected growth. It is not possible to quantify the expected increase in sales as it is very difficult to foresee how new and unexplored markets would react to this product. The oral tobacco producer, Swedish Match, has indicated that the sale of oral tobacco in the EU could potentially generate gross profits to the retail sector in the amount of 3-9 bEUR. Chewing and nasal tobacco producers are likely to be confronted with decreased or stagnating sales in the light of increased competition from oral tobacco and a reduced product portfolio (no STP with characterising flavours, additional health warning). The same could be true for producers of NRT, FMC and RYO if oral tobacco is – as claimed by Swedish Match - used for smoking cessation purposes. However, the evidence to that end is not compelling (see also below on health impacts). Also, products claiming to assist smokers to quit smoking would require authorisation as medicinal products.

All STP producers would be faced with limited additional **compliance costs** in terms of the required additional health warnings and the restrictions in terms of ingredients (ban on STP

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247 This presupposes that potentially reduced sales in Sweden due to labelling and ingredients regulation are compensated through increased sales outside Sweden.

248 Swedish Match, power point from 15 December 2011. See also Swedish Retail Institute (HUI). Snus in the EU The potential economic impact of snus on retail taxes and employment in the EU. Stockholm: HUI; 2010. The lower estimate is based on the assumption that only former smokers would use snus while the higher estimate ("normal consumption scenario") is based on additional recruitment of new users. The calculation does not take into account a corresponding drop in FMC sale.

249 Currently there is a growth potential for STP in the light of smoke-free environments, see option 1.
with characterising flavours). On the other hand, an EU based approach is still cheaper than consecutive changes at national level. Additional compliance costs would consist of familiarisation and understanding of the new requirements, redesign of the packages and reformulation of the products. The industry has estimated the medium range cost for changing the labelling to 6,000 EUR per SKU, although this can vary from company to company. As regards reformulation, an important market development has taken place in STP (including oral tobacco) in recent years. Many products are presented with characterising flavours, including raspberry/pepper and rhubarb/ginger. These products would have to be reformulated or withdrawn from the market. While chewing and nasal tobacco producers are often SMEs, they typically also sell other tobacco products, making them less vulnerable.

Overall, it is expected that the costs linked to compliance with the suggested product standards would be significantly surpassed by the overall increased sales following the lifting of the ban on oral tobacco and thus result in positive economic impact for the oral tobacco industry. Significant product development in novel STP products claimed to imply a lower risk than FMC is also expected under option 1 (see 2.2.1).

As there are no harmonised excise duties on STP in the EU, the impacts on governments in terms of tax incomes would vary among Member States. Governments would be confronted with some additional health risks/costs (see below under health impacts). Overall a broader range of products would be available to EU consumers (including oral tobacco also outside Sweden). However, the option would impact negatively on consumer protection due to the health risks referred to below under health impacts. Prohibiting STP with characterising flavours would be in line with the FCTC guidelines on ingredients. Lifting the ban on oral tobacco would not be in line with the WHO recommendation from 1988 that "countries with no established smokeless tobacco habit should consider a ban on the manufacture, importation, sale and promotion of smokeless tobacco products before they are introduced to market or become established habit".

Social impacts

Option 1 would result in increased employment in STP overall, mainly benefitting the oral tobacco industry (Swedish Match as leading actor) which would have the possibility to explore new markets. It is also expected that the big FMC manufacturers will use this opportunity to enter the STP market on a large scale (further reinforcing this on-going trend). On the other hand, some negative impacts are expected as far as chewing and nasal tobacco employment is concerned following increased competition from oral tobacco. In terms of equality, the already on-going targeting of young people would continue under option 1, but 

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250 As previously explained a common EU approach is expected to be beneficial compared to consecutive amendments of national legislations.
251 Additional costs for changing in printing, packaging and production equipment as well as disposal of existing stocks could be removed if sufficient transposition periods are given.
252 European Smoking Tobacco Industry, facts and figures for DG SANCO, ESTA 2011-2012
253 Swedish Match magazine Inside #2 from May 2008 reports that Swedish Match portfolio has grown from 22 products in 2002 to 180 varieties in 2008 and that 14 new products are launched every year. New tastes flavours have been marketed in recent years, including raspberry/pepper (2011), mint/vanilla (2011), rhubarb/ginger (2010).
254 European Smoking Tobacco Industry: Facts & Figures for DG Sanco, ESTA 2011-2012
255 For example, the US market has seen a diversification of STP in recent years, including marketing of “dissolvable” tobacco products often sold in brightly coloured packaging an in an appearance similar to candy: Krisberg K. New types of smokeless tobacco present growing risks for youth: Survey: Products mistaken for candy. The Nation's Health 2010; 40:1-14. http://thenationshealth.aphapublications.org/content/40/6/1.2.full (accessed 28 Nov 2012).
the attractiveness of the products would be reduced under option 1 due to stricter measures on labelling and ingredients.

**Health impacts**

Studies show that STP are less hazardous to health than FMC and even option 1 would prevent STP with increased toxicity or addictiveness to enter the market.

Therefore, for individuals who replace FMC entirely with STP the overall benefits would outweigh the risks although full abstention from tobacco use would be the most beneficial. The overall health impacts on the population at EU level under option 1 would, however, largely depend on how the EU market reacts on the introduction of oral tobacco and in particular if this measure would contribute to reducing the number of smokers in the EU (substitution) or if it would rather increase the overall tobacco use contribute to initiation and undermine tobacco cessation efforts.

**Health effects of STP**

In terms of health, all STP tobacco products contain nicotine and are addictive. They also contain carcinogenic substances, including tobacco specific N-nitrosamines (TSNA) and polycyclic aromatic hydrocarbons (PAH). The Scientific Committee on Emerging and Newly Identified Health Risks (SCENIHR) concluded, in its opinion on 6 February 2008, that STP in all its forms can cause cancer (with the pancreas as a main target organ) and are addictive. The International Agency for Research on Cancer (IARC) has also classified smokeless tobacco as “carcinogenic to humans”. A study on smokeless tobacco and cancer from 2008 concludes that the cancer risk of smokeless tobacco users is probably lower than that of smokers but higher than that of non-tobacco users.

There are many forms of smokeless tobacco products, which differ considerably in their composition and toxic potential. Some chewing tobacco products, in particular some products used by the South Asian community in the UK, according to a recent study, contain a wide range of toxic substances, such as tobacco-specific nitrosamines (TNSA) chromium, nickel and lead. During the last two decades, the level of tobacco-specific nitrosamines (TNSA), the major group of carcinogens in smokeless tobacco, has been considerably lowered in some STP, including Swedish oral tobacco (snus). This means that the adverse health effects of snus might differ from other non-combustible tobacco products. However, it does not mean that snus or any other oral tobacco product is safe or harmless. Products with lower levels of carcinogenic tobacco-specific nitramines (TNSA) have also been on the market for too short time for any convincing support in favour of the presence or absence of a lower cancer risk.

The WHO Study Group on Tobacco Product Regulation concludes in its report from 2009


258 SCENIHR, 2008


that existing evidence has not established that lowering TSNA or PAH levels in smokeless tobacco will lower cancer risks.263

The link between STPs and pancreatic cancer has been discussed by the research community in recent years. Based on a number of case-control and cohort studies, the two authoritative international research groups SCENIHR and IARC have concluded that there is sufficient evidence that STPs cause pancreatic cancer in humans.264 A recent case-control study suggests, however, that there is no significant association between pancreatic cancer and smokeless tobacco.265 The discrepant results of this study with other case-control studies have been questioned by a number of researchers calling for a cautious interpretation in view of existing strong cohort data supporting an association between STP and risk of pancreatic cancer.266

Risk of oral cancer have been found for various smokeless tobacco products, including some of the chewing tobacco products (e.g. areca nut and betel quid) used by ethnic minorities in the UK.267 There are also suggestions that nasal tobacco increases the risk of certain cancers, e.g. oral cancers.268 The risk for oral cancer is less clear as regards Swedish oral tobacco (snus).269

SCENIHR concluded in 2008 that published studies support a causal role of STP in the etiology of oesophageal cancer.270 According to IARC, there is now sufficient evidence that there is a causal association between smokeless tobacco and oesophageal cancer.271

In addition, there is evidence for an increased risk of fatal myocardial infarction among STP users.272 Hansson concluded in a study from February 2012 that current snus users had a higher probability of dying from acute myocardial infarction (AMI) as compared to non-users,

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268 SCENIHR 2008
269 Idem
270 Idem
and that this increase may be explained in confounding factors, although a small increased risk of sudden death from AMI among snus users cannot be ruled out.\textsuperscript{273}

Some data also indicate that STP use is associated with several \textit{pregnancy complications}, including pre-term birth, intrauterine growth restriction, placenta abruption and still birth.\textsuperscript{274}

In conclusion, despite differences in composition and carcinogenic potential, there is scientific evidence that all STPs are addictive and harmful to health. As shown above, some of the epidemiological data are questioned by studies (partly sponsored by the industry) inconsistent but this does not put into question the overall conclusion. In any event it justifies the application of the precautionary principle, i.e. it justifies not allowing market entry of products, which are addictive and harmful.\textsuperscript{275}

\textbf{Cessation, initiation, dual-use}

\textbf{In terms of substitution}, some studies suggest that oral tobacco (snus) can play a role in smoking cessation\textsuperscript{276} or that oral tobacco users are more likely to quit smoking than users of medicinal smoking cessation products.\textsuperscript{277} Most of the studies are based on observational data, which makes it difficult to draw reliable conclusions as to the relative effectiveness of smokeless tobacco in smoking cessation.\textsuperscript{278} On the other hand, a randomised controlled trial showed that use of STP in cessation did not have any long-terms efficacy.\textsuperscript{279} Swedish Match recently sponsored two clinical trials comparing the effectiveness between oral tobacco (snus)


These findings are challenged by Barrett SP, Campbell ML, Temporale K, Good KBP. The acute effect of Swedish-style snus on cigarette craving and self-administration in male and female smokers. Human psychopharmacology 2011; 26(1):58-62. The study suggests that Swedish snus is effective in acutely suppressing craving and smoking in at least some smokers, but that its acceptability may be limited. Also, Kotlyar M, Hertsgaard LA, Lindgren BR, Jensen JA, Carmella SG, Stepnov I et al. Effect of oral snus and medicinal nicotine in smokers on toxicant exposure and withdrawal symptoms: a feasibility study. Cancer Epidemiol Biomarkers Prev 2011; 20(1):91-100 found that Camel Snus and Taboka use was not superior to medicinal nicotine in reducing withdrawal symptoms.

\textsuperscript{278} SCENIHR 2008, p 110

and placebo products in smoking cessation in Serbia and the US. The studies suggest that smokers using Swedish snus were 2-3 times more likely to quit smoking than those using placebo-products. However, the studies took place over a relatively short time (24 and 28 weeks) and it is impossible to say whether the relatively few people quitting smoking in these studies would have done so also without oral tobacco or also, or even more, with the assistance of NRT. In this context it should also be considered that 2/3-3/4 of smokers quit un-aided.

Sweden's low smoking prevalence in combination with the availability of oral tobacco (snus) is sometimes referred to as an indication of snus as an effective cessation method and there are some data indicating that snus has been used by Swedish smokers as an alternative to smoking. On the other hand, SCENIHR concluded in 2008 that the overall smoking prevalence in Norway, as well as in young Norwegians, had decreased at the same rates in men and women during the last decade, whereas a marked increase in oral tobacco (snus) use during this time period has only occurred in young men. In California, both the prevalence of smoking and smokeless tobacco use have decreased concurrently. Some countries which have invested heavily in preventive measures have also managed to reduce smoking rates without the availability of STP. These data imply that the association between patterns of STP tobacco use and smoking cessation differs between populations and is likely to be affected by cultural, societal and other factors. In this context, SCENIHR has concluded that it is not possible to extrapolate the trends in prevalence of smoking and use of oral tobacco from countries where oral tobacco is available to EU-countries where oral tobacco is not currently available. This is highly relevant, as the sale of the product cannot be limited to people who wish to stop smoking, unless the product is a medicinal product available only on prescription.

Option 1 is also expected to result in uptake of STP use among individuals (including among young people) who would otherwise not have used tobacco. A survey undertaken by the Swedish National Institute of Public Health reveals that four out of ten oral tobacco (snus) users started using tobacco with oral tobacco. In Norway, recruitment of oral tobacco (snus) users among young people, including recruitment of those with no previous experience

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281 The Fagerstrom study (2011) reported that only 4% (5 persons) abstained from week 6-28.


283 Stenbeck M, Hagquist C, Rosén M. The association of snus and smoking behaviour: a cohort analysis of Swedish males in the 1990s. Addiction 2009;104(9):1579-85. See also SCENIHR 2008 although SCENIHR also concludes that it is not clear whether or by how much the availability of snus has influenced smoking prevalence. This conclusion by SCENIHR is based on Norwegian data up until 2006. Data after 2006, however, indicates a significant increase of oral tobacco (snus) uptake also among women: http://www.helsedirektoratet.no/folkehelse/tobakk/tall-ogundersokelser/sinus/Sider/unge-snuser.aspx (accessed 28 Nov 2012). Scheffels (2012) also concludes that Norwegian men frequently use snus as a method for quitting smoking whereas women are more likely to use NRT.

284 SCENIHR 2008


286 Holm LE, Fisker J, Larsen BI, Puska P, Halldórrsson M. Snus does not save lives: quitting smoking does! Tob Control 2009;18(4):250-1. See also Swedish Retail Institute (HUI). Potential savings on social costs by the use of Swedish snus as a harm reduction device in the EU: HUI; 2010. In this study, 5 million new snus users are estimated if snus was allowed in the EU.
of smoking, is increasing.⁴⁸⁹ Results from cross-sectional studies from Norway show that over 40% of young people (16-20 years old) of daily snus users had no previous smoking experience.⁴⁹⁰ Considering the current marketing strategies described under the problem identification (e.g. STP with distinctive tastes) and the obvious interest of the industry to recruit new users, a non-negligible uptake rate is expected under option 1. Smoke-free environment also play an important role in this respect.

Although there is currently limited evidence regarding novel STP, which are yet to be marketed to consumers, there may also be a risk of uptake of these products among new users and smokers who would otherwise have quit smoking altogether. Despite the claim that these products are reduced risk products, they are addictive and harmful to health. As BAT states on their web site: "Cigarette smoking is a cause of serious and fatal diseases and the only way to avoid the health risks associated with tobacco products is to not use them."⁴⁹¹ There is also uncertainty as regards STPs' potential as a "gateway" to future smoking. Evidence from the US indicates that oral tobacco use may lead to subsequent FMC smoking, while some Swedish data do not support this hypothesis.⁴⁹² The SCENIHR opinion of February 2008 suggests caution in translating these data.⁴⁹³ Moreover, there is a risk of "dual use". One study of snus as a cessation method found that 20% of unsuccessful quitters continued to use snus on a daily basis (dual use).⁴⁹⁴ A recent Norwegian study has also found that around 30% of daily snus users were smoking at least occasionally.⁴⁹⁵ Oral tobacco (snus) use in early adolescence has also been associated with increased risk of taking up occasional smoking in addition to snus in late adolescence.⁴⁹⁶ There is also a risk that consumers taking up STP will become chronic users.⁴⁹⁷ Finally, there is a risk under option 1 that lifting the ban on oral tobacco could have a negative impact on overall tobacco control policies. Norway has in its response to the public consultation on the TPD pointed to difficulties from a communication point of view of advocating non-use of oral tobacco (snus) among young people and at the same time

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²⁹¹ http://www.bat.com/group/sites/uk_3mnfen.nsf/vwPagesWebLive/DO52AMGZ?opendocument&SKN=1


²⁹³ SCENIHR 2008


²⁹⁵ Lund KE, McNeill A. Patterns of Dual Use of Snus and Cigarettes in a Mature Snus Market.


advocating the use of the same product as a smoking cessation tool for another group. The same is true for other types of STP, including novel non-combustible products. In addition, the introduction of oral tobacco could potentially weaken cessation policies, in particular as it would allow people to keep up their nicotine addiction in situations where smoking is not allowed (e.g. smoke-free environments) and subsequently resume smoking.

Summarising the findings on oral tobacco, it is not possible at this stage to draw the conclusions oral tobacco is an effective smoking cessation aid in the long term. Any impacts therefore on smoking-related diseases remain uncertain under option 1. On the other hand, it is likely that new oral tobacco users would be recruited under option 1 who would otherwise not have used tobacco (entry gate) and current smokers who would otherwise have quit using tobacco altogether might switch to oral tobacco or use both products (dual use). This would lead to increased adverse health effects (see section 2.2.1). In this light, it appears difficult to reconcile lifting the ban with the precautionary principle.

5.2.1.2. **PO2: Maintain the ban on oral tobacco, subject all novel tobacco products to a notification obligation and all STP to stricter labelling and ingredients regulation**

- a) The current ban on oral tobacco is maintained (except for Sweden which has an exemption), but oral tobacco needs to comply with stricter labelling and ingredients regulation (e.g. health warnings on both sides of the package and a ban of products with characterising flavours and increased toxicity or addictiveness).
- b) The placing on the market of chewing and nasal tobacco continues to be allowed, subject to the same rules as set out under a. A clearer definition of chewing tobacco is inserted.
- c) A notification obligation is introduced for novel tobacco products (glossary) and a report on the market development in these products will be issued by the Commission five years after the transposition of the TPD. Novel tobacco products placed on the market must respect the rules on labelling (health warnings on both sides) and ingredients regulation (ban on products with characterising flavours).

Delegated/implementing power as in option 1.

**Economic impacts**

This option would contribute to a more homogenous development in terms of labelling and ingredients regulation of all STP. The insertion of a clearer definition of chewing tobacco (compared to tobacco products for oral used) would increase legal certainty and prevent circumvention of the TPD provisions. Compared to the baseline scenario, option 2 is considered to have a positive impact on the internal market. The differential treatment in terms of regional impact on Åland would remain neutral under option 2, i.e. ships under Swedish flag would be allowed to sell tobacco whilst vessels under the flags of Finland and Åland serving the same routes would not be allowed to do so. However, this differential treatment is an effect of the Swedish exemption on oral tobacco rather than the ban itself. The differential treatment of oral tobacco and other forms of STP would also persist, as oral tobacco producers would not have the option to expand sales outside Sweden, but as the Court

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299 This is also in line with the Courts reasoning in Case C-434/02 Arnold André GmbH & Co. KG v Landrat des Kreises Herford [2004] ECR I-11825, p 41
ruled, the differential treatment was justified in 2001. The underlying analysis has not changed.

The impact in terms of compliance costs for the economic stakeholders involved in STP manufacturing would be similar to the once described under option 1, i.e. costs linked to the change of labelling and possible reformulation of additives. Option 2 also implies an obligation for manufacturers to report novel tobacco products prior to their placing on the market. The notification would also be accompanied by available scientific and market studies as well as other available and relevant information, including risk/benefit analyses of the products. This obligation would imply an additional burden for manufacturers intending to place novel tobacco products on the market. However, the burden is considered limited as it focus on already available information and information already required under the ingredients reporting requirements (policy area 3). In terms of indirect costs linked to consumption, the sale of oral tobacco would remain limited to Sweden and the sales potential to other Member States allowed for under option 1 cannot be explored under this option. The prohibition of STP with characterising flavours would also affect negatively sales of STP as well as the increased labelling requirements providing consumers with more visible information about the health risks associated with the products.

The administrative burden for Member States of introducing a notification obligation for novel tobacco products under option 2 would be limited, if any. Option 2 would also result in reduction of the tax income for Governments following the drop in STP consumption, but they are expected to benefit from better health outcomes. EU consumers would have a more limited choice of STP than today due to the ban on STP with characterising flavours, but in particular young potential consumers could be better protected against initiation and better informed. The ban on STP with characterising flavours would be in line with FCTC guidelines on ingredients (also covering STP). Overall the option would lead to a consistent approach for all tobacco products that are likely to be used by young people (tobacco consumption initiation).

Social impacts

Option 2 would have some limited negative impact on employment due to the forecasted reduction in use and also because no new STP with characterising flavours attractive to young people could be developed. For the chewing and nasal tobacco industry, this would particularly affect certain regions where chewing and nasal tobacco products currently are produced (mainly in Germany, Denmark, Belgium and Poland). This negative impact, however, is more limited than the impact under option 4 as the products could still remain on the market. Overall, it is expected that some redistribution will take place as money not spent on STP is spent on other economic activities (see reasoning in Annex 5).

Health impacts

Under option 2, chewing and nasal tobacco would lose part of their current appeal and the future development of STP with distinctive tastes or aroma would be prohibited. Option 2 would result in decreased use of STP. Also the additional health warning would reduce the appeal of these products and better inform consumers. As in option 1, option 2 would also

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300 Member States would also have to ensure that sales are limited to territory, i.e. they would have to prohibit cross-border distance sales. Taking into account that the number of traditional STP is limited, the administrative burden for enforcement is limited.
prevent STP with increased toxicity or addictiveness to be placed on the market. Under option 2, Member States would be placed in a better position to monitor market developments with regard to novel STP.

5.2.1.3. **PO3: Maintain the ban on oral tobacco, restrict the sale of other STP to areas of traditional use only and subject all STP to stricter labelling and ingredients rules**

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<td>a)</td>
<td>The current ban on oral tobacco is maintained (except for Sweden which has an exemption), but oral tobacco needs to comply with stricter labelling and ingredients regulation (e.g. health warnings on both sides of the package and a ban on products with characterising flavours and increased toxicity or addictiveness).</td>
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<tr>
<td>b)</td>
<td>The placing on the market of chewing and nasal tobacco is banned unless traditionally used (glossary) in Member States. The placing on the market of traditionally used STP is limited to the relevant territory/Member State and must comply with the same labelling and ingredients rules as set out under a. Member States will have to prove traditional use and notify to the Commission.</td>
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<tr>
<td>c)</td>
<td>The placing on the market of novel tobacco products is banned. Delegated/implementing power as in option 1.</td>
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**Economic impacts**

Option 3 would impact positively on the internal market by removing current regulatory disparities for chewing & nasal tobacco and by ensuring equal treatment between different STP. It would reduce the growth potential (e.g. smoke-free environments) of STP with adverse health effects and prevent the launch of novel tobacco products with adverse health effects.

The impact for the economic stakeholders would be that the sale of oral tobacco would remain limited to Sweden (having an exemption from the ban) and the sale of other STP would be limited to Member States with traditional use. All STP industries would have to comply with stricter rules on labelling (an additional health warning to be added to the packages) and on ingredients (prohibit STP with characterising flavours). The compliance costs would be linked to the change of labelling and possible reformulation of additives. Again, an EU approach is more cost effective than consecutive changes at national level. The companies affected are mainly SMEs although they are also involved in other tobacco products, which means that most of them would not have to exit the market altogether. The negative impact would be further reduced by allowing Member States exemptions for traditionally used STP.

Option 3 would result in reduction of the tax income for Governments following the drop in STP consumption, but they would benefit from better health outcomes. EU consumers would have a more limited choice of STP than today due to the ban on STP with characterising flavours but in particular young (potential) consumers would be protected and/or at least better informed. In terms of international impacts, option 3 could have some negative impact on international trade due to the current import of certain STP from third countries (e.g. import for ethnic minorities). However, considering that chewing and nasal

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301 European Smoking Tobacco Industry, facts and figures for DG SANCO, ESTA 2011-2012
302 Member States would also have to ensure that sales are limited to territory, i.e. they would have to prohibit cross-border distance sales. Taking into account that the number of traditional STP is limited, the administrative burden for enforcement is limited.
tobacco products count for less than 0.1% of the total tobacco market, this impact is considered limited. Also, the exemption of traditional use could apply. The ban on STP with characterising flavours would be in line with FCTC guidelines on ingredients (also covering STP). Overall the option would lead to a consistent approach for all products that are likely to be used by young people (tobacco consumption initiation).

**Social impacts**

Option 3 would have some negative impacts on employment due to the forecasted reduction in use and also because no new STP with characterising flavours attractive to young people could be developed. For the chewing and nasal tobacco industry, this would particularly affect certain regions where chewing and nasal tobacco products currently are produced (mainly in Germany, Denmark, Belgium and Poland). This negative impact is mitigated by the fact that most of these manufacturers are also involved in the production of other tobacco products as well and that Member States can ask for derogations for traditionally used STP for their own territory. In macroeconomic terms the potentially negative impact will be linked to the size of the operators concerned and the expected derogations. Overall, it is expected that some redistribution will take place as money not spent on STP is spent on other economic activities (see reasoning in Annex 5).

Option 3 would remove the current targeting of young people and thus contribute to increased equality. The option could impact on certain ethnic minority groups using specific chewing tobacco products (gutkha, zarda, khaini) if Member States do not ask for an exemption of these products.

**Health impacts**

Under option 3, chewing and nasal tobacco would lose part of their current appeal and the significant market potential of novel STP would be suppressed. Option 3 would result in decreased use of STP and remove the circumvention potential that certain STP have following the ban of oral tobacco. The option also has the potential of impacting positively on smoking cessation by motivating current smokers to free themselves from nicotine addiction altogether rather than using STP in environments where smoking is not allowed. Also Member States exempting STP for traditional use, the ban on products with characterising flavour and the additional health warning would reduce the appeal of these products and better inform consumers. In terms of toxicity and addictiveness, reference is made to options 1 and 2.

5.2.1.4. **PO4: Ban all STP with the exception of oral tobacco in Sweden which would be subject to stricter labelling and ingredients rules**

| a) The current ban on oral tobacco is maintained (except for Sweden which has an exemption) |
| b) The placing on the market of chewing and nasal tobacco is banned given their circumvention potential, in particular as regards chewing tobacco. |
| c) The placing on the market of novel tobacco products is banned. |

**Economic impacts**

Similarly to option 2, option 4 would remove the legal uncertainty as to the classification of STP, which is currently based on the mode of use (“sucking” vs “chewing”). Option 4 would thus prevent circumvention of the ban by products which are comparable (see section 2.2.1). This would improve the effectiveness of the TPD and thus impact positively on the **internal market**. It would also remove the growth potential (e.g. smoke-free environments) of products with adverse health effects and prevent innovation and launch of novel STP,
including non-combustible tobacco products claimed to be less harmful than traditional FMC. The impact on Åland would be similar to options 2 and 3.

The compliance costs for economic stakeholders involved in oral tobacco would mean some costs for complying with labelling and ingredients regulation while the actual sales of oral tobacco in Sweden could continue. For chewing and nasal tobacco, it is expected to result in negative impact as all STP are prohibited under this option. This negative impact would exceed the impact outlined under options 2 and 3. For novel tobacco products, the effects of options 3 and 4 are identical.

Option 4 would result in reduction of the tax income for Governments following the estimated drop of chewing and nasal tobacco use. The administrative burden would be limited. EU consumers outside Sweden would have no access to STP (except for private import, but excluding purchases via internet). In terms of international impacts, option 4 could have some negative impact on international trade due to the current import of certain STP from third countries. However, considering that chewing and nasal tobacco products count for less than 0.1% of the total tobacco market, this impact would be limited.

Social impacts

Employment in oral tobacco would somewhat decrease compared to the baseline scenario, while a negative impact is expected for the chewing and nasal tobacco following the ban of these products. This would particularly affect certain regions where chewing and nasal tobacco products currently are produced (mainly in Germany, Denmark, Belgium and Poland). This negative impact is mitigated by the fact that most of these manufacturers are involved in the production of other tobacco products as well. In macroeconomic terms the potentially negative impact will be linked to the size of the operators concerned. As option 4 (opposite to option 3) does not allow derogations for traditionally used STP, the impact on economic stakeholders would be more important under option 3. The option would also have a negative impact in terms of innovation of novel STP products. Overall, it is expected that some redistribution will take place as money not spent on STP is spent on other economic activities (see reasoning in Annex 5).

In terms of equality, option 4 would remove the current targeting of young people. It would also impact negatively on certain ethnic minority groups using specific chewing tobacco products (gutkha, zarda, khaini).

Health impacts

A ban of all STP would remove access to STP (except for oral tobacco in Sweden) and thus result in less adverse health effect linked to STP use.

5.2.1.5. The views of stakeholders

The tobacco industry (including the STP industry) has expressed support for lifting the current ban on oral tobacco and subject all STP to product standards which already apply to oral tobacco (snus). In this context it has been argued that the current ban on oral tobacco is discriminatory, that oral tobacco is less harmful than FMC and other STP and could help current smokers quitting. Sweden (and the Finnish island Åland) is also in favour of lifting the current ban on oral tobacco (snus), while most of the other Member States have expressed

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303 European Smoking Tobacco Industry, facts and figures for DG SANCO, ESTA 2011-2012
support for keeping the current ban or extend it to all STP (see Annex 1).

On the other hand, the Swedish National Institute of Public Health supports keeping the ban on oral tobacco with the argument that the introduction of every new tobacco product on the EU market increases the dependency and all forms of tobacco is damaging to health. Also health NGOs have argued in favour of keeping/extend the current ban on oral tobacco. One FMC manufacturer has also suggested a new legal framework for regulating potentially reduced harm products.

5.2.1.6. Comparing the options and preferred option

In terms of effectiveness, options 1, 3, and 4 would formally ensure equal treatment of all STP categories (chewing, nasal, oral) in terms of placing the products on the market, but these options do not take into account the significant differences between the products, e.g. in terms of consumption patterns and growth potential. Accordingly, options 1, 3 and 4 do result in the achievement of policy objectives A1 and A2. The ban on STP with characterising flavours under option 2 would contribute to a consistent approach for all tobacco products capable of encouraging young people to take up smoking because of their distinct taste or flavour. This would partly contribute to objectives A1 and A2 (equal treatment and level playing field). Options 1-3 would all contribute to the achievement of objective B1 as they would set common requirements for labelling and ingredients. This objective is less relevant for option 4, implying a ban of STP. Option 1 would facilitate market surveillance (objective B2) as it implies a lifting of the ban on oral tobacco and thus removes the circumvention potential for such products. Also options 3 and 4 would contribute to objective B2 as they would result in a ban of all STP (except in Member States of traditional use in option 3). Also option 2 would contribute to this objective as it would include a clearer definition of chewing tobacco and thus prevent circumvention and facilitate surveillance. Policy consideration C1, regulating hazardous products, would be fully fulfilled under policy options 3 and 4 as these options allow no new STP on the market. This policy consideration (C1) is partly fulfilled by option 2 as this option prevents oral tobacco to be placed on new markets and limits the development of novel STP in the sense that it prevents products with characterising flavours to be developed. Options 1, 2 and 3 are well in line with policy considerations C2 and C4 as they prohibit STP with characterising flavours and strengthen the labelling provisions, i.e. increase the visibility of the information to consumers. Option 3 and in particular 4 would also contribute to policy consideration C5 as they would reduce/remove access to STP. Option 2 would partly fulfil this consideration, but only as regards oral tobacco and STP with characterising flavours.

Lifting the ban on oral tobacco in option 1 provides market opportunities and thereby advantages for certain economic operators (others are likely to suffer). Options 2, 3 and 4 would mean that STP producers have to comply with the requirements in terms of labelling.

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304 Member States (including Sweden's) positions can be found on the public consultation web site: [http://ec.europa.eu/health/tobacco/consultations/tobacco_cons_01_en.htm](http://ec.europa.eu/health/tobacco/consultations/tobacco_cons_01_en.htm) (accessed 28 Nov 2012). The Swedish position has also been reiterated in a letter from the Minister of Trade on 14 December 2010. The Swedish Ministers for Children and the Elderly and for Trade also asks for a uniform and responsible regulation covering all tobacco products in a letter of 24 October 2012.


307 Only the objectives/considerations relevant for this policy area are discussed in this section.

308 See the Courts reasoning in Case C--434/02 Arnold André GmbH & Co. KG v Landrat des Kreises Herford [2004] ECR I-11825, p 41.
and ingredients and the growth potential for oral tobacco would remain unexplored but an EU solution is preferable to consecutive changes at national level. Option 3 and 4 would also limit novel tobacco products from being placed on the market and require some chewing and nasal tobacco manufacturers to exit the markets, but the effect on existing chewing and nasal tobacco producers would be mitigated under option 3 by exemptions for traditionally used STP.

Lifting the ban on oral tobacco (option 1) would have adverse health effects. It could also attract new tobacco users who would otherwise not have taken up tobacco consumption. Moreover, it would have a negative health impact on smokers who would otherwise have quit smoking, but who continue to use both FMC/RYO and STP (dual use) and for smokers taking up STP use who would otherwise have quit using tobacco altogether. For individuals replacing FMC/RYO with STP completely, the health effects would be positive. Considering the uncertainty in relation to substitution, an overall negative outcome under option 1 cannot be excluded. Therefore, this option raises doubts in terms of coherence with the precautionary principle.

Preferred policy option 2: Maintain the ban on oral tobacco, subject all novel tobacco products to a notification obligation and all STP placed on the market to stricter labelling and ingredients regulation

The preferred option would provide a common framework for STP in terms of ingredients and labelling while keeping the current ban on placing of the market of oral tobacco untouched. The proposed ban of placing on the market of STP with characterising flavour would address recent market development and discourage young people from taking up STP use. Putting health warnings on both sides of the package will increase the visibility and thus better inform consumers about the health risks associated with STP use. The preferred option is well in line with FCTC guidelines on ingredients (ban on STP with characterising flavours) and labelling (warnings on both sides). The preferred option would also contribute to increasing the knowledge base as regards novel tobacco products following Member States’ notifications of these products.

It is not considered justified to lift the current ban of placing on the market of oral tobacco which was introduced already in 1992 and which was justified from an internal market point of view since three Member States had already banned or announced bans of the product due to its harmful and addictive effects. At that point in time oral tobacco had also started to be distributed on the market of certain Member States in such a way as to attract young people. Given the introduction of smoke-free environments and the continuous development of oral tobacco, although limited to the Swedish market, the risk of uptake in new population groups (including young people) remains. The industry has also confirmed that oral tobacco has huge market potential, referring to gross revenues to the retail sector in the amount of 3-9 bEUR per year, if the ban was lifted (see section 2.2.1). Although industry emphasises that oral tobacco is less harmful than FMC, they do not claim that it is harmless.309 Adverse health effects were indeed confirmed by several risk assessment bodies, including one of the Commission's advisory Committees (SCENIHR).310

310 SCENIHR 2008
Maintaining the ban of placing on the market of oral tobacco is considered to be the only effective measure to contain the use of this product and avoid uptake in new population groups, among non-smokers and by young people. Reinforcing the health warnings and ban oral tobacco with characterising flavours would not be a sufficient measure in this regard. Although it could have some effect on discouraging young people from taking up use of oral tobacco, the product would still be promoted for use in smoke-free environments and as described under option 1, in particular given the significant growth potential referred to by manufacturers. The establishment of general product regulation (option 1) could reduce, to a certain extent, the hazardous effect of STP, but as illustrated in the assessment section (option 1), the product remains addictive and has adverse health effects. The current ban was also seen as proportionate by the Court in 2003 due to the harmful effects of oral tobacco, the uncertainty of oral tobacco as a substitute for FMC, the addictive and toxic properties of nicotine, oral tobacco's risk potential for young people and the novelty of the product. This reasoning is still valid today. The current ban was moreover deemed non-discriminatory by the Court in view of the fact that oral tobacco was new to the markets of the Member States, authorising a difference in treatment with regard to other STP. This reasoning is also still valid today.

The addictive properties and the adverse health effects are also an issue as far as other STP are concerned (see option 1), but despite some market increase (see section 2.3.1) these product have still very small markets in the EU although a certain diversification in terms of flavours used and promotion of these products could be observed in recent years (see section 2.3.1). Moreover, the cost-intensive production method (partly done by hand) and the very limited number of SMEs who do not have the financial strength to enter new markets where STP has no traditional use reduce the export potential of other STP (e.g. chewing and nasal tobacco) and preserves a small scale manufacturing.

For these products, it is therefore considered that further development and recruitment of new users can be contained without a general EU ban of placing these STP on the market. However, the preferred option would also ban STP other than oral tobacco with characterising flavour and make the health warnings more visible. These considerations provide another reason why maintaining the ban of oral tobacco with its significantly greater market potential (see above in this section) is considered compatible with the principle of proportionality.

There are no less stringent alternative measures that would guarantee the same level of health protection as those suggested under option 2. Subjecting STP to general product standards would not discourage young people from using these products although it could somehow reduce the hazardous effects of STP. The option is also a proportionate limitation of STP companies' freedom to conduct business (Article 16 of the Charter of Fundamental Rights of the European Union) justified in order to protecting public health.

The preferred option would provide an EU added value as it would ensure a harmonised approach throughout the EU and create a level playing field for operators involved in STP while ensuring a high level of health protection. By definition, only an initiative at EU level is capable of ensuring a harmonised approach. Unilateral actions by Member States can contribute to a certain extent to the protection of health, but would certainly result in divergences and unequal health protection.

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312 Idem, paras 68-70.
313 SANCO discussions with stakeholders involved in STP.
1.1.13. **Nicotine containing products (NCP)**

5.2.1.7. **PO1: Subject NCP to labelling and ingredients requirement under TPD**

| NCP placed on the market are subject to adapted health warnings, ingredients reporting and a prohibition to place on the market NCP with characterising flavours (glossary). Delegated/implementing power to adapt health warnings and act on products with characterising flavours, on products with increased toxicity or addictiveness and to regulate additives that cause characterising flavours. |

**Economic impacts**

Option 1 would contribute to a more homogenous development, but some Member States are expected to continue to consider NCP as medicinal products by function, which would maintain legal uncertainty and two parallel legal systems.\(^{314}\) Overall, it is therefore expected that the functioning of the *internal market* is not improved in a satisfactory manner. The use of NCP for smoking cessation purposes (see section 2.1.3) can be expected to continue to a certain extent under option 1. This would result in maintained negative impact on the internal market of smoking cessation products authorised under the medicinal products' legislation.

In terms of compliance costs for the *NCP industry*, option 1 implies a saving compared to the baseline scenario. This is mainly due to the reduced one-off costs related to familiarisation, redesign of the package/labelling and possible reformulation of the products following from an EU-wide measure rather than several consecutive measures at national levels. The costs for adding a textual health warning is estimated to around 7,000-9,000 EUR per.\(^{315}\) Additional costs for new equipment and disposal of stocks costs are disregarded as these costs could be addressed by granting sufficiently transitional periods. Variable costs for changes in printing, packaging and production are estimated to be marginal and in any case passed on to the subsequent levels of trade. The same reasoning is applicable as far as ingredients regulations (ban on NCP with characterising flavours) are concerned. Continuous increase in the sale of NCP is expected under this option although at a lower pace due to increased information and stricter ingredients regulation.

Option 1 could influence some *Governments* to adapt taxes on NCP similar to tobacco products and thus contribute to increased tax income. Current increase in health care risks/costs associated with NCP is expected to slow down due to stricter ingredients regulation and increased awareness. Enforcement would remain an issue in light of many small companies active in the business and internet sales. Option 1 would allow EU *consumers* to maintain a wide choice, with some limitations due to the ban on products with characterising flavours.

**Social impact**

\(^{314}\) Following the EU case-law, for the purposes of determining whether a product falls within the definition of a medicinal product by function, the national authorities, acting under the supervision of the courts, must decide on a case-by-case basis, taking account of all the characteristics of the product, in particular its composition, its pharmacological properties to the extent to which they can be established in the present state of scientific knowledge, the manner in which it is used, the extent of its distribution, its familiarity to consumers and the risks which its use may entail (Case C-319/05, Commission of the European Communities v Federal Republic of Germany, ECR [2007] p I-09811, para 55)

\(^{315}\) Impact Assessment Report from the European Commission on General Food Labelling Issues, 30 January 2008. As for tobacco products, RAND Europe has estimated an annual cost of 2000-9720 EUR per SKU related to administrative burden of labelling (i.e. not only costs related to redesign). As these products are currently not subject to labelling requirements it has been estimated more appropriate to apply these food related figures than figures available for tobacco products.
The current increase in employment opportunities in NCP is likely to continue at a pace in line with the expected consumption. Option 1 would contribute to more equal protection, but targeting of young people cannot be excluded, although the marketing of products with characterising flavours would be removed and health warnings would be added. The prohibition of advertising for tobacco products pursuant to Directive 2003/33/EC does not apply to these products.

Health impacts

A health warning would provide more accurate information and increase awareness. The current increase in the use of NCP is expected to continue, but at a slower pace due to improved information and less attractive products.

The uncertainty in terms of adverse health effects associated with NCP use outlined in the problem identification (section 2.2.1) would persist under option 1. In terms of substitution, there is no conclusive evidence at this stage that NCP can be effectively used in smoking cessation (see section 2.2.1) and, as described in section 2.1.3, manufacturers and distributors of these products normally make no such claims because they know it would oblige them to ask for authorisation as medicinal products. On the other hand, option 1 would allow people to keep up their nicotine dependence also in situations where they cannot smoke and then resume smoking when this is allowed.

If, on the basis of the reporting obligation, data is collected on the content of NCP, this option would allow for further content regulation over time based on toxicity and addictiveness consideration through delegated/implementing act. Such a measure could remove some of the health hazards associated with NCP.

5.2.1.8. PO2: Establish a new authorisation scheme for NCP

| Only NCP that have been authorised under a new authorisation procedure (risk/benefit analysis) set up under TPD are allowed to be placed on the market. Otherwise, placing on the market of NCP is prohibited. The authorisation procedure would also cover labelling and additives control. |

Economic impacts

Option 2 would result in two parallel authorisation schemes for NCP: one scheme which would apply if the product falls under the medicinal products’ legislation by presentation or by function and another one for consumer products. Competition between these two categories cannot be excluded. Some of the current legal uncertainty would persist under this policy option due to this dual approach which means that similar (or even identical) products could be subject to different schemes. Such uncertainty does not favour the functioning of the internal market. It raises also the question of equal treatment with existing nicotine replacement therapies (NRT), which are subject to medicinal products' authorisations.

In terms of impact on economic stakeholders, option 2 would imply fees linked to the authorisation scheme established under this option. The level of the fees would depend on the nature of the scheme and the authorisation procedure foreseen, including the criteria used in the assessment. The estimations used under the medicinal products’ framework can give an indication, but it is likely that the costs would be lower under this policy option as no efficacy in terms of smoking cessation would have to be proven.
As far as Governments are concerned, option 2 is expected to result in significant costs in terms of establishing and hosting the new authorisation system. The new structure would require important human resources to assess the products, ensure follow up and provide secretariat and support functions. This would only be partly compensated by the fees paid by industry for the actual assessment. The identification of assessment criteria (risk/benefit analysis) is somehow difficult. The pharmaceutical framework provides a safety/efficacy assessment where the efficacy is seen in terms of benefit from smoking cessation. As NCPs falling under this scheme do not claim to assist in smoking cessation, the efficacy assessment is therefore different in particularly as the safety consideration needs to take into account that nicotine is a toxic and addictive substance and therefore, per se, not safe. Another open question is whether the assessment can be limited to a current smoker using NCP or whether also uptake/use among other population groups, including non-smokers and risk for dual use should be taken into account. Another approach would be to assess the “attractiveness” in order to avoid that, for example, NCP with distinctive flavours are put on the market. Such an approach however, would not target the safety or quality of the product as such.

The impact on Governments’ tax revenues is expected to be similar to the impact described under option 1. A certain reduction of adverse health effects associated with NCP use is expected due to the authorisation requirement, but no impact is foreseen in terms of health costs linked to smoking attributable diseases. Consumers would enjoy a higher degree of health protection due to the prior authorisation requirement removing the most hazardous products from the market. However, nicotine as such is a toxic and addictive substance.

Social impacts

In terms of employment, a shift from SMEs to larger companies is expected under option 2, mainly as a result of the additional requirements and the fees to be paid for assessment/authorisation. However, this is already part of an on-going trend. It is likely that the targeting of young people could develop under option 2 as the products would be distinguished from pharmaceuticals through the establishment of a parallel authorisation scheme.

Health impacts

Option 2 would reinforce the character of NCP as “leisure products” rather than a product used in smoking cessation. This is inherent in the very nature of establishing a separate authorisation scheme. Option 2 is therefore expected to lead to increased consumption among young people, and people wishing to experiment with new products or to use alternatives to FMC where smoking is not allowed. Products authorised under the scheme suggested under option 2 are expected to be less interesting for people wishing to quit. On the other hand, a reduced health risk is expected following the requirement for prior authorisation as well as better control and knowledge of the NCP put on the market. No impact is foreseen in terms of tobacco related diseases as, to the contrary, NCP might develop into an entry gate to smoking initiation and dual use might prevent smokers intending to quit from quitting.

5.2.1.9. PO3: Subject NCP over a certain nicotine threshold to the medicinal products’ legislation and the remaining NCP to labelling requirements

NCP with a nicotine level over a certain threshold may only be placed on the market if they have been authorised as medicinal products on the basis of their quality, safety and efficacy, and with a positive risk/benefit balance under the medicinal products legislation. NCP with nicotine levels below this threshold will be subject to an adapted health warning. The nicotine threshold identified under this policy option should be established by considering the nicotine
content of medicinal products (NRTs) for smoking cessation which have already received a market authorisation under the medicinal products' legislation.

Delegated/implementing power to adapt the health warning and the identified nicotine threshold taking into account scientific and technical developments.

**Economic impacts**

Option 3 would consolidate current developments in Member States and ensure a harmonised approach for NCP with nicotine levels above a certain threshold, to be based on nicotine content in medicinal products authorised for smoking cessation. This would increase legal certainty and respond to Member States' requests for clarification. It is difficult to estimate with precision how many of the currently marketed NCP would have a nicotine level above the identified threshold and how many of those products would pass a prior authorisation, but considering that many consumers are reported to use the products for cessation/limitation purpose and presuppose a pharmacological reaction it is believed that the majority of products currently on the market would be affected. However, if a product is authorised under the medicinal products' framework, this would enable application of the mutual recognition procedure (MRP) and thus facilitate sale of the authorised products throughout the internal market. The approach also ensures equal treatment with NRT which are authorised smoking cessation aids with a positive risk/benefit analysis. Option 3 would also approximate the labelling requirements as regards NCP with nicotine levels below the identified threshold.

Option 3 would imply costs for economic players related to the authorisation scheme under the medicinal products' legislation. The costs will vary between Member States (e.g. 10,400 EUR-47,230 EUR per application in the UK, Netherlands, Germany and Denmark plus additional costs for each Member States where the applicant wants to enter thereafter under Mutual Recognition Procedure). However, this would be no change to the current situation. As set out in Annex 2, a significant number of Member States see NCP as medicinal products by function. The already on-going development towards bigger companies is expected to continue and at a higher speed due to innovation opportunities and a clear legal framework. Also, bigger companies are more likely to have resources for obtaining market authorisations. In addition, some adaptations in terms of composition and/or design might be needed to ensure compatibility with the medicinal products framework.

Products below the identified threshold would be subject to national legislation and could be placed on the market without prior authorisation. This would imply less cost for economic stakeholders, but on the other hand these “low-nicotine” NCP would not respond to users cravings for nicotine and could thus be less interesting from a marketing point of view. The potential of NCP in smoking cessation under this policy option (see below under health impacts) is expected to reduce, to some extent, indirect costs linked to tobacco consumption.

The impact on Governments is expected to be positive as NCP could develop their potential as smoking cessation aid and thus lead to improved health outcomes (less premature mortality, health care costs, productivity). Governments' tax revenues are expected to be neutral but they could no longer charge excise duties for NCP (for indirect impacts, see section 5.7 and Annex 5). The administrative burden is limited due to making use of an established authorisation regime with adequate fees. Consumers would have a more limited choice, but a higher degree of health protection. Some unauthorised low-nicotine NCP are likely to remain on the market, but these are expected to be of less interest for smokers trying to quit or smokers using NCP as an alternative where smoking is not allowed. In addition,

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316 Matrix 2012
consumers would be informed, through the labelling, that these products contain nicotine and can have adverse health effects. In terms of international impacts, it should be recalled that many of NCP are imported from third countries (notably China). However, given the estimated market value (section 2.1.3) this impact is considered limited overall. The option would be in line with the recommendations of the WHO Study Group on Tobacco Product Regulation that has recommended that these products are regulated as combination drugs and medical devices.\(^{317}\)

**Social impacts**

In terms of employment, a shift from SMEs to larger companies is expected. However, this is already part of an on-going trend. Option 3 would lead to improved equality within and between Member States, in particular as it would remove NCPs' appeal for young people. In particular NCP with characterising flavours are unlikely to be authorised under the medicinal products' legislation. NCP with nicotine contents below the identified nicotine content are not limited in terms of flavours.

**Health impacts**

Option 3 is expected to reinforce the character of NCP as smoking cessation rather than “leisure” product, in particular as the nicotine threshold would be set based on nicotine levels already authorised under the medicinal products' legislation. A significant drop in consumption of NCP among young people and people wishing to experiment with new products or to use alternatives to FMC where smoking is not allowed is expected. It would also provide the consumer with more appropriate information, including through the patient leaflet and better controlled marketing activities. Improved information about the adverse health effects associated with NCP is expected following the requirement for prior authorisation.

Under option 3, NCP would primarily target people wishing to quit smoking. Further research on the efficacy of NCP (notably electronic cigarettes) in smoking cessation would be encouraged and products put on the market would be safer and more adapted for smoking cessation. It is estimated that option 3 would lead to some reduction of tobacco related diseases and mortality due to the potential of NCP in smoking cessation under this option. Post-authorisation obligations, including pharmacovigilance, would provide further protection. It would put NRTs and NCPs on a level playing field.

**5.2.1.10. PO4: Subject all NCP to the medicinal products' legislation**

Only NCP that are authorised as medicinal products on the basis of their quality, safety and efficacy, and with a positive risk/benefit balance are allowed to be placed on the market. Otherwise, the placing on the market of NCP is prohibited. The authorisation procedure is described under option 3.

**Economic impacts**

Option 4 would go one step further than option 3 and provide a fully harmonised approach in the sense that all NCP placed on the internal market (regardless of the nicotine content) would need a prior authorisation as a medicinal product. However, as it is not certain whether

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\(^{317}\) World Health Organization (WHO) Study Group on Tobacco Product Regulation. Report on the Scientific Basis of Tobacco Product Regulation. WHO Technical Report Series, no. 955. Geneva: WHO; 2010. If the nicotine containing part of the NCP/the nicotine cartridge/refill bottle would be (was?) considered a medicinal product, the device designated to administer the nicotine would most likely be considered a medical device under Directive 93/42.
all NCP would fall under the medicinal products definition under Directive 2001/83 it is possible that some products would be excluded from the market.

The costs related to authorisation and the market developments are expected to be the same as under option 3. However, it is possible that some NCP would have to be taken off the market as they do not fit under the regulatory framework for medicinal products (no efficacy).

The impact on Governments and the international impact are very close to the impact under option 3. The consumer would have a more limited choice than under option 2, but a somehow higher degree of protection considering that only authorised NCP (regardless of nicotine level) would be put on the market.

**Social impacts**

The social impact is expected to be similar to the impact under option 3.

**Health impacts**

Similarly to option 3, option 4 is expected to reinforce the character of NCP as smoking cessation rather than “leisure” product. This is expected to lead to a drop in consumption similar to the impact under option 3.

5.2.1.11. **The views of stakeholders**

The tobacco industry and consumer focused advocacy groups have argued that NCP are different from tobacco products and therefore require a separate regulation. The argument of consumers’ choice was reiterated. Part of the pharmaceutical industry also supported this approach, while another part argued that NCP should be regulated as medicinal products. The European Respiratory Society (ERS) is opposed to the use of all (tobacco and) unapproved nicotine delivery products, including electronic cigarettes.318 The approach of subjecting NCP to the pharmaceutical legislation was also supported by health NGOs expressing concerns about the health risks of NCP. Among Member States responding to the public consultation, the views were divided. Some presented arguments for regulating NCP as medicinal products (which is already the situation in about half of the Member States) and others for the inclusion of electronic cigarettes in the TPD. Electronic cigarettes industry trade (ECITA) referred to self-regulation and claimed that no further regulation is needed.

5.2.1.12. **Comparing the options and preferred option**

In terms of effectiveness, regulating NCP under the medicinal products' legislation (partly policy option 3 helps to remove differential treatment between NCP and NRT (policy objective A1) and facilitate a level playing field (policy objective A2) as it would subject all economic stakeholders involved in NCP and NRT to the same rules. Regulating NCP under the TPD (options 1 and 2) would lead to the creation of different legal systems, associated with legal uncertainty. Policy option 1 would contribute to the achievement of objective B1 as it would result in harmonised labelling and the same is true to a certain extent for NCP under the identified nicotine threshold. Options 2, 3 and 4 would facilitate market surveillance (objective B2) as it would set up requirements for prior authorisation. Options 3 and 4 are well in line with policy consideration C1 as it would require prior authorisation and thus regulate NCP in the sense that only products which have passed a risk benefit balance would be allowed on the market. Option 2 would partly contribute to this consideration as also this option 4.

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option would require an authorisation scheme although the benefit criteria is less clear under this option. Adding health warnings and banning NCPs with characterising flavour (option 1) or authorising these products under pharmaceutical legislation (options 3 and 4) would help to remove NCP that could be attractive to young people (policy consideration C2). Regulating NCP under the TPD (policy options 1 and 2) or under the medicinal products' legislation (policy options 3 and 4) would also contribute to improved consumer information (policy consideration C4) due to labelling and/or the patient leaflet. In any event, for all products covered by the TPD, the pharmaceutical legislation is lex specialis.

Regulating NCP under the TPD (policy option 1) is a cost effective solution given that no prior authorisation is required and the products could thus make use of the already existing tobacco legislation. Requiring authorisation of NCP (policy options 2, 3 and 4) would imply additional costs, but these assessment costs would primarily be paid through fees from industry. Setting up of a new structure for authorisation under option 2 entails however important costs for public administrations, while options 3 and 4 take advantage of already existing structures under the pharmaceutical legislation.

Authorising NCP under the medicinal products' legislation (policy options 3 and 4) would be coherent with the medicinal products' framework, but option 4 would fully exclude some NCPs from the market as they might not fit under the legal framework.

Preferred policy option 3: Subject NCP over a certain nicotine threshold to the medicinal products' legislation and the remaining NCP to labelling requirements.

The preferred option would remove current legislative divergence between Member States and the differential treatment between NRTs and NCP, increase legal certainty and consolidate the on-going development in Member States based on the function of these products. It would encourage research and innovation in smoking cessation with the aim of maximising health gains.

It appears appropriate to focus, at this stage, on NCP above a certain nicotine threshold, in particular considering these products' similarities with already existing medicinal products for smoking cessation. In addition, most consumers use these products for cessation/limitation purpose, which presupposes a pharmacological reaction, i.e. a certain level of nicotine.

For NCP below the identified nicotine threshold, a less stringent measure appears more proportionate, in particular as there are not sufficient evidence that these products would fit under the medicinal products' legislation. The economic burden on industry is also taken into consideration in the proportionality assessment concluding that the suggested labelling requirement is sufficient for NCP below the identified nicotine threshold.

There are no less strict measures available to obtain the objectives. Subjecting all NCP to labelling requirements and ingredients regulation only (option 1) would have a positive impact on health, but to a lesser extent than the preferred option. A separate authorisation scheme (option 2) could address some safety concerns, but this option would not have the same potential in smoking cessation.

In terms of economic costs and burden weighed against the benefits, it is likely that some of the current NCP would be removed from the market or would have to change composition or design under the preferred option, but this burden is considered appropriate due to the health concerns associated with NCP. In any case, the economic impacts on industry under the
preferred option are lower than the additional burden of a separate authorisation mechanism (option 2) or the option of subjecting all NCP to the medicinal products’ regime (option 4).

The preferred option provides a clear added value as it contributes to a level playing field for operators involved in NCP and NRT. Only an initiative at EU level is capable of preventing further diversity and legal uncertainty.

1.1.14. **Herbal products for smoking**

5.2.1.13. **PO1: Subject all herbal products for smoking to labelling requirements under TPD**

Adapted health warnings are required for herbal products for smoking. Delegated/implementing power to adapt health warnings.

**Economic impacts**

Option 1 would ensure increased convergence of national rules and removal of some of the existing discrepancies on the internal market.

The benefit for economic players (cost savings compared to baseline scenario) would be similar to that described above for NCP because taking action at EU level in one go is less costly than consecutive changes at national level. In terms of sales, the current increase is expected to slow down due to stricter requirements and less demand due to more accurate health information. The expected slowdown in sales, would affect Governments' tax income, but taking into account the market size the decrease would be moderate if any. In return, the Governments would benefit from improved public health as herbal products for smoking are associated with some health concerns as indicated in the problem identification (section 2.2.1). Option 1 would have a positive impact on consumers as it would provide adequate protection in terms of health warnings while maintaining a wide choice of products.

**Social impacts**

Current increase in herbal products for smoking (see section 2.1.3) is likely to slow down, having a limited negative impact on employment. Option 1 would lead to improved and more equal protection of citizens throughout the EU, in particular by reducing some of the products potential of attracting/misleading consumers, including young people.

**Health impacts**

The labelling requirements under option 1 would contribute to increased awareness. It would also contribute to removing some misleading features, including wording such as natural, no additives etc. and better inform consumers of the health risks associated with herbal products for smoking e.g. herbal cigarettes and tobacco-free filling to water-pipes (sisha). This is expected to lead to reduced consumption which will, over time, result in less health problems related to herbal products for smoking.

5.2.1.14. **PO2: Phase out the placing on the market of herbal products for smoking**

Placing on the market of herbal products for smoking is phased out.

**Economic impacts**

Option 2 provides full harmonisation of national rules, but removes the current cross-border trade of herbal products for smoking on the internal market. Under the assumption that the product could not be regarded as medicinal products, option 2 would result in reduced sale
and possible closure of businesses involved in herbal products for smoking, mainly affecting SMEs involved in this sector. This limits the freedom to conduct business need to be balanced against health considerations. A reduction of Governments’ tax income as well as health care costs/concerns is expected under option 2, but due to the modest market size, this impact would be limited. Consumers would enjoy a higher level of health protection, but less choice.

Social impacts

Option 2 would have a negative impact on employment in herbal products for smoking, including possible regional impact, but redistribution is expected to take place. Option 2 would provide full and equal protection of EU consumers.

Health impacts

A ban would remove the current appeal of herbal products for smoking and reduce significantly the consumption, resulting in reduced health risks from herbal products for smoking. No impact on tobacco consumption is foreseen as it has not been demonstrated that the two categories are interchangeable.

5.2.1.15. The views of stakeholders

Health NGOs argued for the inclusion of herbal cigarettes into the TPD framework, referring to the harmful aspects following the combustion of these products. Most Member States which submitted contributions to the public consultation were either in favour of extending the scope of the TPD or did not refer to the question in a detailed manner. Other key stakeholders have not addressed specifically this area.

5.2.1.16. Comparing the options and preferred option

In terms of effectiveness, both regulating herbal products for smoking under the TPD or phasing them out (policy options 1 and 2) would create a level playing field and remove current national disparities (policy objectives A2 and A3). Option 1 would also partly fulfill objective B1 as it would unify the labelling rules, while this objective is not relevant as far as a full phasing out (option 2) is concerned. Phasing out herbal products for smoking (policy option 2) would provide regulation of products harmful to health and thus be fully in line with policy consideration C1, while option 2 (labelling rules) only partly would achieve this consideration. The labelling requirements suggested under option 2 would be well in line with policy consideration C4 as it would contribute to better information to consumers, while phasing out these products (option 2) would reduce the availability of herbal products for smoking and thus fulfil policy consideration C5.

Regulating herbal products for smoking under the TPD (policy option 1) would provide a cost-effective measure to better inform consumers about the health risks of these products, while phasing them out (policy option 2) would have more important consequences for economic stakeholders having to exit the market altogether (except export possibilities).

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<thead>
<tr>
<th>Preferred policy option 1: Subject all herbal products for smoking to labelling requirements under TPD</th>
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The preferred option would ensure a more homogenous development in the EU and create a safety net for consumers. The preferred option would also provide consumers and potential consumers with more appropriate information about the adverse health effects of herbal products for smoking and thus allow them to make informed choices.

A stricter measure (option 2) would further harmonise Member States’ legislations, and fully remove the adverse health effects associated with herbal products for smoking. However,
considering that the main health concern is the frequent misperception about the health risks and that herbal products for smoking entail more limited risks of developing an addiction compared to tobacco products, a full ban (option 2) is not considered proportionate. In addition, such a measure would be unnecessarily burdensome for economic stakeholders.

The objective of harmonising the labelling of herbal products for smoking is to ensure a harmonised approach throughout the EU and to create a level playing field for operators involved in these products while ensuring a high level of health protection. By definition, only an initiative at EU level is capable of ensuring a harmonised approach and thus added value. Unilateral actions by Member States can contribute to the protection of health, but would certainly result in divergences.

5.3. **PACKAGING AND LABELLING**

1.1.15. **PO1: Mandatory enlarged picture warnings**

| Combined warnings (picture plus text) of 75% displayed on both sides of the packages of tobacco products, presented in rotation. TNCO levels on the packages are replaced with descriptive information on content, emissions and risks. Display of cessation information (e.g. quit-lines, websites) is added to the packages. Tobacco products other than FMC and RYO are exempted (current TPD rules apply). Delegated/implementing power to remove/extend the exemption for these products if there is a change of circumstances and to adapt the health warnings. |

**Economic impacts**

Increased convergence of national rules on labelling and packaging, including mandatory pictorial warnings, would facilitate the functioning of the internal market in terms of cross border trade and improve the level playing field for economic stakeholders.

The actual one-off cost for introducing pictorial warnings for economic players (manufacturers) changing the labelling on FMC is estimated to range between 14,500 and 50,000 EUR per SKU and for RYO between 2,500 and 9,000 EUR per SKU (see section 5.1). However, considering the economies of scale referred to in section 5.1, it should substantially reduce the burden for manufacturers if changes are done in one go across the EU instead of individual adaptation to different national legislations. In particular SMEs could benefit from a harmonised approach in one go as they might have less resources to adapt to many different legal systems in case they would like to expand their activities to other Member States.

In addition, the package producers have reported that they have already made the investments necessary to comply with the expected regulation on pictorial health warnings. Therefore, there would be no additional costs for buying new equipment (e.g. printing equipment). Costs for disposal of old stocks are also disregarded as a transitional period would allow manufacturers to adapt the production process and retailers to manage their inventory accordingly.

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319 Matrix 2012. ESTA estimations for RYO (fine cut tobacco), Facts and figures for DG SANCO 2011-2012. In addition to the one-off cost, FMC industry has reported an increase of 1,3-1,5% in variable costs.

In terms of **running costs** (e.g. use of more expensive ink to produce combined instead of text-only warnings), the packaging industry has reported that the on-going costs related to pictorial warnings would be "slightly above" the costs for packages with text-only warnings. It has been estimated that costs could increase by 1.3-1.5% following the introduction of EU-wide pictorial warnings. Assuming this reported increase can be extrapolated across the four biggest tobacco companies, this would imply an annual cost increase for them of between 59 mEUR and 68 mEUR. Comparing these estimates to data from impact assessments in other jurisdictions (e.g. the UK, the US, Canada and Australia), compliance costs, in particular recurrent costs as presented here, appear to be relatively high. Taking into account that the printing is generally outsourced, the costs would not occur in the first place with tobacco manufacturers, but rather with the packaging and paper industry. If the packaging industry manages to charge higher prices from the tobacco manufacturers as has been suggested it is likely that the costs would be passed on by the tobacco industry to the subsequent levels of trade. In terms of **innovation**, the option maintains the possibilities of the tobacco industry to redesign packages and labels, albeit with limited space.

Under the provisions considered under option 1, **manufacturers** would be confronted neither with commoditisation of the market nor with a prohibition of the use of trademarks on the package. The measures proposed would continue to allow brand recognition albeit the space of the trademarks would be somewhat limited. However, this is considered to be a proportionate limitation of the right of property (trade marks).

Overall, the option would have a neutral or even positive direct effect on economic players (manufacturers) as possible minor increases in variable costs would be counterbalanced by savings in familiarisation costs due to implementation in one go. The same applies to the packaging industry. The indirect impact in terms of reduced profit following the estimated drop in consumption is further described in section 5.7 (and Annex 5).

As all Member States already have cessation services in place, only a limited additional burden is expected for **Governments** as a result of the introduction of references to such cessation services (quit-lines or other services) on the packages, e.g. in form of an increase in call volume. Outsourcing this role to NGOs with an expertise in providing such services

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321 Idem. This is in line with the estimates given in the context of industry interviews carried out, Matrix 2012.
322 Matrix 2012
325 There is no absolute right to use a trademark and the Court has confirmed the validity of Articles 5 (health warnings) and 7 (ban on misleading descriptors) of the current TPD which constitutes a limitation to use the trademarks. BAT case: C-491/01 The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd.[2001] ECR I-11453, para 131-132. Although not precedent for the Union, , the High Court of Australia ruled, on 15 August 2012 the Government's tobacco plain packaging legislation to be constitutionally valid.
could be an effective and efficient approach to reduce costs. Other administrative burdens associated with this policy option are rather limited. For indirect impacts (e.g. improved public health and reduced tax revenues), see section 5.7 and Annex 5. Furthermore, the adoption of uniform rules across Member States could reduce the burden of developing individual solutions. The proposed measure would improve consumer protection across the EU, including providing assistance to smokers wishing to quit (e.g. quit lines, web sites), protecting consumers from misleading information on the health impact of smoking (replacement of TNCO levels) and providing appropriate information (strengthened labelling). At the same time, it is expected that the measures would affect neither the range of products available on the EU-market nor their quality. It can be expected that the tobacco industry will try to pass on the additional compliance costs to consumers, but these are expected to be very small if any (see above). In terms of international impact, the labelling requirements under option 1 would bring the EU more in line with international developments and the FCTC commitments.

Social impacts

The impact on employment would follow the expected drop in prevalence. This is further described in section 5.7 and Annex 5, but in the light of the input/output model, money not spent on tobacco is spent on other goods and services. No major impact is expected on package manufacturers, as these generate only 10% of their turnover with tobacco industry. Positive impact is expected in terms of equality. Pictorial warnings appear to be more effective than texts among persons with lower levels of literacy and particularly in young people. References to cessation services would have a positive impact on vulnerable groups.

Health impacts

Overall, it is expected that option 1 would make a substantial contribution to the expected decrease in consumption and prevalence over five years (see section 5.7 and Annex 5).

Evidence suggests that the effectiveness of health warnings depends on their size, location and design. In general, prominent pictorial warnings placed on the front of the packages are seen to be the most effective in increasing perceptions of risk and promoting behavioural change. Enlarged picture warnings on both sides of the package are expected to result in greater noticeability and salience for consumers, stronger beliefs about the health risks of smoking and increased quit attempts.

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smoking, as well as increased motivation to quit smoking.\textsuperscript{330} There is evidence that the warnings are more visible if placed on the front panel in the upper part of the package.\textsuperscript{331} Furthermore, messages and images which elicit strong emotional reactions are considered particularly effective.\textsuperscript{332} Finally, evidence suggests that the impact of health warnings tends to decrease over time (wear-out effect) whereas regular rotations and updates of the health warnings and messages, already prescribed in the current TPD, are associated with increased effectiveness.\textsuperscript{333}

Research comparing pictorial warnings with text-only warnings from several countries (e.g. Australia, France, Spain, the UK and US) also demonstrates that pictorial warnings increase consumers’ awareness of warnings, knowledge and credibility of health risks, depth of processing and also cessation behaviours such as forgoing FMC, quit intentions and quitting.\textsuperscript{334} They also contribute to reduced consumption as indicated above.\textsuperscript{335} Research conducted for the Canadian Government also found that larger warnings are more effective at eliciting negative reactions, conveying information about the health risks of smoking and reducing consumption.\textsuperscript{336} Consumers living in countries that have introduced graphic tobacco health warnings have a greater knowledge on the health effects of smoking\textsuperscript{337} and may have

\begin{thebibliography}{99}
fewer disparities in health knowledge across educational levels. In countries with picture warnings on one side of the package the products are typically displayed in such a way that the side of the text warnings is visible to consumers.

The replacement of TNCO levels with descriptive information on contents/emissions under option 1 would help to better address a possible confusion among consumers regarding possible effects of the product (see problem identification 2.2.2). The inclusion of concrete quitting information is strongly supported by smokers and has been shown to increase the use of these services dramatically.

1.1.16. PO2: PO1 plus harmonise certain aspects of packets and prohibit promotional and misleading elements

| Option 1 plus: |
| 1) The tobacco labelling and packaging and the tobacco product itself shall not include any promotional and misleading elements (e.g. misleading colours, symbols, slim FMC), |
| 2) setting certain requirements for packages (e.g. cuboid shape, minimum number of and FMC per package) as well as for the size of the warnings |

Member States are allowed to regulate the area not regulated by the TPD or other Union legislation, including adopting provisions providing full standardisation of packaging of tobacco products (i.e. plain packaging) as far as these provisions are compatible with the Treaty. The Commission will report on experiences gained with respect to surfaces not governed by the TPD five years after the transposition of the TPD.

For Point 2) Tobacco products other than FMC and RYO are exempted (current TPD rules apply). Delegated/implementing power to remove the exemption for these products if there is a change in circumstances.

Economic impacts

Option 2 would further advance the functioning of the internal market following e.g. harmonisation of additional elements including the removal of promotional/misleading elements where some Member States already have taken actions (2.2.2). The standardisation of the size of the health warnings as well as the shape of the package would contribute to effective display of the warnings and thus maximise the effect of the TPD.

Option 2 would further reduce compliance costs for tobacco manufacturers and it would result in even larger economies of scale, including standardised package size. The

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harmonisation of legislation in one go under option 2 rather than consecutive changes at national level would be cost efficient for the industry. The prohibition of promotional and misleading elements, including inserts prepared by several manufacturers for marketing reasons\textsuperscript{340} would lead to some cost savings. For indirect impacts linked to consumption, see below references to section 5.7 and Annex 5. According to industry reports, the total ongoing administrative burden for producing inserts from reduced factory efficiency (i.e. higher production cost per unit of output) and the costs of the insert itself (i.e. paper and printing ink) would be likely to be between 42.2 and 75.5 mEUR a year in the EU-27.\textsuperscript{341} Regulating the appearance of FMC should not place a major burden on manufacturers as most FMC currently on the market comply with a standard format, but products with a misleading size ("slim") would be affected. No impact beyond option 1 is expected on package producers and retailers. For tobacco products other than FMC and RYO the rules of the existing TPD could continue to apply (e.g. size and location), but with some minor modifications of the health warnings.

The further harmonization of packages and labelling requirements will support and save costs for Governments who would not have to invest in developing their own legislation. The measures reduce the administrative burden as the measures are self-executing and mutually reinforcing (package shape/sized pictures). For the indirect impact on Governments following the estimated additional drop in consumption reference is made to section 5.7 and Annex 5. The protection of consumers would be reinforced by regulating promotional and misleading aspects of the packaging (see glossary). Option 2 would complement those measures foreseen under policy area 3 (ingredients), which aim at banning characterising flavours (see glossary), as these provide tobacco products with a misleading taste and smell. As in option 1, it is expected that the measures would affect neither the range of products available on the EU-market nor their price and quality. Option 2 provides a further step towards implementing the FCTC guidelines. It does not prevent Member States from regulating surfaces not regulated by TPD or other legislation and thus implement full standardisation (plain packaging).

**Social impacts**

The impact on employment following the estimated drop in prevalence is further described in section 5.7 and Annex 5, including redistribution effects. No additional impact is foreseen for package manufacturers. Option 2 is expected to strengthen the effects of option 1 on equality, in particular as it would further limit the possibility of developing packages and FMCs particularly appealing to young people.

**Health impacts**

The envisaged measures under option 2 are expected to significantly contribute to the projected decrease of consumption and prevalence. Reducing the packages' potential to mislead consumers would have a positive impact on awareness.

As illustrated in the problem identification (section 2.2.2), many studies indicate that package design influences the perception of risks and has the potential of misleading consumers.

The measures proposed under this option are expected to reduce these problems. A ban on promotional/misleading elements would reduce the packages' potential to mislead consumers and increase the noticeability of health warnings. Setting certain requirements for package


\textsuperscript{341} RAND 2010
appearance and regulate the size of health warnings would reduce the promotional and misleading potential of the package, ensure that labelling requirements are not undermined and that the enlarged picture warnings can be appropriately and fully displayed on the package in order to ensure their full effectiveness. A minimum number of FMC in each package and a minimum weight of tobacco in RYO packages would ensure that the size of the package allows a sufficient space for health warnings.

1.1.17. PO3: PO2 plus full plain packaging

Option 2 plus: standardised colour, font, size and position of brand name and brand variant on packages (plain packaging) and a readable health warning on each FMC stick.

Other tobacco products than FMC and RYO will be exempted from the requirements (current TPD rules apply). Delegated/implementing power to remove the exemption if there is a change in circumstances.

Economic impacts

The precise economic effects of plain packaging in real life are difficult to quantify at present due to lack of empirical data and experience with plain packaging in Member States or other countries.

Option 3 would maximise the effects on the internal market. Product would have a homogenous appearance throughout the EU and all discrepancies between national legislations would be removed. Again, the direct (compliance) costs for economic stakeholders would be further reduced under option 3. The industry claims that the initial burden of introducing mandatory plain packaging would be the same as the burden of mandatory pictorial warnings, i.e. 32.5-125.4 mEUR. However, this does not appear to be realistic. It is rather expected that companies would spend far less resources on the development of packages, considering that their possibility to develop individual packages are excluded under this option. Therefore, it is estimated that this option would result in overall cost savings as packages would remain the same and regular changes would become obsolete. Also, the production of a plain package is expected to be cheaper as e.g. fewer, simpler and less expensive colours are used on the package itself. Finally, the harmonisation of legislation in one go would be cost efficient for the industry. In terms of on-going costs, the full standardisation package appearance would also lead to economies of scale in the production of packages and FMC. Printing a health warning on a cigarette stick would result in some on-going costs for the printing process (colours, maintaining equipment), but no quantitative figures are available.

On the other hand, it is expected that plain packaging would result in reduced possibilities of brand differentiation, affecting in particular high margin/premium brands which could, over

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342 According to self-reported data of one major cigarette producer in Europe that provided quantitative estimates to RAND Europe (RAND 2010). The company disclosed that the initial administrative burden would be approximately 40 to 45 million EUR for the company if considerable time were not allowed for compliance. This translates into 18,100–20,400 EUR per SKU (no data available on costs for printing a warning on a cigarette stick).

343 According to information from one tobacco product manufacturer, the cost of a general brand re-design which tends to happen about every two to three years is more than 20,000 EUR per SKU (Matrix 2012), which would be saved by manufacturers under this option.

time, result in price competition and commoditisation of the market. Whilst it would be very difficult to establish new premium brands, entry barriers for non-branded products would be lowered.345 Considering the high degree of market concentration observed at present there is an increased risk of collusion between producers of premium brands. The industry has argued that plain packaging will increase illicit trade. However, no convincing evidence has been submitted and the argument appears counter-intuitive taking into account that counterfeiters try to benefit from the brand recognition associated with premium brands (currently recognisable from their trade marks which is not possible with plain packaging). Anyway, the actual and alleged additional risk of illicit trade is addressed in policy area 5 (traceability and security features). Regarding FMC papers/sticks, representatives from the fine paper industry confirmed that tipping paper is subject to very frequent changes346 and it is therefore not expected that this would have any significant negative impact.

As regards other economic stakeholders, package producers could be slightly affected by the provisions under this option, as the production of the packages would become cheaper and could result in a drop in price per package, and possibly also in profit margins. However, this impact would be limited, in particular as package producers supply various sectors.347 Retailers have claimed that plain packaging would make it more difficult for sales persons to identify packages. Therefore, management of stock and identification of products would become more cumbersome, increase transaction time and slow down the selling process, resulting in complaints from customers and higher business costs, affecting SMEs and small family enterprises disproportionally.348 However, this problem is expected to be solved after an initial period of familiarisation, while staff is getting used to the new system, and by an appropriate reorganisation of the display.

For the indirect impact on economic players and Governments following the estimated additional drop in consumption, reference is made to section 5.7 and Annex 5. It is very likely that any shift from premium to low cost brands. However, Member States are likely to counter this with tax increases given the fact they can, in the context of the mixed structure of the EU excise duty system, make use of both the specific and the ad valorem component as needed.

Consumer protection would be further reinforced by regulating additional promotional and misleading aspects of the packaging and highlighting the health warnings, including also a warning on the stick. Consumer choice may be limited in the long run as new premium brands would have more difficulties to enter the market and second tier brands might exit the market. On the other hand, entry barriers of non-branded products would be lowered. No relevant impact on quality and price are expected as potential price drops could be compensated by tax increases. Option 3 is fully in line with the FCTC guidelines. Legal proceedings have been initiated following Australia’s adoption of a similar measure as of 1 December 2012. While the High Court of Australia ruled the Government's tobacco plain packaging legislation to be constitutionally valid on 15 August 2012,349 discussions are still on-going in the WTO.

347 Related figures were not delivered by the industry despite repeated requests.
348 Information provided by EU retailer association, CEDT, to DG SANCO
following requests for consultations by Ukraine, Honduras and the Dominican Republic. Legal proceedings are also on-going under the Australia-Hong Kong Bilateral Investment Treaty.

**Social impacts**

The impact on employment due to further reduction in consumption is described in section 5.7, including the input / output model. No major impact on package manufacturers is foreseen as they are also serving multiple other industries. Option 3 would, overall, improve equality, in particular as it would further limit the possibility of developing packages appealing to young people. However, standardising the font and size of brand and product name could to a certain degree make products less easily identifiable to people with lower levels of literacy.

**Health impacts**

Option 3 is expected to strengthen further the effects of option 2. Although, no studies based on real life experiences are available at this stage, many recent studies indicate that plain packaging not only increases the noticeability and effectiveness of health warnings, but also reduces substantially the attractiveness and appeal of tobacco packaging, the product, particular brands, the user and smoking (both to smokers and potential smokers) as well as false beliefs about the risks associated with different brand variants. These effects are expected to be particularly strong in young people and for initiation. For example, a recent eye tracking study showed that plain packaging increased the salience of health warnings among non-smokers and light (i.e. non-established) smokers. These results have recently been replicated in adolescents. As a consequence, plain packaging may help to reduce tobacco consumption and smoking prevalence, in particular by discouraging young people from taking up smoking, by reducing tobacco consumption among young adult smokers, by keeping the "in-between-category" of occasional smokers from

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becoming regular smokers and by encouraging - in particular young and occasional – smokers
to give up their habit.355

1.1.18. The views of stakeholders

The tobacco industry has argued that bigger pictorial health warnings or plain packaging have
no effect on smoking prevalence, that plain packaging increases illicit trade and that it
undermines their intellectual property rights (trademarks). On the other hand, at least one of
the four big FMC manufacturers expressed no major concerns with the introduction of
mandatory picture warnings, replacement of TNCO-levels display and mandatory printing of
cessation services, although the importance for keeping appropriate space for trademarks was
underlined.356 Concerns about increased illicit trade following a plain packaging regime were
also expressed by suppliers to the tobacco industry (packaging, fine paper, ingredients
industries), but again they were not able to fully substantiate their arguments. Cigar
manufacturers have referred to the specificity of cigars as an argument against strengthened
labelling requirements and highlighted that SMEs would be affected disproportionately by
stricter regulation/plain packaging. Health NGOs have argued that large mandatory pictorial
warnings and plain packaging reduce the attractiveness and do not facilitate illicit trade. Most
of the Member States responding to the public consultation were in favour of enlarged
mandatory pictorial warnings, while the positions on plain packaging were more diverse.

1.1.19. Comparing the options and preferred option

Full plain packaging (policy option 3) would be most effective in terms of removing national
disparities (policy objective A3), but requiring mandatory pictorial warnings and/or
harmonising package shape and prohibiting promotional elements (options 1 and 2) would
also help to achieve this aim, albeit to a lesser extent. Additionally, full plain packaging
(policy option 3) would help to reduce administrative burdens by fully unifying labelling rules
(policy objective B1), whereas requiring mandatory pictorial warnings and/or harmonising
certain aspects of the package shape and prohibiting promotional and misleading elements
(options 1 and 2) would contribute to a lesser extent to this objective. Harmonising certain
aspects of the package shape and banning promotional elements or introducing full plain
packaging (policy options 2 and 3) would remove the misleading potential of packages from
the market (policy consideration C2). Introducing bigger mandatory pictorial warnings (policy
options 1-3) would help improve consumer information (policy consideration C4).

The assessment above has illustrated that all options 1-3 would lead to cost savings compared
to status quo in terms of compliance costs and because action is taken at one go at EU level
rather than many consecutive changes by Member States. Overall, however, it is expected that
plain packaging (option 3) would also reduce the possibilities for brand differentiation, in
particular affecting high margin/premium brands and impact more negatively on
revenues/profits due to the drop in consumption (see section 5.7 and Annex 5), which is
expected to be more important than under options 1 and 2.

prepared for the UK Department of Health. Stirling: Centre for Tobacco Control Research; 2009. Germain D,
Wakefield MA, Durkin SJ. Adolescents’ perceptions of cigarette brand image: Does plain packaging make a
Estimating the impact of pictorial health warnings and “plain” cigarette packaging: Evidence from experimental

356 Minutes from the meeting with FMC manufacturers, 2 December 2011:
Preferred policy option 2: Mandatory enlarged picture warnings plus harmonise certain aspects of packets and FMC appearance and prohibit promotional and misleading elements.

The preferred option addresses the current heterogeneous development in Member States, in particular as regards pictorial warnings and cessation information where different regimes affect the internal market negatively. The suggested standardisation of the package appearance will ensure effective display of the health warnings. The option provides an ambitious and balanced approach taking into consideration the concerns of some stakeholders. It addresses in particular the concerns of the tobacco industry as it leaves a certain space on the package for manufacturers to display their trademark. The limitation of the product scope to FMC and RYO in a first stage would also take into consideration the concerns expressed by cigar manufacturers (often SME’s). This is justified also because cigars (and pipe tobacco) are primarily used by adult smokers (see section 2.2.1). The option is based on new evidence showing that bigger and pictorial warnings are more effective and current indications of TNCO level and other aspects of packaging and labelling are misleading. The exact size of the warning (75%) has been suggested after thorough analysis of scientific evidence and international experience and developments as well as the impact on economic stakeholders.

There is no less stringent measure available to reach the objective of improving the internal market while protecting public health. All elements included in option 2 would contribute to the objective, but none of them can, on their own, be expected to achieve the objective as effectively as the package of measures proposed in the preferred option. It is assumed that the measures proposed in the preferred option reinforce each other. Finally, some of the measures proposed need to be regulated jointly to avoid circumvention of legislation (e.g. picture warnings and package shape). Also, the costs associated with the preferred options are considered proportionate compared to the overall benefits. On the other hand, a stricter measure (option 3, full plain packaging), is expected to achieve the policy objectives even more effectively, but, given the current lack of real life experience, pending legal disputes regarding the plain packaging and serious concerns expressed by some stakeholders it appears most appropriate at this stage to choose option 2, while providing an opportunity to evaluate the situation after 5 years with a view to considering further standardisation (plain packaging). The option would not prevent Member States to regulate surfaces of the packages not covered by TPD or other legislation and thus implement full standardisation (plain packaging) provided it is compatible with the Treaty.

As illustrated above, only an EU action can bring some of the current provisions in line with international and scientific development as Member States are prevented from taking actions on their own (e.g. TNCO levels display). Other aspects of labelling (e.g. pictorial warnings) are associated with a lot of diversity and only an EU action can ensure a homogenous development facilitating the functioning of the internal market.

357 75% on both sides in Canada, 30% and 90% in Australia and New Zealand, 80% of both sides in Uruguay, 60% and 70% in Mauritius, 30% and 100% in Mexico.
358 Plain packaging entered into force in Australia as of 1 December 2012.
5.4. REPORTING AND REGULATION OF INGREDIENTS

1.1.20. PO1: Common reporting format on a voluntary basis. Prohibit toxic, addictive and attractive additives in tobacco products.

Member States are free to decide whether they oblige manufacturers to report additives in tobacco products. If Member States decide to make the reporting obligatory, the common reporting format must be used. Member States shall prohibit additives based on the general criteria toxicity, addictiveness and attractiveness.

Economic impacts

A harmonised reporting format would remove current disparities and facilitate the monitoring and analysis of ingredients data across the EU, but a voluntary reporting regime would provide an incomplete picture of ingredients used across the EU, which is not beneficial for the functioning of the internal market. Regarding ingredients regulation, Member States would have to ban additives which increase toxicity, addictiveness and attractiveness. This is expected to have a positive impact on the internal market, especially as a result of increased convergence as regards additives with toxic and addictive properties. Following the obligation to ban toxic and addictive additives, Member States are also expected to re-evaluate their current ingredients lists (where such lists exist) and update those according to scientific developments. Also this, would remove some of the current divergence. In terms of “attractive” additives, the effect is expected to be more limited as this concept is more subjective and no further guidance would be given in the legislation. Member States could be expected to take the FCTC guidelines for implementing Article 9 and 10 as guidance, but they give significant discretion to Parties and are expected to be interpreted in different ways.

The optional and harmonised reporting system would lead to cost savings for economic stakeholders, Governments (in particularly those opting for no reporting) and the Commission as it would reduce the compliance costs compared to the baseline scenario. Also the harmonised ingredients regulation under option 1 is expected to lead to some cost savings compared to the baseline scenario due to the increased convergence in terms of additives allowed or banned in tobacco products. However, as indicated above, some discrepancies can still be expected under option 1, in particular in terms of “attractive” additives where Member States are expected to interpret the concept differently. For the indirect impact on economic players and Governments following the estimated marginal drop in consumption, reference is made to section 5.7 and Annex 5.

Under option 1, consumers would benefit from a somewhat increased protection due to the ban on toxic, addictive and attractive additives. The consumer choice is expected to be remain stable. Even if some of the tobacco products currently on the market would have to be reformulated, it is likely that alternative additives or mixture of additives would be used in order to maintain the product or a similar one on the market.

Social impacts

The indirect impact on employment is expected to follow the limited drop in consumption. As indicated above, it is likely under option 1 that the rather general criteria, and in particular on attractiveness could be interpreted differently in Member States. This could have a negative impact on equality between Member States, i.e. lead to different level of protection.

359 For definition, see glossary
As the criteria suggested under option 1 are related to the additives as such and not the tobacco products, it is likely that the producers would find alternative additives (instead of those banned because of their attractiveness) to create a similar product. Therefore, it is expected under option 1 that attractive tobacco products, including fruit and candy-flavoured FMC would continue to be marketed, at least in some Member States. This would particularly affect young people who are more vulnerable in relation to these products.

Health impacts

Option 1 is expected to remove some of the most hazardous additives from tobacco products. However, as explained previously in this impact assessment, tobacco consumption as such is linked to a number of health risks which would remain despite the regulation foreseen under this policy option. It is also likely that an additives-based approach only in terms of attractiveness is less effective than a product-based approach (see option 2), in particular considering the risk that alternative additives could be used to produce a similar (distinctive) flavour.

1.1.21. PO2: Mandatory reporting in harmonised format. Prohibit tobacco products with characterising flavours and products with increased toxicity or addictiveness

Manufacturers are obliged to electronically report ingredients (glossary) of tobacco products in accordance with a common format and provide supporting data (e.g. marketing reports). Fees charged by Member States for handling the information submitted to them shall not exceed the cost attributable to those activities. Placing on the market of new or modified tobacco products must not take place before the submission of ingredients data. Reported data, excluding confidential information, is published. Delegated/implementing power to specify the reporting format.

Tobacco products with characterising flavours (glossary) are prohibited (this is similar to the US model). Test panels assist in the decision making process. Additives associated with energy and vitality (e.g. caffeine and taurine) or creating the impression that products have health benefits (e.g. vitamins) are prohibited. No flavourings are allowed in filters, papers or packages. Tobacco products with increased toxicity or addictiveness shall not be placed on the market. Member States shall remove from the market tobacco products that include ingredients not complying with REACH. Delegated/implementing power to set limits for additives imparting a characterising flavour, toxicity and addictiveness.

Tobacco products other than FMC, RYO and STP (i.e. cigars, cigarillos and pipes) are exempted from the prohibition of products with characterising flavour and the prohibition of additives associated with energy and vitality or health benefits. Delegated/implementing powers to remove the exemption for these products are foreseen if there is a substantial change of circumstances.

Economic impacts

Mandatory reporting of additives in combination with a harmonised format would ensure a level playing field and facilitate analysing and monitoring of data. Also the ingredients regulation suggested under option 2 would further harmonise national legislations, prevent a patchwork of national regulations, facilitate cross-border trade and thus improve the functioning of the internal market. The approach in terms of attractiveness, addictiveness and toxicity would rather focus on the tobacco product itself than the additives used in option 1. In addition, some easily determined additives, such as vitamins and caffeine would be banned throughout the EU. This would result in less discrepancy for Member States. Also, implementing/delegated powers are foreseen to ensure a consistent approach. Test panels would assist in determining products with characterising flavour, which would further
increase convergence. Overall, option 2 would increase further the positive impact on the internal market compared to option 1.

The EU guide on harmonised reporting format and the already developed tool for data submission (EMTOC) would provide a basis for a mandatory harmonised reporting system. The costs for introducing such a system on a mandatory basis would therefore be marginal and largely off-set by the savings generated by the use of one single format across the EU. A robust electronic system as the one suggested would also address the concern expressed by tobacco manufacturers as regards trade secrets as publication for general public could take into account legitimate confidentiality requests. If Member States charge a fee for the reporting, the economic players have indicated in the context of the public consultation that they would be ready to participate to the financing of an electronic reporting system. Additional market studies and scientific studies (e.g. on toxicity) would need to be submitted.

In terms of ingredients regulation, the economic actors (the tobacco industry) would benefit from cost savings under option 2. For products sold across various Member States the decision is adopted once across the EU rather than several times consecutively, which reduces the compliance costs, including understanding of the new requirements and possible reformulation of products. In order to quantify these costs, a question was submitted to four major FMC manufacturers to know the average cost per brand of changing the composition of cigarettes. The replies provided were incomplete which made it difficult to quantify the extra costs for changes of ingredients regulation in a given Member State. One company stated, however, that a major product redesign would cost in excess of 1 mEUR per brand. Costs included in this figure are development work, specification maintenance, pre-trials, pilot plant trials, analytical work, as well as the cost impact on stock holdings and factory efficiency. It is estimated, under option 2, that a limited part of the market would be affected by the change (including the menthol FMC counting for 4% of the market). Also the ban of flavourings added to the filters, paper and packages would affect the (limited) segment of products currently including such flavourings. Moreover, the regulation of products with increased toxicity or addictiveness is harmonised under this policy option, which could result in limited additional costs savings for economic stakeholders. The possibility for the Commission to adopt harmonised rules (delegated/implementing acts) should further reduce compliance costs compared to option 1. The production costs would also decrease under option 2 as less ingredients are expected to be used. Ultimately, option 2 would allow manufacturers to produce one product which could be sold across borders in all Member States. The measure would primarily affect a specific segment of the market, namely products with characterising flavours or additives giving the misleading impression of energy or vitality (such as caffeine and taurine) or associated with health benefits (such as vitamins).

Menthol-flavoured FMC are more common than other FMC with characterising flavours. They account for 4% of all FMC sales over the past ten years ranging from 25% in Finland

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360 CECCM submission to the public consultation on the possible revision of the Tobacco Products Directive 2001/37/EC.
361 Matrix 2012. One company stated, in the context of an interview carried out by Matrix, that a single regulation across Europe might be less expensive than Member State level changes with different limits across each of the EU Member States, as long as ingredient regulation is proportionate and evidence based.
362 Certain flavourings are used in relatively high quantities, such as cocoa and liquorice, without necessarily resulting in a product with a characterising flavour, whilst other products includes flavourings in lower quantities giving the product a characterising flavour (distinguishable taste or aroma). Often, the characterising flavour is indicated on the package. Tobacco products overtly marketed as containing additives (e.g. menthol cigarettes) command a relatively small market share in EU (SCENIHR 2010)
and 0.1% in Greece.\textsuperscript{363} As indicated under health impacts below, it is estimated that flavours have an effect on smoking initiation.\textsuperscript{364} It is impossible to predict with certainty the response of already established smokers of menthol FMC if these products were not allowed. However, it appears likely, in particular taking into account addictiveness that a significant part of these smokers would switch to traditional FMC without characterising flavours. Sales lost from menthol cigarettes would therefore be partially off-set by sales of other FMC or by efforts to acquire the products outside the EU or illicitly. However, some current menthol smokers are also expected to quit others are expected not to start. The impact on growers is limited to the drop in consumption and there is no risk of distortion between growers of different tobacco types. The impact on flavouring suppliers is expected to be limited as they normally supply multiple actors and multiple flavourings. The indirect impacts on the economic stakeholders following the expected drop in consumption are further described in section 5.7 and Annex 5.

Regarding the reporting obligations, the main impact expected for Governments is that enforcing and verifying the reporting obligations is facilitated. National authorities would also be able to charge a fee for the ingredients reporting. Regarding ingredients reformulation, however, there may be some additional administrative burden for Governments and the Commission when analysing the market studies submitted by industry or when taking decisions on characterising flavours. From an international perspective, option 2 is well in line with the FCTC guidelines on ingredients and similar to the US approach. All products with characterising flavours (e.g. menthol and clove) are treated the same way to avoid unjustified differences of treatment.\textsuperscript{365} The indirect impacts directly linked to consumption/prevalence are further developed in section 5.7 and Annex 5.

\textbf{Social impacts}

In terms of employment, the impacts of option 2 are in line with trends outlined above for the economic stakeholders, i.e. they would broadly correspond to the foreseen reduction in consumption (see section 5.7 and Annex 5). This applies to upstream suppliers (e.g. growers and producers of additives), tobacco industry and distributors. Reference is also made to the redistribution effect which is based on the input/output model (see Annex 5). The option would not affect additives essential for the manufacturing of tobacco products (other than those imparting a characterising flavour). It would not discriminate between different tobacco varieties. Option 2 would provide improved and more equal protection due to the removal of certain appealing products. This is expected to have a particular impact on young people.

\textbf{Health impacts}

As described in the problem identification, scientific studies have demonstrated an influence of flavourings on smoking initiation.\textsuperscript{366} Tobacco products may also be designed in such a way that they are easier to start smoking with. This may be attained by making it easier to inhale the smoke in the lungs and by creating a sweeter, milder or “colder” smoke. By reducing and changing the harshness of the smoke, special target groups may be reached.\textsuperscript{367} In a number of

\textsuperscript{363} Matrix 2012. Based on Euromonitor (volume ) data for 24 Member States (no data available for Cyprus, Malta and Luxembourg)

\textsuperscript{364} SCENIHR 2010

\textsuperscript{365} See WTO Appellate Body, AB-2012-1, United States – Measures Affecting the Production and Sale of Clove Cigarettes (DS406)

\textsuperscript{366} SCENIHR 2010

countries, sweet and tasteful tobacco products are the most preferred tobacco products among children and adolescents as well as experimenting smokers.\textsuperscript{368} A study by Hersey et al., using data from the National Youth Tobacco Survey in the US, found that menthol FMC use was significantly more common among newer, younger smokers.\textsuperscript{369} A recent study analysing data of more than 47,000 US-pupils US found a greater risk of progression to regular smoking and nicotine dependence for those who start smoking menthol cigarettes compared to those starting with non-menthol cigarettes.\textsuperscript{370} The US FDA Tobacco Products Scientific Advisory Committee has also confirmed, on the basis of the extensive review of all available information, that the evidence was sufficient to conclude that it is more likely than not that the availability of menthol FMC increases the likelihood of experimentation and regular smoking beyond the anticipated prevalence if such FMC were not available.\textsuperscript{371} Overall, it is expected that option 2 would result in lower appeal and increased \textit{awareness}. Additives that give the impression that FMC are less harmful, contain healthy properties or are associated with energy and vitality (including fruits, vitamins and caffeine) would no longer be allowed and if new attractive products appeared on the market, the Member States and the Commission would be able to react, if necessary, through a delegated/implementing act. This would contribute to the projected decrease in \textit{consumption}. The option is likely to reduce smoking uptake rather than affecting habits of established smokers although a certain impact is also expected for established smokers. Option 2 would be further reinforced by stricter requirements in the area of packaging and labelling (see policy area 2), namely by the removal of misleading and promotional elements under that policy area. Over time, it will result in reduced morbidity/mortality from smoking and a higher level of well-being.

\begin{center}
1.1.22. \textbf{PO3: Mandatory reporting in harmonised format. Prohibit all additives not essential for manufacturing (similar to Canada model)}
\end{center}

Ingredients reporting as in option 2. In terms of ingredients regulation, all additives in tobacco products, except those essential for manufacturing are prohibited. Maximum limits are set for sugar and sweeteners.

Other tobacco products than FMC, RYO and STP are exempted from the ingredients regulation. Delegated/implementing power to remove the exemption for these products if there is a change of circumstances.

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**Economic impacts**

As regards ingredients **reporting**, the impacts would be similar to option 2, although the volume of reporting could be reduced following the restriction of ingredients allowed.

Regarding ingredients regulation, option 3 would further improve the functioning of the **internal market** due to the removal of the remaining discretion for Member States. Products could then circulate freely across borders without additional health concerns. Compared to the previously described options, **manufacturers** would be faced with significantly higher costs associated with reformulation and removal of products from the market. Using the same cost estimated as under option 2 means that a major product redesign would cost in excess of 1 mEUR per brand, but compared to option 2, most brands across the EU are expected to be affected by option 3. The costs associated with a change at EU level would however still be lower than consecutive changes at national level.\(^{372}\)

Option 3 would also impact negatively on product differentiation opportunities for the industry and is therefore likely to reduce turnover and profits, see section 5.7 and Annex 5. Option 3 would have a particular impact on economic players involved in Burley and Oriental tobacco (growers, manufacturers of American blend FMC) compared to Virginia as more additives, including sugar, are used in the manufacturing of FMC based on these tobacco types. Some additional impacts on flavouring suppliers can be expected compared to option 2, although they normally supply multiple sectors.

On balance, the option is not expected to have any additional direct impacts on **Governments**. Compared to option 2, the costs of option 3 associated with ingredients regulation are on the one hand lower (less administrative burden), but on the other hand result in significantly reduced choice for **consumers** although they would benefit from a more comprehensive protection. In terms of **international impacts**, option 3 is well in line with the FCTC guidelines and similar to the Canadian and Brazilian approaches which have been discussed in the framework of the WTO. It could have some impact on EU trade, considering that it would affect primarily Burley and Oriental producers (mainly in the EU), while Virginia producers (including non-EU) would be less affected.

**Social impacts**

The indirect impact on **employment** linked to the drop in consumption is described in section 5.7 and Annex 5. In terms of distributional effects, option 3 is expected to have a greater impact on employment related to Burley and Oriental tobacco (American blend FMC) as explained above. In this context, it should be recalled that more than 80% (69,000) of the EU farmers are involved in these tobacco types. These are often small family businesses concentrated in specific regions, including rural areas in Bulgaria, Poland, Greece, Spain and Italy. In terms of **equality**, option 3 would provide similar (or marginally improved) protection of young people compared to option 2 as it would remove additional products from the market that could be attractive for young people.

**Health impacts**

Option 3 would further reinforce the effect on **awareness** and prevalence outlined under option 2, although only to a limited extent as most of the particularly appealing products are already banned under option 2. The indirect impacts are further developed in section 5.7 and Annex 5.

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\(^{372}\) This is certainly true if one Member State opts for a full ban on additives (Canadian/Brazilian model).
1.1.23. The views of stakeholders

All stakeholders have expressed support for mandatory reporting on ingredients in a common format. The tobacco industry has referred to commercially sensitive data and the importance of keeping this data confidential. In addition, FMC manufacturers have indicated that they agree to participate to the financing of a reporting system with regard to running costs.

In relation to ingredients regulation, the tobacco industry as well as the flavouring industry have emphasised that there is no scientific basis to regulate attractiveness arguing it would be a "subjective concept". Growers have expressed particular concern that a ban on all additives would discriminate against certain types of tobacco (Burley and Oriental) and have very negative impact on employment. Cigar manufacturers have argued against further requirements on ingredients, in particular stressing that cigars are mainly used by adult users and not by young people taking up smoking. The four big FMC manufacturers indicated, however, that prohibiting certain characterising fruit and candy-like flavours for products such as pina colada and strawberry can be further explored. Growers indicated that a ban on dominant flavour could be acceptable. Health NGOs have expressed support for an approach addressing the attractiveness of tobacco products. A majority of the Member States that responded to the public consultation supported some kind of ingredients regulation (Annex 1).

1.1.24. Comparing the options and preferred option

In terms of effectiveness, prohibiting products with characterising flavours (policy option 2) or (even more so) prohibiting all additives not essential for manufacturing (policy option 3) would remove national disparities and unify national rules (policy objectives A3 and B1). Option 1 would partly contribute to the achievement of these objectives. The mandatory and electronic reporting system under options 2 and 3 would also be well in line with objective B2 as it would facilitate enforcement and verification by Governments as regards reporting obligations. All options 1-3 would contribute to policy consideration C1 as they would regulate the most hazardous products on the market. Options 2 and 3 would fulfil consideration C2 as they would remove products which are particularly attractive, including to young people.

Prohibiting products with characterising flavours (policy option 2) appears to be a manageable and efficient measure to regulate tobacco products appealing and attractive to young people, therefore taking into account the concept of “attractiveness”. The measure would affect a limited number of products, while prohibiting all additives not essential for manufacturing (policy option 3) would affect most of the products on the market and thus have more significant economic impact. A ban of all products with characterising flavours, rather than just a limited number of identified flavours would ensure coherence and equal treatment and avoid unjustified differences of treatment (e.g. menthol vs clove).

Preferred policy option 2: Mandatory reporting in harmonised format. Prohibit tobacco products with characterising flavours and products with increased toxicity or addictiveness.

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373 Based on the existing system, EMTOC.
374 The European Association for Cigarette Manufacturers (CECCM) considers the introduction of fees for operating costs acceptable in its contribution to the public consultation on a possible revision of the Tobacco Products Directive.
A harmonised reporting format and mandatory reporting will create a level playing field and facilitate collection, analysing and monitoring of data. It will also reduce the administrative burden of the industry, Member States and the Commission and provide a more robust system to handle sensitive data. All stakeholders support this approach and the European Association for Cigarette Manufacturers (CECCM) considers the introduction of fees for operating costs acceptable (see above).

The preferred option also addresses the heterogeneous development in Member States in relation to ingredients regulation and takes into account international developments (FCTC). It allows industry to adapt the production lines in one go whilst allowing industry some margin to differentiate products. It focuses on products particularly attractive to young people and is estimated to reduce smoking initiation. It addresses recent market developments, including the new technology of inserting additives (e.g. menthol) in the filters of the cigarettes, and allows for further guidance and developments through implementing/delegated acts. A general ban on characterising flavours (instead of regulating individual additives) makes it more difficult to circumvent the ban by developing alternative chemical combinations with the similar properties (taste/aroma). The option provides a balanced approach, taking into consideration the concern of some stakeholders, in particular regarding demand for certain tobacco types (Burley and Oriental) and EU growers have indicated that they could accept the suggested approach.376

The preferred option would contribute to the achievement of the policy objective, i.e. improve the functioning of the internal market through further alignment of Member States’ regulations. In terms of health, it would in particular target smoking initiation among young people. A stricter approach (option 3) would go beyond the focus on young people and affect a substantial number of products which could be seen as less proportionate to the expected additional impact on smoking initiation. A less stringent measure (option 1) would not improve the functioning of the internal market. Also, when balancing the benefits with the costs for the economic operators, option 2 appears to be a proportionate solution given the number of products affected compared to option 3.

Only an initiative at EU level is capable of removing the current and expected diversity in terms of regulation and provide a standardised format for reporting of additives.

5.5. CROSS-BORDER DISTANCE SALES OF TOBACCO

1.1.25. PO1: Notification and age verification system

Retailers of tobacco products intending to engage in cross-border distance sales shall notify their cross-border activities to the Member States where the company has its seat and where it intends to sell. Member States may require the retailer to appoint a natural person, who ensures compliance with the TPD of products delivered to customers in Member States concerned. Mandatory age verification mechanism is foreseen.

Economic impacts

Option 1 would provide more equal market conditions for internet retailers operating on the EU market. The requirement to notify internet retail under option 1 is expected to ensure/improve compliance with the law (including TPD) and is expected to reduce the amount of illegal sales of tobacco products in favour of legal business (traditional retailers

and legal internet retailers) operating in compliance with the TPD (e.g. labelling and ingredients regulation). This would maximise the effect of TPD and have positive impact on the internal market.

The notification requirement and the age control mechanism are expected to impose some compliance costs on tobacco internet retailers. The cost of notification is expected to be minor as the actual requirement will be limited to sending an e-mail/letter to the competent authorities of the host Member State as well as Member States of destination. This must be seen in the context of the specificities of tobacco products responsible for almost 700,000 deaths in the EU each year. The designation of a natural person to ensure compliance with the TPD is optional for Member States and it is expected that the retailer can make use of resources already required by other legal frameworks (tax legislation). The age control verification is also expected to result in minor costs. Option 1 is expected to result in a reduced number of internet retailers as many of the current retailers would close their business rather than notify their (illegal) activity. This reduction would partially reduce consumption, partially lead to a return of consumption to the legal supply chain. Overall, the impact on economic stakeholders is expected to be positive following the estimated shift from illegal to legal sale, despite the somehow reduced consumption.

The shift towards legal sale would also have the unintended but welcome side effect of increased tax revenues for Governments and would facilitate criminal investigations, as internet sales by non-notified businesses would be automatically illegal. On the other hand, option 1 would imply additional, but limited costs for national authorities in terms of setting up the notification system. It is expected that 0.5 full time equivalent (FTE) would be more than adequate whilst law enforcement would be simplified compensating largely for the additional costs. Consumers would continue to have access to several sales channels, including cross-border distance sale and have access to products not available on their domestic markets. They could verify before purchase whether the internet seller is a legal entity competent to engage in such activity. The protection against illicit supply and protection of young people would be improved and more consumers would obtain products complying with the TPD, including with warnings in their own language. Consumer prices are expected to increase as a result of the shift from illegal to legal supply. Option 1 provides a step towards, but is not fully in line with the FCTC guidelines recommending a full ban.

**Social impacts**

As most of the consumers currently purchasing from illegal internet sources are expected to divert to other legal distribution channels, option 1 is likely to have a positive overall impact on employment in the legal supply chain. Option 1 would lead to improved protection of young people as internet retailers would be obliged to control the age of the purchaser.

**Health impacts**

Option 1 is expected to further reduce availability and access to tobacco, in particular for under-aged people. The option would improve awareness as the safeguards of the TPD would be better respected and would contribute, to some extent, to a drop in consumption/prevalence, particularly in the illicit segment of the market. Most of the consumers currently using the internet as a source for tobacco products are expected to turn to alternative, legal sources, but some are also expected not to start, quit or reduce their consumption. It would also limit young people's access to promotional aspects often visible

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377 Some retailers are also expected to continue illegal, non-notified activities.
on the internet and could discourage uptake of tobacco use, in particular among young people. Overall this is beneficial for public health.

1.1.26. **PO2: Prohibit cross border distance sales of tobacco products**

| Cross-border distance sales of tobacco products are prohibited in the EU. |

**Economic impacts**

A ban on cross-border distance sales of tobacco would remove current divergences on the internal market and ensure that certain provisions concerning protection of the internal market (e.g. labelling and ingredients rules set out in the TPD) are fully respected/not circumvented.\(^{378}\) The measure would also facilitate enforcement.\(^{379}\) The current disadvantage of traditional retailers having to compete with lower prices applied by (mainly) illegal internet retailers would be removed. This would have the positive impact on the **internal market.** However, option 2 would also remove legal cross-border distance sale.

Option 2 could imply a one-off compliance cost of lost stocks and of equipment used for the tobacco **internet retail** activity, although these costs would be very limited considering the transposition period. The products can also still be sold domestically and there is no legitimate expectation for illicit activities to continue. Option 2 would interfere with the freedom to conduct business which would need to be balanced against the principle of health protection.\(^{380}\) The shift towards sales in the legal supply chain is also expected, as an unintended side effect, to improve Governments’ ability to collect taxes resulting in increased tax revenues. Indirect impacts directly linked to consumption are further described in Section 5.7 and Annex 5. Option 2 would limit the number of sales channels for consumers, but provide a higher level of protection as it would reduce the availability of illicit tobacco products not necessarily complying with the TPD. The measure is fully in line with the **FCTC** guidelines, but needs to be justified under WTO rules.

**Social impacts**

Option 2 would further reinforce the positive impact on legal **employment** in tobacco outlined under option 1. It would also remove the current negative impact on **young people** and apply in an equal manner to all EU citizens.

**Health impacts**

Many of the current internet sites include promotional aspects and the products are unlikely to comply with required labelling or content rules. This has a negative impact on availability and **awareness.** Option 2 is expected to further reduce access, both to under-aged people who might be tempted to purchase on the internet due to less strict age controls and cheaper prices and adults who wish to avoid paying the applicable taxes. This would lead to a drop in **consumption** exceeding the one described in option 1.

1.1.27. **The views of stakeholders**

The FMC manufacturers expressed support for distance selling of tobacco products to adults provided that such sale is regulated and proper tax payments are ensured (see Annex 1). Retailers have indicated that internet sale of tobacco is associated with problems in relation to

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\(^{378}\) Compare the reasoning in Case C-491/01, The Queen v Secretary of State for Health, ex parte British American Tobacco (Investments) Ltd and Imperial Tobacco Ltd, ECR [2002], p I-11453, para 81-91.

\(^{379}\) Whilst it is true that distance purchases can also take place from countries outside the EU, remaining customs control offer better chances of detection.

\(^{380}\) Article 16 v. Article 35 of the Charter of Fundamental Rights of the European Union.
under age control and collection of taxes. They argue that stricter measures, if any, should be introduced at national levels. Health NGOs have asked for a ban of internet sales of tobacco as a logic extension of the advertising ban.

1.1.28. Comparing the options and preferred option

In terms of effectiveness, both a notification system combined with age verification (option 1) and prohibiting cross-border distance sale (option 2) would facilitate the creation of a level playing field (policy objective A2). Both options would also facilitate market surveillance (policy objective B2). Moreover, both options (and even more so, option 2) would contribute to reduced access for young people (policy consideration C5) and minimum level of health protection (policy consideration C6).

In terms of costs, the establishment of a notification system under option 1 would imply additional, however limited, costs for national authorities whilst facilitating law enforcement. Prohibiting cross-border internet sales (policy option 2) would fully remove one sales channel and thereby limit market opportunities. It would deprive consumers of the possibility to buy tobacco products not available in the Member States of location. Both options are also expected to lead to reduced illicit trade and thereby benefit economic players involved in the legal supply chain.

Preferred policy option 1: Notification and age verification system

The preferred option will facilitate legal activity without removing any sales channel, while allowing consumers legitimate access to tobacco products not available on their domestic market. The option reinforces the effect of the TPD by preventing purchasing of products not complying with the Directive, including health warnings in the right language and ingredients regulation. An unintended side-effect is that the preferred option also will reduce the availability of cheaper products not respecting national price policies, address underage purchasing and allows Governments to better collect applicable taxes.

The preferred option complies with the principle of proportionality. It contributes to the achievement of the internal market objective by ensuring TPD safeguards and facilitating trade in products complying with TPD. The additional burden for retailers involved in cross-border distance sale of tobacco products is considered justified taking into consideration the specific characteristics of tobacco products (responsible for almost 700,000 death in the EU each year) not comparable with any other product on the internal market. The preferred option will also contribute to a drop in consumption/prevalence. Given that option 1 will contribute to better compliance with TPD, a full prohibition of cross-border distance sale (option 2) would go beyond what is necessary to obtain the objective. There is no less strict measure available to attain the objective. The costs associated with the preferred option both for economic operators and Governments are considered limited (if any) compared to the benefits following from reduced consumption, reduced illicit trade in tobacco and facilitated enforcement.

Cross-border distance sales can only be addressed effectively at EU level as the sales, by definition take place between Member States. Therefore, an EU action in these areas is of added value.
5.6. TRACEABILITY AND SECURITY FEATURES

1.1.29. PO1: EU tracking and tracing system

An EU tracking and tracing system at packet level for tobacco products throughout the supply chain (excluding retail) is introduced. Tobacco manufacturers shall conclude contracts with independent third parties that provide data storage capacities for such system ensuring full transparency and accessibility by Member States at all times. Tobacco products other than FMC and RYO are granted a transitional period of five years. Delegated/implementing power to adopt technical specification to ensure compatibility between the systems used.

Economic impacts

Ensuring full traceability and a strong control of the supply chain by a tracking and tracing system will contribute to the functioning of the internal market by reducing the volume of illicit supply, which is undermining the safeguards of the TPD. The system would create a level playing field for all operators on the internal market and remove the current situation where only the four biggest tobacco companies are bound by obligations as regards tracking and tracing via the Agreements with the EU and the Member States (see section 2.2.5). Currently, there is no harmonisation of national legislations on tracking and tracing in the EU although, in practice, there is increasing convergence in terms of technical standards used by tobacco manufacturers.381 Option 1 would thus ensure a harmonised approach in Member States.

Tracking and tracing systems are generally seen as an effective means to address the risk of diversion of genuine products into the illicit market and such systems have already been developed and used in recent years. Based on industry data an effective tracking and tracing system reduces illicit contraband by 30% in five years. The main part of these additional revenues will go to the legal supply chain (FMC/RYO manufacturers and distribution chain), but some smokers are also expected to stop smoking/not to start smoking.

Due to the existing Agreements between the four largest tobacco manufacturers and the EU and participating Member States (see section 2.2.5), the largest tobacco manufacturers are already implementing some of the requirements foreseen under option 1 and additional costs (e.g. associated with the outsourcing of the data storage are considered proportionate when compared with the existing contractual obligations (tracking and tracing at packet level). It should also be noted, that the Agreements will start expiring as of 2016 (unless they are extended). The proposed measure will be more burdensome for smaller operators who will possibly have to build up a tracking and tracing system from scratch. However, there are possibilities to benefit from existing experience382 and in any case it will be easier if such a system is introduced at EU level rather than in 27 Member States individually. Operators involved in tobacco products other than FMC and RYO products (e.g. cigars and pipe tobacco) will benefit from longer transitional periods.

It is not easy to fully estimate the costs associated with the introduction of a tracking and tracing system. However, the following cost estimates, which were provided in the context of

381 Divergence might still exist regarding the package level and the level of trade.
382 For example, PMI declared its readiness to grant royalty free licenses to third parties that want to use PMI's tracking and tracing software. Presupposing that the PMI system is compatible with the international obligations this would reduce the costs for introducing such a system substantially.
the international negotiations for an Illicit Trade Protocol (based on information from PMI) are a good indication:

For the currently used carton level tracking and tracing, the industry's indications are that the actual one-off costs include machines for printing the label for master cases (5,000–20,000 EUR per machine) and for cartons (10,000–35,000 EUR) as well as for carton coding system (10,000-35,000 EUR) and carton tracking system (30,000-50,000 EUR). To this should be added hardware (10,000-30,000 EUR), IT infrastructure for the retention of tracking and tracing data (10,000-150,000 EUR) and hosting and maintenance of data (50,000-150,000 EUR). All costs for the equipment etc. would be covered by the manufacturers (including equipment used in the distribution chain). The only additional costs at distributor level are handling costs, but it is believed that the system would be highly automated.

Taking into account the significant cross-border trade, only an EU system can be effectively enforced. On the positive side, it also needs to be noted that the expected decrease of illicit trade would have the effect of increasing the sales in the legal supply chain. The total illicit market at retail level (excluding taxes) is estimated to amount to 3 billion EUR, of which 30% are attributed to contraband products. Based on industry data an effective tracking and tracing system reduces illicit contraband by 30% in five years. Part of these additional revenues will go to the FMC/RYO manufacturers (other parts to the distribution chain, and some smokers are also expected to stop smoking/not to start smoking).

For Governments, option 1 would have positive impacts in that it strengthens the legal supply chain and ensures that consumers benefit from the safeguards of the TPD. A tracking and tracing system which gives the authorities access to the data storage of the independent third party, will help the authorities (including the Commission) to monitor systematically the movement of tobacco products from the place of their manufacture, through the distribution chain to the intended market of retail sale ("tracking"). It will also enable the authorities, at the time of an audit or seizure of a product, to recreate the route taken by the product from the place of its manufacture, through the distribution chain to the point at which the product was diverted into illegal trade channels ("tracing"). On the other hand, some administrative costs are expected from the monitoring of the system. The measure would thus facilitate market surveillance. An unintended but welcome side effect would be that the measure would increase Governments' tax revenues by decreasing tax evasion. For consumers, option 1 would mean a higher level of protection. It would also mean that the availability of cheap illicit FMC and RYO that do not comply with the TPD is reduced. Option 1 improves compliance with the FCTC.

Social impacts
The shift from illegal to legal trade would result in increased employment opportunities in the legal tobacco sector despite the predicted overall prevalence drop (which would essentially come from the illicit segment). There would also be increased employment opportunities in the setting up and operating tracking and tracing systems (including data storage). In terms of equality, the measure would contribute to removing illicit tobacco products' potential to target young and deprived people.  

Health impacts

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383 FCTC/COP/4/INF.DOC./1 of 15 September 2010.
The proposed measure would ensure that the intended increase in awareness is not circumvented by illegal **products which do not comply with the relevant requirements** (such as labelling and ingredients regulation). Part of the demand previously met by illicit products would shift to the legal supply chain, whereas another part is expected not to be substituted, i.e. would result in reduced consumption, which would in turn lead to reduced health risks and increased well-being in the long term. For the indirect impacts, see section 5.7 and Annex 5 which apply, however, primarily to the legal supply chain (as the illegal part of the supply chain has no legitimate expectations).

1.1.30. **PO2: Tracking and tracing system, complemented by security features**

| Option 1 plus: security features against counterfeiting and against illicit/cheap whites (glossary) on all tobacco products (e.g. holograms). Tobacco products other than FMC and RYO are granted a transitional period for five years. Delegated/implementing power to adopt technical specifications for the security features. |

The subsequent assessment only covers the additional elements beyond option 1.

**Economic impacts**

Adding special security features to limit sales of counterfeits and illicit whites (glossary) at EU level would help limiting sales of these products and further facilitate the functioning of the **internal market** of legal tobacco products. It avoids disparities and differences as regards security features and prevents further heterogeneous development in coming years following national measures to address illicit trade. It would also improve compliance of tobacco products with the safeguards of the TPD. Adding a security feature would imply additional costs for the **industry** to comply with the requirements. These costs depend largely on the technology chosen and might reach 0.005 EUR per package, for example for a hologram.\(^{385}\) According to Euromonitor, 608 billion sticks of FMC were sold in the EU in 2010. This corresponds to approximately 30.4 billion packages of FMC (assuming a package size of 20 sticks per package). The costs for applying holograms on all cigarette packages would thus amount to **approximately 150 mEUR**\(^{386}\). It is assumed that these costs would be passed on to subsequent distribution level and ultimately to the consumer.

On the other hand, the industry would benefit from reduced illicit trade. In addition to the reduction of illicit trade (contraband) estimated under option 1, it is assumed that counterfeit and illicit white products would be reduced by 10% under option 2. Again, the larger part of these additional revenues is expected to go to the legal supply chain (FMC/RYO manufacturers, distribution chain), but some smokers are also expected to stop smoking/not to start smoking. In addition, the economic operators would benefit from a harmonised approach as they do not have to adapt to different regimes in different Member States. The calculations of the indirect impacts linked to a change of consumption patterns are set out in Section 5.7 and Annex 5.

**Governments** would benefit from a common system for security features. Whilst only seen as an unintended side effect for the purpose of the TPD revision, option 2 would also contribute to the collection of taxes. As the costs for the security features will be borne by industry, no

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\(^{385}\) This is based on costs for a self-adhesive frangible (tamper evident) polyester hologram closure seal to be machine-applied at the packing or filling station, which is considered one of the more sophisticated security measures. (indication by the International Hologram Association)

\(^{386}\) The calculation presupposes that the actual application of the security features on the package is included in the price.
additional costs beyond option 1 are foreseen. Increased tax income and facilitated market surveillance as well as effective implementation of the FCTC are further benefits for Governments. Moreover the security feature would assist consumers in determining whether the product is authentic and ensure a higher level of protection against products not complying with the TPD.

**Social impacts**

The positive impacts on employment outlined in option 1 would apply and be reinforced under this option. In addition, some employment would also be created in the production and application of security features. The option would also further strengthen equality due to the reduction of counterfeit and illicit white products (or at least by slowing down their growth). Availability of cheap (illegal) products would be reduced, which are through their lower price particularly appealing to young or deprived persons.

**Health impacts**

Security features will increase the awareness of the problems around illicit tobacco products and thus indirectly lead to increased awareness of the risks associated with tobacco consumption in general. Option 2 would further reinforce the positive health impacts outlined in option 1 and is expected to contribute to a significant reduction in consumption.

1.1.31. **The views of stakeholders**

Policy area 5 was introduced in reaction to concerns expressed by the industry (in particular FMC and RYO manufacturers, upstream suppliers and downstream distributors) in the context of the public and targeted stakeholder consultations. They argued that illicit trade already accounts for a large part of the market and is expected to increase. They pointed in particular to the risks associated with some of the policy options on plain packaging and display ban at PoS (which was subsequently disregarded in this impact assessment, see section 4.1) (*nota bene*: the evidence submitted by the economic stakeholders to support this argument is not considered compelling/convincing). No concrete proposals on how to effectively address illicit trade were made, but tracking and tracing is believed to provide the appropriate response and is already standard practice for the big four tobacco manufacturers. Providers of security systems (e.g. holograms) said that such a measure would reduce counterfeit to a certain degree (up to 30%) but would not eliminate it all together (*nota bene*: for the purpose of this impact assessment reduction of 10% was assumed). Member States are generally supportive, in particular in view of their tax revenues. Health NGOs contested that other planned TPD measures, e.g. on packaging would increase illicit trade, but nevertheless expressed support for additional measures to fight illicit trade.

5.6.1.1. **Comparing the options and preferred option**

In terms of effectiveness, an EU tracking and tracing system complemented by security features (policy option 2) would contribute the most to facilitating a level playing field (policy objective A2), although a tracking and tracing system on its own (policy option 1) would also go into this direction. Both options would contribute to better market surveillance (policy objective B2). A tracking and tracing system complemented by security features (policy option 2) would assist consumers in verifying the authenticity and ensure a minimum level of health protection (policy considerations C3 and C6). Both options would contribute to reducing accessibility to young people (policy consideration C5).

An EU tracking and tracing system without security features (policy option 1) would provide a less costly solution to set up, in particular as a large part of the market already has a tracking and tracing system in place, at least at carton level. However, it would not specifically address
the issue of counterfeiting which is important in terms of ensuring a high level of health protection in line with the TPD. The costs associated with these two measures are expected to be outweighed by the significant benefits in terms of reduced illicit trade, resulting in increased sales for economic stakeholders (for more details, see Annex 5).

| Preferred policy option 2: Tracking and tracing system, complemented by security features |

The preferred option maximises the effect of the TPD and ensures that consumers throughout the EU benefit from a minimum level of protection when purchasing tobacco products (e.g. health warnings and ingredients regulation). The option creates a level playing field between different operators (currently only the biggest four tobacco manufacturers are bound to develop and use tracking and tracing systems) and would be beneficial for them (unless they are themselves involved in illicit trade). The option ensures compliance with Article 15 FCTC and ensures a common EU approach. It also facilitates market surveillance and empowers consumers in verifying the authenticity of tobacco products. The option addresses concerns of some stakeholders claiming that revision of the TPD would result in increased illicit trade (however without substantiation). The option also foresees a longer transitional period for manufacturers of tobacco products other than FMC and RYO (e.g. STP, cigars and pipe tobacco). This will allow small business to adapt and learn from bigger companies.

There is no less restrictive measure available, because only if the market is effectively protected against illegal supply can the measures foreseen in the TPD achieve their objective. The two elements included in the preferred option are mutually reinforcing and address two separate aspects of illicit trade. A tracking and tracing system addresses contraband products non-complying with the TPD, while the security features deal with counterfeit and illicit whites.

The costs associated with these two measures are expected to be outweighed by the benefits in terms of reduced illicit trade which partially benefit the legal supply chain and leads to reduced consumption.

By definition, only an initiative at EU level is capable of ensuring a harmonised approach. Unilateral actions by Member States can contribute to the protection of health, but result in many legal and factual divergences and reduced effectiveness taking into account the significant cross-border nature of the market. The preferred option would provide an EU added value and an EU wide measure also facilitates enforcement in Member States. The preferred option is a necessary addition to all other measures in this proposed TPD, as only strong measures can provide for a protection of the legal market with all the necessary safeguards for consumers, manufacturers and retailers.

In designing the legislation, due attention will be paid to the freedom to conduct business and to the protection of personal data in line with existing legal requirements.

5.7. INDIRECT EFFECTS / HEALTH IMPACTS

As explained the various policy options are expected to impact the economic stakeholders and Governments not only in a direct manner (e.g. costs/benefits associated with the implementation of the measure), but also in an indirect manner. Over time the proposed measures are expected to impact on peoples' awareness on the risks associated with tobacco products, which in turn will lead to a change in behaviour. Less young people will start smoking and some adults will successfully quit smoking. This is expected to lead to a reduction of smoking consumption/prevalence.
When comparing with international experiences, it is assumed for the purpose of this impact assessment that the combination of the preferred policy options will lead to a **reduction of consumption of around 2%** (1.7-2.6% see figure 14 below) within a five year period after transposition beyond the baseline for FMCs and RYO. This corresponds to a reduction of 2.4 million smokers in the EU. It has to be stressed that this figure is a best effort estimation.

The assumption of 2% is mainly based on experiences and estimations from other jurisdictions. All policy areas are expected to make a contribution to the overall consumption drop, albeit not to the same degree. The main contributions are expected from the policy areas on packaging and ingredients which are mutually reinforcing.

Several independent studies have assessed and attempted to quantify the impacts of **packaging and labelling measures**. Practically all reached the conclusion that such measures impact on the awareness of consumers, which over time changes also smoking behaviour but there was some divergence as to the exact level. The prevalence of adult smoking in Canada has declined approximately 6% since the implementation of large pictorial warnings in 2001, which is at least partially attributable to the picture warnings. Another study prepared for Health Canada assessing a set of policy measures on labelling similar to the ones proposed in option 1 under packaging and labelling estimated a rather small reduction of 0.3 percent to 0.8 percent in the number of smokers within ten years. Assuming uniform quit rates across smokers with different consumption levels this corresponds to a comparable drop in consumption of about 0.3-0.8% resulting from those quitting. It should be emphasized that at the time of the Canadian assessment picture-based warnings were already in use which could limit the additional impact to be expected. A cost-benefit analysis prepared for the Australian Government estimated that introducing pictorial warnings covering 50% of the front and back of the packets would result in a 1.3% decline in smoking prevalence rate per annum (12.3% decrease in 10 years) and a 3% decrease in tobacco consumption per capita per annum (26.3% decrease in 10 years). The UK Department of Health’s impact assessment (2007) estimated that the introduction of pictorial warnings would result in a 0.5% decrease in smoking prevalence in the long term compared to 0.05 percent decrease in consumption if the status quo (text warnings only) were maintained. Finally, a more recent US impact assessment estimated the reduction in smoking prevalence as a result of introducing nine pictorial warnings, occupying 50% of both display areas, including a mandatory reference to a toll-free quitline. The reduction in the US smoking population in 2013 was estimated at 213,000 persons (corresponding to a prevalence reduction of 0.45%), with small subsequent effects.

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389 As a result of the measures proposed, an additional reduction in consumption levels can also be expected for those who keep smoking.


In its opinion from 2010, SCENIHR concluded that the use of fruit and candy flavours seems to favour smoking initiation in young people and that some additives decrease the harshness and increase the smoothness of the smoke.\textsuperscript{393} The US FDA Tobacco Products Scientific Advisory Committee confirmed, on the basis of the extensive review of all available information, that the evidence was sufficient to conclude that it is more likely than not that the availability of menthol FMC increases the likelihood of experimentation and regular smoking beyond the anticipated prevalence if such FMC were not available.\textsuperscript{394}

The contribution to the reduced smoking prevalence from policy area “\textit{STP and extension of the product scope}” is primarily expected to result from the possibility that e-cigarettes can develop a potential as a smoking cessation aid under the preferred option.

Also the measures in the policy areas dealing with \textit{cross-border distance sales}\textsuperscript{395} and \textit{traceability & security features} are expected to contribute to a drop in consumption, in particular in the illicit segment of the market. Part of this demand will return to the legal supply chain, which is however more expensive and therefore it is expected to encourage some consumers not to start smoking/stop smoking or smoke less, in particular in parts of society with lower revenues such as young people. Also, consumers are better informed by the health risks of tobacco products fully compatible with the TPD.

Figure 14 provides a tentative break-down of the contributions of individual policy areas. More information is provided in Annex 5. While 2\% is estimated reasonable, it needs to be underlined that even a lower drop in consumption remains beneficial from a macro-economic and Governmental/societal perspective (see section 6.2.4 and Annex 5). It needs also to be stressed that conclusive empirical data is lacking for some of the measures, including NCP (where no electronic cigarettes have been authorised, at this stage, under the medicinal products’ legislation).

\begin{table}[h]
\centering
\begin{tabular}{|c|c|}
\hline
\textbf{Policy area} & \textbf{Foreseen contribution to the decrease in \%} \\
\hline
Scope & \\
(STP) & 0.2-0.3 \\
NCP & \\
(Herbal) & \\
Packaging & Labelling & 1-1.5 \\
Ingredients & 0.5-0.8 \\
\hline
\end{tabular}
\caption{Tentative contributions of individual policy areas to the projected decrease of cigarette/RYO consumption}
\end{table}

\textsuperscript{393} SCENIHR 2010.


\textsuperscript{395} In an experimental Californian study from 2004 96.7\% of participating minors were successful in finding and ordering tobacco on the internet. Jensen JA, Hickman NJ 3rd, Landrine H, Klonoff EA. Availability of tobacco to youth via the Internet. JAMA 2004; 291(15):1837.
<table>
<thead>
<tr>
<th>TOTAL</th>
<th>1.7-2.6%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cross-border distance sales + Illicit trade</td>
<td>Additional decrease of consumption, however not in the legal supply chain. (Decreases in illicit consumption could mitigate the decrease in the legal chain).</td>
</tr>
</tbody>
</table>

In addition to the predicted drop in smoking prevalence, the preferred options are also likely to result in reduced uptake of **STP and herbal products for smoking**. In particular, it is expected that STP use will remain limited to specific population groups already using these products and that recruitment of new users will become more difficult. However, it is difficult to quantify this effect, also in the light of increased use of smoke-free environments.

### 1.1.32. Economic Stakeholders

For the **tobacco industry** (the manufacturers of FMC and RYO) a reduction of consumption in the range of 2% within five years would mean that their revenues would decrease by 376 mEUR per annum. This figure is a “**worst case scenario**” for the industry, as it does not take into account the effects of the policy options developed in policy areas 4 and 5, which have the effect that certain sales in the illicit supply channel return to the legal supply chain. In terms of employment, the reduction of consumption is also expected to reduce the work force of the tobacco industry. Based on the input output model explained in Annex 5 the reduction is expected to amount to 1500 employees. On the other hand it should be noted that money not spent on tobacco products will be spent on other goods and services. In this respect the “input-output model” used for the purpose of the impact assessment indicates **net gains in employment of around 2,200**. Industries that are most likely to benefit are food/beverage, textiles manufacture and the service sector.

Annex 5 also presents what a reduction of tobacco consumption would mean for the **upstream suppliers** (e.g. growers, ingredients suppliers, paper industry) and **downstream distributors** (wholesale, retail). Overall the impacts associated with the reduction in consumption do not seem to be of a significant nature, as the economic stakeholders typically sell other products (e.g. package industry makes only around 10% of its turnover with the tobacco industry) or generate a significant part of their turnover with export sales outside the EU which would not be affected. Even the impact on specialised retailers does not seem to be significant as – according to the EU association of the retailers (CEDT) – these specialised shops nowadays generate about half of their turnover with non-tobacco products. The impact on the revenues for tobacco growers would mean an annual reduction in income of 158 EUR per farmer, which would however, not affect all farmers alike. Rather it is expected in line with previous trends (discontinuation of direct subsidies) that certain farmers will discontinue their activities irrespective of the TPD revision.

### 1.1.33. Governments/society

The main socioeconomic impact for Governments/society – associated with a reduction of tobacco consumption - is the improvement of public health. Non-smokers live longer and benefit from more **healthy life years**. It has been estimated that smokers who die as a result of their tobacco consumption pass away 14 years earlier than people who never-smoked (see Annex 5, 5.2.3.1.1). A reduction of consumption of 2% corresponds to 2,4 million smokers

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stopping the behaviour and 16.8 million life-years (not quality adjusted) gained. The gain of (healthy) lifeyears is a value in its own right and of greatest importance for the persons concerned as well as their families. On the basis of the impact guidelines it is possible to monetise the gain in public health. In this light, a reduction of the tobacco consumption by 2 % would correspond to a benefit for society of 10.3 bEUR annually.

In addition, a reduction of tobacco consumption will reduce health care expenditure by 506 mEUR annually. Also society would benefit from a reduction of productivity losses (less absenteeism and premature retirements) in the region of 165 mEUR annually (see Annex 5). As explained these are conservative estimates as other studies report about significantly larger cost savings for health care and gains in productivity.

Another impact for Governments would be that they are confronted with a risk of reduced tax revenues by 1.6 bEUR annually because of reduced consumption. It is important to note from a macro-economic perspective the reduction of taxes is not a cost, but only a question of allocations within society (State/companies). It also needs to be taken into account that the reduction of taxes is an unlikely "worst case scenario". The figures do not yet take into consideration the positive effects of the policy options developed in policy areas 4 and 5 which would increase Governments' tax revenues. Additionally, a reduction of tobacco consumption does not necessarily lead to lower tax incomes as tax rates can be increased (like in the past). As a matter of fact despite decrease of tobacco consumption since 2000 tax revenues for Member States increased very significantly in the same period. In addition, money not spent on tobacco would be spent on other sectors also benefitting the Governments in terms of taxes.

In summary it can thus be concluded that the impact of reduced tobacco consumption would lead to overall benefits of 9.4 bEUR per annum. The calculation is based on present values and thus in principle expressed in current prices.

Social discounting allows comparison of benefits and costs that occur at different times based on the rate at which society is willing to make such trade-offs. This is also relevant in the case of tobacco control as some of the expected benefits will only develop over time whilst certain impacts (e.g. on tax revenues) would materialise earlier. Different scenarios for social discounting have been developed in Annex 5 (5.2.3.4). Under the most likely scenario (i.e. when decrease in tax revenues and health care/absenteeism savings materialise in the period of 5 years, while on average the benefits from reduced premature mortality accrue only in 25 years), the annual net benefit of a reduction in tobacco consumption by 2 % would be 4 bEUR.

Figure 15 provides a breakdown of the overall net costs and benefits (including discounted values).

**Figure 15: Overall net costs and benefits (mEUR)**

<table>
<thead>
<tr>
<th>Different percentage reduction in tobacco consumption</th>
<th>1%</th>
<th>2%</th>
<th>3%</th>
<th>4%</th>
<th>5%</th>
</tr>
</thead>
</table>

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397 Population over 15 is covered by EB - according to Eurostat this is 431 million adult citizens. 28% percent of them, thus 120.8 million are smokers. 2% of those correspond to 2.426 million smokers. Assuming than 7 years are gained per smoker (14 years /2 prematurely dying), this result in 16.8 million lifeyears (not quality adjusted).


399 Social discounting renders benefits and costs that occur in different time periods comparable by expressing their values in present terms based on the rate at which society is willing to make such trade-offs.
Decrease in excise tax revenues\textsuperscript{400} & 794 & 1588 & 2382 & 3176 & 3970 \\
Decrease in health care expenditures & 253 & 506 & 759 & 1012 & 1265 \\
Decrease of productivity loss & 83 & 165 & 248 & 331 & 413 \\
- due to early retirement / deaths & 61 & 122 & 183 & 244 & 305 \\
- due to absenteeism & 22 & 43 & 65 & 87 & 108 \\
Decrease in premature mortality costs & 5,167 & 10,334 & 15,501 & 20,669 & 25,836 \\
Overall net benefit & 4,709 & 9,417 & 14,126 & 18,836 & 23,544 \\
Discounted values & 2,016 & 4,032 & 6,048 & 8,064 & 10,080 \\

The preferred options would have a significant positive impact on public health. For example, the current targeting of young people with specifically developed products like candy-flavoured FMC and attractive packages would be removed, young people's access to tobacco would be further restricted and improved health warnings would improve all citizens’ possibilities to make informed decisions. In addition, due to the foreseen reduction in illicit trade smokers would benefit more from the protective rules set up by TPD (e.g. health warnings and ingredients control).

6. **OVERALL CONCLUSION OF THE IMPACT ASSESSMENT**

6.1. **COMPARING THE OPTIONS AND THE PREFERRED OPTIONS**

Following the conclusions in the previous sections, the table below gives an overview of the policy options as well as justifications for choosing the preferred options (which are marked in grey).

\begin{tabular}{|l|l|l|l|l|l|}
\hline
\textbf{PA / Options} & \textbf{1} & \textbf{2} & \textbf{3} & \textbf{4} & \textbf{Justification} \\
\hline
1a. STP & Lift the ban on oral tobacco and subject all STP to stricter labelling and ingredients regulation & Maintain the ban on oral tobacco, subject all novel tobacco products to notification obligation and all STP to stricter labelling and ingredients regulation & Maintain the ban on oral tobacco, restrict the sale of other STP to areas of traditional use only and subject all STP to stricter labelling and ingredients regulation & Ban all STP with the exception of oral tobacco in Sweden. Subject oral tobacco in Sweden to stricter labelling and ingredients regulation. & -achievement of policy objectives/considerations: B1, C2, C4 and partly A1, A2 and C1. -harmonised labelling and ingredients regulation for all STP. -facilitated level playing field. -proportionate to prevent the introduction / expansion of new addictive, harmful products in the internal market -health concerns with all STP -no evidence that STP leads to smoking cessation, risk of entry gate and dual use -risk for market development (ingredients and smoke-free environments) -Trade-off: impact on SME justified due to health risks associated with product development, marketing and expansion to new user groups. Option 3 and 4 would have more positive impact on health, but option 2 was considered more proportionate after a cost/benefit balance. \\
\hline
\end{tabular}

\textsuperscript{400} Disregarding measures taken against illicit trade and possibility to increase tax levels.
<table>
<thead>
<tr>
<th>1b. NCP</th>
<th>Subject NCP to labelling and ingredients requirements under TPD</th>
<th>Establish a new authorisation scheme for NCP</th>
<th>Subject NCP over a certain nicotine threshold to the medicinal products’ legislation and the remaining NCP to labelling requirements</th>
<th>Subject all NCP to the medicinal products’ legislation</th>
<th>-achievement of policy objectives/considerations: A1, A2, B2, C1, C2, C4 -clear and well established legal framework for risk/benefit analysis facilitating the free circulation of duly authorised products, in conformity with their nature -possibility of mutual recognition within the internal market. -same treatment of NCP and NRT. -harmonised approach, consolidating trend in MS -minimum safety standard -potential in smoking cessation -Trade-off: additional burden for application justified by the setting up of a harmonised safety net with potential to reduce smoking.</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA / Options</td>
<td>1</td>
<td>2</td>
<td></td>
<td>Justification</td>
<td></td>
</tr>
<tr>
<td>1c. Herbal products for smoking</td>
<td>Subject all herbal products for smoking to labelling requirements under TPD</td>
<td>Phase out marketing of herbal products for smoking</td>
<td></td>
<td>-achievement of policy objectives/considerations: A2, A3, C4 and partly B1, C1 -facilitates the free circulation of products -remove current misperception on health Trade-off: Removes misperceptions while minimising compliance costs.</td>
<td></td>
</tr>
<tr>
<td>PA / Options</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td>Justification</td>
<td></td>
</tr>
<tr>
<td>2. Packaging and labelling</td>
<td>Mandatory enlarged picture warnings</td>
<td>Option 1 plus harmonise certain aspects of packets and FMC appearance and prohibit promotional and misleading elements</td>
<td>Option 2 plus full plain packaging</td>
<td>-achievement of policy objectives/considerations A3, B1, C2, C4 -removes disparities on internal market and facilitates free circulation -improves awareness and removes misleading elements -in line with FCTC commitments -proportionate: focus on smoking initiation, products attractive for young people -takes into account economic stakeholders’ concerns (e.g. no plain packaging/interference with trademarks) -allows awaiting international and scientific developments -allows Member States to adopt plain packaging to comply with FCTC guidelines as far as it is compatible with the Treaty -Trade-off: Option 3 more effective from health point of view, but appropriate to wait for real life experience.</td>
<td></td>
</tr>
<tr>
<td>3. Ingredients</td>
<td>Common reporting format on a voluntary basis. Prohibit toxic, addictiveness and attractive additives in tobacco products.</td>
<td>Mandatory reporting in harmonised format. Prohibit products with characterising flavours and products with increased toxicity</td>
<td>Mandatory reporting in harmonised format. Prohibit all additives not essential for manufacturing.</td>
<td>-achievement of policy objectives/considerations: A3, B1, B2, C1, C2 -removes disparities on the internal market and facilitates free circulation -reduces administrative burden (reporting).</td>
<td></td>
</tr>
</tbody>
</table>
In terms of effectiveness, the combined preferred options contribute to the overall objective of the revision of the TPD to guarantee a proper functioning of the internal market while ensuring a high level of public health. As illustrated below, the preferred options are also well in line with the objectives and considerations identified in section 3.

As far as the main objective (internal market) is concerned, the options:

- facilitate the free movement of goods in the internal market by removing existing or expected discrepancies between national legislations and/or,
- adapt the level of harmonisation in the current TPD to a new level warranted by scientific development or international obligations/commitments and/or,
- ensure that the safeguards of the TPD are not undermined by illicit products or by cross-border sales not respecting these safeguards.

As far as health considerations are concerned, the envisaged revision focuses on discouraging young people from taking up tobacco consumption and allowing adult consumers to take informed decisions. The revision also allows smokers to benefit from the protective measures.
set by the TPD. In instances where new products aim at exploiting smoke-free policies, this is also tackled.

The combined preferred options would allow/facilitate Member States/EU to fulfil their commitments/obligations in the context of FCTC. In many policy areas (herbal products for smoking, packaging & labelling, ingredients, and cross-border distance sales) a stricter option than the one identified as preferred option would have been more beneficial for health. However, the overall assessment of all applicable assessment criteria, including impact on economic stakeholders, has led to the conclusion that a less strict measure was more appropriate (see table above).

There are also important synergies between individual options increasing the overall effectiveness of the “preferred options package”. For example, prohibiting tobacco products with characterising flavours is reinforced by the removal of misleading and promotional elements on the labelling and the obligation to notify cross-border distance sale will reinforce the reduction of illicit trade expected from the measure on traceability and security features.

The combined preferred options are cost-efficient in the sense that they are expected to result in overall socio economic benefits for society. For economic stakeholders the preferred options are foreseen to lead to reduced direct (compliance) costs which, however, can be outweighed by lost revenues due to a decrease in consumption. The overall impacts of the preferred options are further described in the next section, 6.2.

The identified options constitute a coherent approach consistent with international commitments and fundamental rights and values as well as the overall aim of the EU to promote well-being of its people (Article 3 TEU). The combined options contribute to the Europe 2020 strategy in the sense that keeping people healthy longer will have positive impact on productivity and competitiveness.

A proper stakeholder consultation has been carried out throughout the impact assessment process (section 1.3) and the comments from various stakeholders have been considered carefully. The next section (section 6.2) describes further the overall impact on key stakeholders.

6.2. OVERALL IMPACTS OF THE PREFERRED OPTIONS

1.1.34. Internal market

As explained, it is expected that the combined preferred options would improve the functioning of the internal market by removing current disparities in national legislation, ensure homogenous development and facilitate the creation of a level playing field among economic actors. In contrast to the baseline scenario, the preferred options would also contribute to a consistent approach in the implementation of international commitments (FCTC).
1.1.35. **Economic stakeholders**

6.2.1.1. **FMC and RYO manufacturers**

The impacts on FMC and RYO manufacturers consists of the direct impact from complying with the preferred options (direct effects) and indirect impacts following an expected decrease in consumption.

The **direct impact on FMC and RYO manufacturers is expected, to be positive compared to the baseline scenario**. The benefits stem in particular from reduced one-off costs associated with familiarisation (all product related areas) and redesign/reformulation (labelling and ingredients) as well as acquisition of new equipment (tracking and tracing). With respect to the variable costs it is assumed that no noteworthy additional costs are expected. To the contrary in certain areas such as traceability and security features, cost savings are expected. In any event additional costs are likely to be passed on to consumers taking into account the significant market power of the tobacco manufacturers. Higher revenues are also expected for FMC/RYO manufacturers as a result of the measures against illicit trade. While a precise quantification is not possible, the table below describes the situation in general terms and refers to figures to the extent possible.

<table>
<thead>
<tr>
<th>Cost category (compliance cost)</th>
<th>Preferred option</th>
<th>Baseline scenario</th>
</tr>
</thead>
</table>
| One-off costs                   | -Economic stakeholders only have to get familiar with one legal system, implemented within a limited time frame.  
- Economic stakeholders have to undertake one redesign/reformulation only adapted to the markets of all Member States.  
- Some economy of scale is/can be achieved when acquiring of new equipment due to harmonisation.  
- Disposal of old stocks. This cost can normally be disregarded taking into account transposition and transitional periods.  
**Packaging and labelling:**  
FMC: 14,500-50,000 EUR per SKU  
RYO: 2,500-9,000 EUR per SKU  
**Ingredients:**  
Product redesign: 1mEUR per brand  
**Tracking and tracing:**  
Indicative costs for carton level tracking and tracing:  
Printing machine master case: 5,000-20,000 EUR per machine  
Printing machine carton: 10,000-35,000 EUR per machine  
Carton coding system: 10,000-35,000 EUR  
Carton tracking system: 30,000-50,000 EUR  
Hardware: 10,000-30,000 EUR  
IT infrastructure: 10,000-150,000 EUR  
Hosting and maintenance: 50,000-150,000 EUR | -Economic stakeholders need to monitor 27 different legal systems and get familiar with all unilateral changes adopted by Member States. This can also include seeking legal advice.  
- Economic stakeholders need to adapt to subsequent unilateral changes adopted by Member States. This also includes testing and legal/marketing/scientific advice. Stakeholders also have to adapt production lines to comply with different national legislations.  
- Economic stakeholders might have to buy new equipment to comply with national measures.  
- Disposal of old stocks. This cost can normally be disregarded provided sufficient transitional periods are granted.  
The cost estimates provided for the preferred option would be higher under the baseline scenario due to multiple and consecutive adaptations to national systems under the baseline scenario. |

401 In the latest Eurobarometer 2012 consumers stated that they are ready to accept price increases to fight against illicit trade.
On the other hand, the **indirect impacts following from an expected decrease in consumption will over time lead to a loss of revenue** for the FMC and RYO manufacturers of 376 mEUR ("worst case scenario" as the benefits from reduced illicit trade are not considered in this figure). This negative impact on FMC and RYO manufacturer might outweigh the benefits from the cost savings referred to above. However, money not spent on tobacco is expected to be spent on other sectors which would then benefit.

### 6.2.1.2. Upstream suppliers

**Growers** would only be affected indirectly by the reduction of smoking and the annual reduction in the farmers’ income has been estimated to amount to 158 EUR per farmer. However, as explained under section 5.7.1, it is expected in line with previous trends (discontinuation of direct subsidies) that certain farmers will discontinue their activities irrespective of the TPD revision. The preferred options would not discriminate between growers of different tobacco types (Oriental, Burley, Virginia).

Similarly to the FMC and RYO manufacturers, the **packaging and labelling** industry is expected to benefit from the “one go” solution at EU level compared to multiple adaptations under the baseline scenario. The industry reported to have carried out the main investments already. On the other hand, a negative impact on revenues of around 18.2 mEUR is estimated following the reduced tobacco consumption. This represents 0.2% of the total sector turnover. A similar reasoning applies to **producers of additives and flavours**.

### 6.2.1.3. Downstream sectors and other industries

**Wholesalers** are primarily affected indirectly by the reduction in tobacco consumption resulting in reduced revenues of around 465 mEUR\(^{403}\) and by compliance with the tracking and tracing system (no cost estimate available). This needs to be counterbalanced, however, with the increased revenues stemming from reduced illicit trade benefitting wholesalers. The introduction of EU tracking and tracing systems will call for some technical adjustments in warehouses, but the costs for the equipment should normally be limited as they should be borne by the tobacco industry, which is under the obligation to introduce the tracking and tracing system. The introduction of a common EU wide system is beneficial for the wholesalers, as one system takes less space and is easier to handle for its staff.

\(^{402}\) As pointed out in sections 5.3.1, the cost increases reported by industry seem exaggerated.

\(^{403}\) Obviously, the wholesalers would have reduced purchasing costs in the region of 376 mEUR.
Retailers are in a similar situation as wholesalers regarding the indirect impacts. The measures on tracking and tracing and security features are not expected to lead to additional costs for retailers as retailers are not part of the tracking and tracing system. As even specialised tobacco retailers generate only up to 50% of their revenues from tobacco products (dependence of other retailers on tobacco products, e.g. supermarkets, is even lower), the impact is not expected to be disproportionate.

A variety of other industries are expected to benefit from certain of the preferred policy options, e.g. companies developing tracking and tracing systems or security features.

6.2.1.4. Small and medium sized enterprises (SMEs)

The particular situation of micro-enterprises and SMEs has been taken into account when identifying the preferred option. The preferred policy options are primarily targeting FMC, RYO and STP. Pipe tobacco and cigars which are often manufactured by SMEs are not affected, in the first stage, in most policy areas. As referred to above, the impact on growers is limited. The impact on retailers is also expected to be limited as referred to above, in particular since the policy area/option of restricting display of tobacco at point of sale was discarded (see section 4.1). Products other than FMC and RYO which will be included in the tracking, tracing and security regime will be granted an additional transitional period which will benefit primarily SMEs.

1.1.36. Employment

In terms of employment it is estimated that jobs lost in tobacco will be off-set by jobs gained in other sectors (input/output model), as money not spent on tobacco is spent on other goods/services. This will be a gradual process as tobacco consumption is expected to decrease over time (2% in five years). The input/output model suggests that a 2% reduction of tobacco consumption would lead to a net gain in employment of 2,235 employees in the EU taking into account that tobacco is not a labour intensive industry whilst a number of the sectors benefitting from the redistribution are (e.g. hospitality sector). Possible regional employment impacts, as well as the specific situations of SMEs and micro-enterprises, have been carefully considered when formulating the preferred options. Obviously it cannot be excluded that some tobacco manufacturers will use the TPD revision as a welcome excuse to further automate production/concentration and would therefore lead to factory closure. On the other hand, keeping people healthy and active for longer has a positive impact on the productivity and competitiveness in general and is an integral part of Europe 2020 strategy. In the light of an ageing population in the EU it also appears important to assist young people staying healthy.

1.1.37. Governments/Society

The main benefit for Governments is the improvement of health which is a value in its own right. The preferred options are expected to result in a 2% drop in consumption within a five year period, corresponding to 2.4 million smokers. The expected socio-economic benefits (reduction of health care costs, productivity losses and monetised life years saved) represents an annual amount to the EU of 9.4 bEUR (see section 5.7) even if one deducts reduced tax revenues. A somewhat lower figure would be achieved if discounting is applied.

The preferred options are also associated with a number of administrative costs for national authorities. With respect to the measures envisaged in the scope section (STP, NCP and herbal products for smoking), a limited additional burden is expected when assessing NCP under the medicinal products' framework, however in a well-established legal framework and against fees. For labelling, no major impact is expected for Governments, possibly with one exception, namely that additional resources might be needed for cessation services. Regarding
ingredients reporting, costs savings associated with the improved reporting system and additional revenues in terms of fees from the industry are expected. Regarding ingredients regulation some investment might be needed to develop further – in conjunction with the Commission – the concept of attractiveness. Regarding cross-border distance sales the notification system would lead to very limited additional costs, but will also facilitate law enforcement. Also regarding traceability and security features, the main direct impact is on law enforcement, but overall it is assumed that law enforcement is facilitated/simplified by the proposed measures, i.e. the costs for Governments should be reduced.

7. **MONITORING AND EVALUATION**

The successful implementation of the revised TPD will depend on several factors:

*Transposition of the Directive*

Member States need to transpose the revised TPD correctly and in time. The European Commission should assist Member States in this exercise through meetings where Member States report about transposition progress and can discuss problems and seek clarifications. Member States should also be encouraged to communicate their draft transposition measures to the Commission.

An implementation plan will be developed to ensure effective implementation of the Directive.

*Indicators*

A key indicator for achievement of the objectives outlined in this impact assessment is the implementation by Member States, infringement cases launched and the number of complaints.

In addition, the following indicators will be monitored on a regular basis
- the smoking consumption and prevalence in the EU, including smoking uptake in young people
- the awareness of the harmful effects of tobacco
- the number of novel tobacco, nicotine and niche products as reported by Member States
- the frequency of use of quit lines/cessation services
- the number of electronic cigarettes authorised as medicinal products
- the consumption of herbal products for smoking
- the number of seizures of illegal tobacco products reported by Member States
- the level of additives used in marketed products and their variations between Member States
- the number of cross-border distance sale notifications.

*Consultation and reporting*

A network of Member States will provide a regular platform to discuss issues related to the implementation of the Directive.

The Commission should report to the European Parliament and the Council about the implementation of the revised TPD five years after transposition. The report should address the impact of the new rules in respect of the internal market, public health and international developments. In particular, the Commission should report on international, legal and scientific developments in terms of labelling and packaging as well as on novel tobacco products and, if appropriate, provide suggestions for further EU action in this area.