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EDITOR’S PREFACE

Pre-merger competition review has advanced significantly since its creation in 1976 in the United States. As this book evidences, today almost all competition authorities have a notification process in place – with most requiring pre-merger notification for transactions that meet certain prescribed minimum thresholds. Additional jurisdictions, particularly in Asia, are poised to add pre-merger notification regimes within the next year or so. In our endeavour to keep our readers well informed, we have expanded the jurisdictions covered by this book to include the newer regimes as well.

Given the ability of most competition agencies with pre-merger notification laws to delay, and even block, a transaction, it is imperative to take each jurisdiction – small or large, new or mature – seriously. For instance, in 2009, China blocked the Coca-Cola Company’s proposed acquisition of China Huiyuan Juice Group Limited and imposed conditions on four mergers involving non-China-domiciled firms. In Phonak/ReSound (a merger between a Swiss undertaking and a Danish undertaking, each with a German subsidiary), the German Federal Cartel Office blocked the entire merger, even though less than 10 per cent of each of the undertakings was attributable to Germany. It is, therefore, imperative that counsel for such a transaction develops a comprehensive plan prior to, or immediately upon, execution of an agreement concerning where and when to file notification with competition authorities regarding such a transaction. To this end, this book provides an overview of the process in 41 jurisdictions, as well as a discussion of recent decisions, strategic considerations and likely upcoming developments. Given the number of recent significant M&A transactions involving media, pharma and high-technology companies, we have included chapters that focus on the enforcement trends in these important sectors. In addition, as merger review increasingly includes economic analysis in most, if not all, jurisdictions, we have added a chapter that discusses the various economic tools used to analyse transactions. The intended readership of this book comprises both in-house and outside counsel who may be involved in the competition review of cross-border transactions.

Some common threads in institutional design underlie most of the merger review mandates, although there are some outliers as well as nuances that necessitate careful consideration when advising a client on a particular transaction. Almost all jurisdictions vest exclusive authority to review transactions in one agency. The United States and China may
end up being the exceptions in this regard. Most jurisdictions provide for objective monetary size thresholds (e.g., the turnover of the parties, the size of the transaction) to determine whether a filing is required. Germany, for instance, provides for a *de minimis* exception for transactions occurring in markets with sales of less than €15 million. There are some jurisdictions, however, that still use ‘market share’ indicia (e.g., Bosnia and Herzegovina, Colombia, Lithuania, Portugal, Spain, Ukraine and the United Kingdom). Most jurisdictions require that both parties have some turnover or nexus to their jurisdiction. However, there are some jurisdictions that take a more expansive view. For instance, in Poland, a notification may be required even though only one of the parties is present and, therefore, there may not be an impact on competition in Poland. Turkey recently issued a decision finding that a joint venture (JV) that produced no effect on Turkish markets was reportable because the JV’s products ‘could be’ imported into Turkey. Germany also takes an expansive view by adopting as one of its thresholds a transaction of ‘competitively significant influence’. Although a few merger notification jurisdictions remain ‘voluntary’ (e.g., Australia, Singapore, the United Kingdom and Venezuela), the vast majority impose mandatory notification requirements. Moreover, in Singapore, the transaction parties are to undertake a ‘self-assessment’ of whether the transaction will meet certain levels, and, if so, should notify the agency to avoid potential challenge by the agency.

Although in most jurisdictions the focus of the competition agency is on competition issues, some jurisdictions have a broader mandate. For instance, the ‘public interest’ approach in South Africa expressly provides for consideration of employment matters, local enterprises and procurement, and for economic empowerment of the black population and their participation in the company. Many of the remedies imposed in South Africa this year have been in connection with these considerations. Although a number of jurisdictions have separate regulations and processes for addressing foreign entity acquisitions when national security or specific industrial sectors are involved, in Romania, for example, the competition law provides that the government can prohibit a merger if it determines that such merger could have a potential impact on national security.

The potential consequences for failing to file in jurisdictions with mandatory requirements vary. Almost all jurisdictions require that the notification process be concluded prior to completion (e.g., pre-merger, suspensory regimes), rather than permitting the transaction to close as long as notification is made prior to closing. Many of these jurisdictions can impose a significant fine for failure to notify before closing, even where the transaction raises no competition concerns (e.g., Austria, Cyprus, India, the Netherlands, Romania, Spain and Turkey). In France, for instance, the competition authority imposed a €4 million fine on Castel Frères for failure to notify its acquisition of part of the Patriache group. In Ukraine, the competition authority focused its efforts on discovering consummated transactions that had not been notified, and imposed fines in 32 such cases in 2015 alone.

Some jurisdictions impose strict time frames within which the parties must file their notification. For instance, Cyprus requires filing within one week of signing of the relevant documents and agreements; Serbia and India provide for 15 days after signing of the agreement; and Hungary, Ireland and Romania have a 30-calendar-day time limit for filing the notification that commences with entering into the agreement. Some jurisdictions that mandate filings within specified periods after execution of the agreement also have the authority to impose fines for ‘late’ notifications (e.g., Bosnia and Herzegovina, Indonesia, India and Serbia). Most jurisdictions also have the ability to impose significant fines for failure to notify or for closing before the end of the waiting period, or both (e.g., Austria, Canada,
China, Greece, Portugal, Ukraine and the United States). In Macedonia, the failure to file can result in a misdemeanour and a monetary fine of up to 10 per cent of the worldwide turnover. In Belgium, the competition authority fined a party for late submission of information.

In addition, other jurisdictions have joined the EC and the United States in focusing on interim conduct of the transaction parties. Brazil, for instance, issued its first ‘gun-jumping’ fine in 2014 and recently issued guidelines on gun-jumping violations. In most jurisdictions, a transaction that does not meet the pre-merger notification thresholds is not subject to review or challenge by the competition authority. In Canada – like the United States – however, the Canadian Competition Bureau can challenge mergers that were not required to be notified under the pre-merger statute. In Korea, Microsoft initially filed a notification with the Korea Fair Trade Commission (KFTC), but when it faced difficulties and delays in Korea the parties restructured the acquisition to render the transaction nonreportable in Korea and consummated the transaction. The KFTC, however, continued its investigation as a post-consummation merger investigation and eventually obtained a consent order.

In almost all jurisdictions, very few transactions undergo a full investigation, although some require that the notification provide detailed information regarding the markets, competitors, competition, suppliers, customers and entry conditions. Most jurisdictions that have filing fees specify a flat fee or state in advance a schedule of fees based upon the size of the transaction; some jurisdictions, however, determine the fee after filing or provide different fees based on the complexity of the transaction. For instance, Cyprus is now considering charging a higher fee for acquisitions that are subjected to a full Phase II investigation.

Most jurisdictions more closely resemble the EC model than the United States model. In these jurisdictions, pre-filing consultations are more common (and even encouraged); parties can offer undertakings during the initial stage to resolve competitive concerns; and there is a set period during the second phase for providing additional information and for the agency to reach a decision. In Japan, however, the Japan Federal Trade Commission (JFTC) announced in June 2011 that it would abolish the prior consultation procedure option. When combined with the inability to ‘stop the clock’ on the review periods, counsel may find it more challenging in transactions involving multiple filings to avoid the potential for the entry of conflicting remedies or even a prohibition decision at the end of a JFTC review. Some jurisdictions, such as Croatia, are still aligning their threshold criteria and processes with the EC model. Some jurisdictions even within the EC remain that differ procedurally from the EC model. For instance, in Austria, the obligation to file can be triggered if only one of the involved undertakings has sales in Austria, as long as both parties satisfy a minimum global turnover and have a sizeable combined turnover in Austria.

The role of third parties also varies across jurisdictions. In some jurisdictions (e.g., Japan), there is no explicit right of intervention by third parties, but the authorities can choose to allow it on a case-by-case basis. In contrast, in South Africa, registered trade unions or representatives of employees must be provided with a redacted copy of the merger notification from the outset and have the right to participate in merger hearings before the Competition Tribunal: the Tribunal will typically also permit other third parties to participate. Bulgaria has announced a process by which transaction parties even consent to disclosure of their confidential information to third parties. In some jurisdictions (e.g., Australia, the EC and Germany), third parties may file an objection to a clearance decision. In some jurisdictions (including Canada, the EC and the United States), third parties (e.g., competitors) are required to provide information and data if requested by the antitrust authority. In Israel, a third party that did not comply with such a request was recently fined by the authority.
In almost all jurisdictions, once the authority approves the transaction, it cannot later challenge the transaction's legality. The United States is one significant outlier with no bar for subsequent challenge, even decades following the closing, if the transaction is later believed to have substantially lessened competition. Canada, in contrast, provides a more limited time period of one year for challenging a notified transaction (see the recent CSC/Complete transaction). Norway is a bit unusual, where the authority has the ability to mandate notification of a transaction for a period of up to three months following the transaction’s consummation. In ‘voluntary’ jurisdictions, such as Australia and Singapore, the competition agency can investigate and challenge unnotified transactions.

It is becoming the norm in large cross-border transactions raising competition concerns for the United States, Canadian, Mexican and EC authorities to work closely together during the investigative stages, and even in determining remedies, minimising the potential of arriving at diverging outcomes. The KFTC has stated that it will engage in even greater cooperation with foreign competition authorities, particularly those of China and Japan, which are similar to Korea in their industrial structure. Regional cooperation among some of the newer agencies has also become more common; for example, the Argentinian authority has worked with Brazil’s CADE, which in turn has worked with the Chilean authority. Competition authorities in Bosnia and Herzegovina, Bulgaria, Croatia, Macedonia, Montenegro, Serbia, Slovenia and Turkey similarly maintain close ties and cooperate on transactions. Taiwan is part of the Asia-Pacific Economic Cooperation Forum, which shares a database. In transactions not requiring filings in multiple European jurisdictions, Member States often keep each other informed during the course of an investigation. In addition, transactions not meeting the EC threshold can nevertheless be referred to the European Commission in appropriate circumstances. The United States has signed cooperation agreements with a number of jurisdictions, including most recently Peru and India. China has ‘consulted’ with the United States and the EC on some mergers and entered into a cooperation agreement with the United States authorities in 2011.

The impact of such multijurisdictional cooperation was very evident this year. For instance, the transaction parties in Applied Materials/Tokyo Electron ultimately abandoned the transaction due to the combined objections of several jurisdictions, including the United States, Europe, and Korea. In Office Depot/Staples, the FTC and the Canadian Competition Bureau cooperated and both jurisdictions brought suits to block the transaction (although the EC had also cooperated on this transaction, it ultimately accepted the undertakings offered by the parties). In the GE/Alstom transaction, the United States and the EC coordinated throughout, including at the remedies stage. Additionally, in the Halliburton/Baker Hughes transaction, the United States and the EC coordinated their investigations, with the United States suing to block the transaction while the EC’s investigation continued. Also, in Holcim/Lafarge, the cooperation between the United States and Canada continued at the remedies stage, where both consents included assets in the other jurisdiction’s territory. The United States, Canada and Mexico coordinated closely in the review of the Continental/Veyance transaction. In fact, it is becoming the norm for coordination among the jurisdictions in multinational transactions that raise competition issues.

Although some jurisdictions have recently raised the size threshold at which filings are mandated, others have broadened the scope of their legislation to include, for instance, partial ownership interests. Some jurisdictions continue to have as their threshold test for pre-merger notification whether there is an ‘acquisition of control’. Many of these jurisdictions, however, will include, as a reportable situation, the creation of ‘joint control’, ‘negative (e.g., veto) control’ rights to the extent that they may give rise to de jure or de facto control (e.g., Turkey),
Editor's Preface

or a change from ‘joint control’ to ‘sole control’ (e.g., the EC and Lithuania). Minority holdings and concerns over ‘creeping acquisitions’, in which an industry may consolidate before the agencies become fully aware, have become the focus of many jurisdictions. Some jurisdictions will consider as reviewable acquisitions in which only a 10 per cent or less interest is being acquired (e.g., Serbia for certain financial and insurance mergers), although most jurisdictions have somewhat higher thresholds (e.g., Korea sets the threshold at 15 per cent of a public company and otherwise at 20 per cent of a target; and Japan and Russia at any amount exceeding 20 per cent of the target). Others use, as the benchmark, the impact that the partial shareholding has on competition; Norway, for instance, can challenge a minority shareholding that creates or strengthens a significant restriction on competition. The UK also focuses on whether the minority shareholder has ‘material influence’ (i.e., the ability to make or influence commercial policy) over the entity. Several agencies during the past few years have analysed partial ownership acquisitions on a stand-alone basis as well as in connection with JVs (e.g., Canada, China, Cyprus, Finland and Switzerland). Vertical mergers were also a subject of review (and even resulted in some enforcement actions) in a number of jurisdictions (e.g., Belgium, Canada, China, Sweden and Taiwan). Portugal even viewed as an ‘acquisition’ subject to notification the non-binding transfer of a customer base.

For transactions that raise competition issues, the need to plan and to coordinate among counsel has become particularly acute. Multi-jurisdictional cooperation facilitates the development of cross-border remedies packages that effectively address competitive concerns while permitting the transaction to proceed. The consents adopted by the United States and Canada in the Holcim/Lafarge merger exemplify such a cross-border package. As discussed in the International Merger Remedies chapter, it is no longer prudent to focus merely on the larger mature authorities, with the expectation that other jurisdictions will follow their lead or defer to their review. In the current enforcement environment, obtaining the approval of jurisdictions such as Brazil and China can be as important as the approval of the EC or the United States. Moreover, the need to coordinate is particularly acute to the extent that multiple agencies decide to impose conditions on the transaction. Although most jurisdictions indicate that ‘structural’ remedies are preferable to ‘behavioural’ conditions, a number of jurisdictions in the past few years have imposed a variety of such behavioural remedies (e.g., China, the EC, France, the Netherlands, Norway, South Africa, Ukraine and the United States). For instance, some recent decisions have included as behavioural remedies pricing, sales tariffs and terms of sale conditions (e.g., Korea, Ukraine and Serbia), employee retrenchment (South Africa) and restrictions on bringing antidumping suits (e.g., Mexico). Many recent decisions have imposed behavioural remedies to strengthen the effectiveness of divestitures (e.g., Canada’s decision in the Loblaw/Shoppers transaction, China’s MOFCOM remedy in Glencore/Xstrata, and France’s decision in the Numericable/SFR transaction). This book should provide a useful starting point in navigating cross-border transactions in the current enforcement environment.

Ilene Knable Gotts
Wachtell, Lipton, Rosen & Katz
New York
July 2016
Chapter 3

EU MERGER CONTROL IN THE PHARMACEUTICAL SECTOR

Pablo Figueroa and Alejandro Guerrero

I INTRODUCTION

The combination of relatively stagnant markets in the EU with the foreseen extraordinary growth in the emerging markets is likely to result in further consolidation in the pharmaceutical industry. Regulators will be crucial to this consolidation. This chapter summarises the approach of the European Commission to merger control in the pharmaceutical industry.

II MARKET DEFINITION

Defining the relevant market and the calculation of market shares is necessary to analyse market power and the potential impact of behaviour on competition. In the Commission’s words: ‘[m]arket definition is a tool to identify and define the boundaries of competition between firms. It serves to establish the framework within which competition policy is applied by the Commission. The main purpose of market definition is to identify in a systematic way the competitive constraints that the undertakings involved face.’ The calculation of market

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1 Pablo Figueroa is a senior associate and Alejandro Guerrero is an associate at Gibson, Dunn & Crutcher LLP. The usual disclaimers apply.
2 According to a report by PricewaterhouseCoopers, the global pharmaceutical industry could be worth nearly $1.6 trillion by 2020 (see PwC, ‘The Global Pharmaceutical Market could be Worth Nearly $1.6 trillion by 2020’, available at www.pwc.com/gx/en/pharma-life-sciences/pharma2020/market-opportunities-and-outlook.jhtml).
shares, by reference to a previously defined market, is an exercise that the Commission considers to provide ‘useful first indications of the market structure and of the competitive importance of both the merging parties and their competitors’.4

The principles set out in the Commission’s Notice on the Definition of the Relevant Market5 have been applied to the pharmaceutical sector by the European institutions.6

i Product market definition for finished dose pharmaceutical products

The Commission has traditionally resorted to different parameters to define relevant markets for finished dose pharmaceuticals.

The European Pharmaceutical Market Research Association Anatomical Therapeutical Chemical Classification (EPhMRA’s ATC)

In past years,7 the Commission has resorted to the EPhMRA’s ATC8 classification, which has been developed for marketing purposes and is the basis for the pharmaceutical sales Intercontinental Medical Statistics database (IMS Health), which provides data that are often used by global pharmaceutical and biotechnology companies for econometric market analysis.9

The EPhMRA ATC classifies pharmaceutical products10 according to their indications and use,11 distinguishing the following four levels:

a the first level of the code indicates the anatomical main group (i.e., the part of the human body that the medicine intends to address);
b the second level of the code indicates the therapeutic main group (i.e., the main disease groups that the medicine intends to address);
c the third level of the code indicates the therapeutic and pharmacological subgroup (i.e., the different drug actions that will address the disease in question); and
d the fourth level of the code indicates the chemical subgroup.

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7 See Case COMP/M.5865 – Teva/Ratiopharm; Case COMP/M.6613 – Watson/Actavis; Case COMP/M.6969 – Valeant Pharmaceutical International/Bausch & Lomb Holdings; Case COMP/M.7379 – Mylan/Abbott EPD-DM.
8 See ‘EphMRA/PBIRG Classification Committee; who we are; what we do 2008’, available at www.ephmra.org/user_uploads/ephmra%20who%20we%20are%202015%20final.pdf.
9 IMS Health is a private entity that provides information and services for the healthcare industry (see www.imshealth.com/portal/site).
10 This distinguishes it from the WHO classification system, which classifies substances according to the therapeutic or pharmaceutical aspects and in one class only. The main purpose of the WHO classification is for international drug utilisation research and for adverse drug reaction monitoring.
11 It is therefore possible to find the same compound in several classes, depending on the product, e.g., naproxen tablets can be classified in M1A (antirheumatic), N2B (analgesic) and G2C if indicated for gynaecological conditions only.
The Commission starts its analysis at different levels, often depending on whether the transaction involves producers of originators or generics.

When it comes to originator companies, the Commission’s analysis tends to begin at the ATC3 level, which, as indicated in (c) above, groups medicines according to their broad therapeutic and pharmacological indications.\(^\text{12}\)

By contrast, in many recent merger cases involving mature genericised markets (i.e., markets involving products with regard to which originators’ patent protection has expired and generic companies are offering alternative generic products), the Commission has found classification by molecule (e.g., at ATC4 level) to be more accurate for the purposes of defining the relevant market.\(^\text{13}\)

In mergers involving originators and generic producers, the Commission may identify a certain degree of substitutability between the molecules used by originators and generic producers (see Section II.i, Originators, generics and biosimilars, infra). In these scenarios, the Commission might take into account the closeness of substitution between these molecules, and consider the generic molecule as being the closest substitute to the ex-originator drug based on the same molecule.\(^\text{14}\)

**Prescription medicines, over-the-counter (OTC) and dual-status medicines**

The Commission has usually defined separate markets for prescription medicines and for OTC or dual-status medicines.\(^\text{15}\) This is due to the fact that seriousness of disease (i.e., medical indications or dosage, or both, in some cases), strength of products (including possible side effects and harmfulness if misused), legal framework, marketing, distribution, the medical indications (including their possible side effects), legal framework, distribution and rules on reimbursement of drugs all tend to differ between the two categories of medicines, even when the active ingredients are identical.

Certain variants of a drug with the same active ingredient or brand name are sometimes classified as both OTC and prescription-only, depending on the package size, dosage or ‘galenic’ form.\(^\text{16}\) In these cases, the price of the OTC medicine may be a factor determining whether the patient simply purchases this medicine at his or her own expense.

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\(^{12}\) See, e.g., Case COMP/M.5253 – *Sanofi-Aventis/Zentiva*, at paragraphs 12 ff; Case COMP/M.5295 – *Teva/Barr*, at paragraphs 10 ff. In the context of originator products, the Commission has sometimes resorted to the EPhMRA ATC4 level (see Case COMP/M.3544 – *Bayer Healthcare/Roche (OTC Business)*, at paragraphs 15–20).

\(^{13}\) See Case COMP/M.6613 – *Watson/Actavis*, at paragraph 7; Case COMP/M.5295 – *Teva/Barr*, at paragraph 18; Case COMP/M.5865 – *Teva/Ratiopharm*, at paragraph 12; Case COMP/M.7379 – *Mylan/Abbott EPD-DM*, at paragraph 13. However, the Commission has sometimes defined markets on the basis of the molecule level or group of molecules that are interchangeable for a wide range of applications (see Case COMP/M.5865 – *Teva/Ratiopharm*, at paragraph 14).

\(^{14}\) See Case COMP/M.7379 – *Mylan/Abbott EPD-DM*, at paragraph 13. This possibility had already been explored in Case COMP/M.5865 – *Teva/Ratiopharm*, at paragraph 13.

\(^{15}\) Ibid, at paragraphs 22 ff.

\(^{16}\) Ibid.
or visits a doctor to obtain a prescription for a reimbursable alternative. Similarly, in some specific circumstances where the status of a drug is not clearly limited to either OTC or prescription, it may not be excluded that these products compete with each other. Finally, the presence of a product or brand in both the prescription and OTC segments may result in it enjoying a stronger market position.

**Originators, generics and biosimilars**

Broadly speaking, there are two types of pharmaceuticals that aim to offer alternatives to originator drugs: (1) synthetic small-molecule generics; and (2) biosimilar products.

Small-molecule generic products are based on the same active principle as that of their equivalent small-molecule originator drugs and are synthesised by chemical processes. Generic products are produced and offered upon patent expiry of the originator that they aim to reproduce. Small-molecule originators and generic drugs can generally be considered homogeneous products, to the extent that the Commission has recently found that they compete mainly on price, particularly in the case of hospital drugs procured through tenders.

While the Commission’s market investigations have often suggested that there may be differences in demand for originators versus generics, this phenomenon has not been found to justify the definition of two separate product markets. According to the Commission, a number of elements indicate that generic medicines based on the same molecule compete in the same product market as the branded originator medicines on which they are based. These elements include:

- the fact that, in order to obtain regulatory approval to market its product, a generic drug manufacturer must demonstrate that its drug is bioequivalent to the originator

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17 See Case COMP/M.5253 – Sanofi-Aventis/Zeniwa, at paragraphs 58–59. For OTC products, the Commission has also recently taken into consideration the classification used in the IMS Consumer Health’s OTC Review Reports (see, e.g., Case COMP/M.6280 – Procter & Gamble/Teva OTC business, at paragraphs 9 and 11).


19 See, e.g., Case COMP/M.1846 – Glaxo Wellcome/Smithkline Beecham, at paragraphs 98–113.

20 See Case COMP/M.7559 – Pfizer/Hospira, at paragraph 33.


22 See Case COMP/M.5476 – Pfizer/Wyeth, at paragraph 19; Case COMP/M.7379 – Mylan/Abbott EPD-DM, at paragraph 16; Case COMP/M.7559 – Pfizer/Hospira, at paragraph 33. However, generic companies might also compete with products based on other molecules (see Case COMP/M.5865 – Teva/Ratiopharm, at paragraph 13; Case COMP/M.7379 – Mylan/Abbott EPD-DM, at paragraph 13).
drug\(^{23}\) (i.e., the generic is bioequivalent in dosage form, safety, strength, route of administration, quality, performance characteristics and intended use, thus working in essentially the same way as the originator in the patient’s body);\(^{24}\) the fact that generic versions of originator medicines are often designed by generic producers to be copies of those originator medicines on which they are based (indeed, only bioequivalence can ensure generics the fast route to the market, meaning that they main obtain a marketing authorisation by simply showing that they are bioequivalent to the originator drugs, which thus dispenses them from carrying on the full clinical trials that an originator product would have to do); and the applicable regulatory framework, which encourages switching between originator and generic medicines.

Separately, in the past years, growing competition has been observed between originator biological products (biopharmaceuticals) and biosimilar products.\(^{25}\) Unlike small molecule generics, bio-similars are not exact copies of the originator drugs. However, biosimilar drugs aim to have the same therapeutic mechanism as original patented medicines.\(^{26}\) According to the guidelines of the European Medicines Agency (EMA), in order to obtain a marketing authorisation for a biosimilar, its manufacturer needs to demonstrate similarity (in terms of quality, safety and efficacy) to a reference biological product.\(^{27}\)

The Commission has observed that, since the first complex bio-similar was approved in Europe in 2013, the offering of biosimilar products has led to price decreases compared to originator products.\(^{28}\) Biosimilars are expected to allow wider access by patients to biological drugs and to be an important factor in relieving the financial pressure on healthcare systems. Accordingly, in recent cases the Commission has found that an originator biopharmaceutical and its biosimilar product belonged to the same product market.\(^{29}\)

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23 Bioequivalence is defined by the European Medicines Agency as the absence of a significant difference in the rate and extent to which the active ingredient or active moiety in pharmaceutical equivalents or pharmaceutical alternatives becomes available at the site of action when administered at the same molar dose under similar conditions in an appropriately designed study.

24 As regards competition between generic medicines, generic medicines developed to be bioequivalent of the same originator product, and based on the same molecule, may still have some differentiating elements from one another (e.g., inactive ingredients or impurities). However, these differences have not been found sufficient to define separate markets for two generics falling under the same molecule: see Case COMP/M.5253 – Sanofi-Aventis/Zentiva, at paragraph 26.

25 Biopharmaceuticals are originator medicines whose active substance is made by or derived from living organisms.

26 See Case COMP/M.7559 – Pfizer/Hospira, at paragraphs 9 and 34–35.


28 See Case COMP/M.7559 – Pfizer/Hospira, at paragraph 11.

29 See Case COMP/M.7559 – Pfizer/Hospira, at paragraph 25, where the Commission found the relevant product market to comprise infliximab pharmaceuticals, including both the
However, competition among biosimilars and biopharmaceuticals may not always be homogeneous. Because of the possible differences between originator biological products and biosimilar products in terms of clinical evidence available on their efficacy and safety, physicians and pharmacists might not necessarily consider originator and biosimilars based on the same biologic molecule to be fully interchangeable, depending on the Member State. This may equally apply to the interchangeability amongst biosimilars based on the same molecule, which are also not identical in their chemical structure and clinical evidence. Accordingly, competition between original biologic products and biosimilar products, as well as between any pair of biosimilar products, may be characterised by a limited degree of substitutability for patients already undergoing treatment, and a high degree of substitutability for new patients.

**Biosimilar and small molecule traditional generics**
The Commission has also reviewed in detail the distinction between bio-similar products and small molecule traditional generics. The Commission has identified a number of differences between these types of products, which has sometimes led to their individual assessment:

- the development of biosimilars tends to require considerably longer development periods than synthetic generics;
- the development of biosimilars tends to require higher upfront investments than those required for the development of other generics;
- the development of biosimilars entails a higher risk of failure for research and development (R&D);
- the development and manufacturing of biosimilars requires specific biotech know-how and facilities; and
- the R&D process for biosimilars is closer to the R&D of originator than of synthetic generic drugs requiring, for example, clinical trials.

**Galenic formulation**
In recent cases that have involved generics, the Commission has investigated whether a difference in the ‘galenic’ formulation further limits substitutability within or across molecules. The Commission has found this to be the case in a number of cases, to the extent that the launch of a new ‘galenic’ form may take up to two or three years to appear in the market.

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30 See Case COMP/M.7559 – Pfizer/Hospira, at paragraphs 35-38.
31 Ibid.
33 See Case COMP/M. 5865 – Teva/Ratiopharm, at paragraph 29; Case COMP/M.5479 – Lonza/Teva JV, at paragraph 7; Case COMP/M.7559 – Pfizer/Hospira, at paragraph 18.
34 See, e.g., Case COMP/M.5865 – Teva/Ratiopharm, at paragraphs 16–21 and 39–41: ‘[…] in this case, different routes of administration of a medicine are, in general, […] not interchangeable. This may also be the case of the dosage and of the pharmaceutical form […].’ Lack of substitutability was mainly found for oral syrups, rectal forms, and injectable or parenteral forms (see paragraphs 19, 118, 157, 184, 253–256, 267–268, 279, and 336–340).
market, a period that may exclude supply-side substitutability; and that different routes of administration for a medicine are, in general, designed to serve the needs of different patient groups, and are therefore not interchangeable. In these situations, the reference system for distinguishing between medicinal formulations is the typology of form code (also called the New Form Code (NFC)) used by IMS Health and EphMRA.

**Product 'pipelines', innovation and R&D**

In accordance with the Commission’s general guidance, R&D and product pipelines should be considered in markets and sectors, such as the pharmaceutical sector, where innovation is an important competitive force that may be driven or impeded depending on the particular circumstances of each case. In such scenarios, a full competitive analysis requires that the relevant authority examine those products that have not as yet entered the market, but that are at an advanced stage of development (e.g., advanced R&D pipeline products in Phase III of clinical trials). In a number of cases, the Commission has assessed the impact of transactions in ‘pipeline products’. Patents and other IP rights also play an important role in the competitive assessment of current and future markets.

**National registration and reimbursement rules**

In certain cases, the Commission has taken into consideration the influence of national registration and national reimbursement rules on the prescription behaviour of physicians for the purposes of defining markets.

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36 See Case COMP/M.7379 – Mylan/Abbott EPD-DM, at paragraph 24. For example, the liquid form of certain drugs (such as syrups), has been considered to be mainly designed for paediatric patients.
37 See http://www.ephmra.org/New-Form-Codes-Classification.
38 See Guidelines on the Assessment of Horizontal Mergers at paragraph 38.
39 See, e.g., Case COMP/M.1403 – Astral/Zeneca, at paragraphs 43 and 44; Case IV/M.1846 – Glaxo Wellcome/Smithkline Beecham, at paragraphs 150–216; Case COMP/M.3354 – Sanofi-Synthélabo/Aventis, at paragraphs 324–330; COMP/M.5476 – Pfizer/Wyeth, at paragraphs 13, 34–38, 65, 87–95, and 99; and Case COMP/M.5999 – Sanofi-Aventis/Genzyme, at paragraphs 7, 21, 29, and 38–46. However, in Case COMP/M.7275 – Novartis/GlaxoSmithKline Oncology Business, the Commission analysed markets where the parties had ongoing clinical trials at earlier stages, including Phase I and Phase II.
42 See, e.g., Case COMP/M.7275 – Novartis/GlaxoSmithKline Oncology Business, at paragraphs 123-124; Case COMP/M.5865 – Teva/Ratiopharm, at paragraph 90.
Other medicine characteristics
Further segmentation of product markets in the pharmaceutical sector may result from a number of characteristics of medicines (e.g., the medicine’s indications and contraindications, their efficacy and side effects, their frequency of administration and period of action).  

ii Markets upstream and downstream from finished pharmaceuticals
In addition to the markets defined for finished pharmaceuticals set out above, the Commission has also identified the neighbouring markets, some of which are set out below.

Active pharmaceutical ingredients (API) markets and other raw materials
In its precedents, the Commission has identified separate markets for APIs, which is one of the main raw materials used by pharmaceutical companies for the manufacture of drugs. The Commission has generally found that API markets might be as narrow as each individual API (i.e., the relevant molecule). Sufficient supply-side substitutability may, however, justify the inclusion of a number of APIs in the same relevant product market. However, the following elements might lead to the opposite conclusion:

a demand-side substitutability is unlikely in practice, given that a generic company that produces a specific pharmaceutical product needs to buy the relevant API molecule and lacks the ability to use another alternative API; and

b switching to another API source might require variations in the relevant authorisations. Obtaining these can involve devoting significant resources.

Other raw materials used for the manufacture of pharmaceutical products include: (1) excipients (non-active ingredients used in a final drug dosage form, such as binders, fillers, diluents, lubricants, flavours, solvents sweeteners or preservatives, with inter alia lactose, starch, cellulose, magnesium, stearic acid, gelatine, sucrose, talc or sodium); and (2) biopharm ingredients (process chemicals used in biopharmaceutical processes, including buffers and stabilisers – such as amino acids, carbohydrates and polymers).
Contract manufacturing of finished dose pharmaceuticals
The Commission has identified a separate upstream market for the contract manufacturing of finished dose pharmaceuticals on behalf of third-party pharmaceutical companies. In the past, the Commission has considered, but ultimately left open, the possibility that this market be further segmented by reference to the function of the pharmaceutical form (e.g., solids, powder, liquids, sterile liquids) or by reference to the conditions of manufacture (e.g., types of APIs involved, toxicity, involvement of a sterile environment, etc.).

Out-licensing
In the out-licensing market, one party (the licensor) out-licenses a pharmaceutical product to one or more third parties (the licensee or licensees). During the duration of the licence, the licensee will generally buy the finished product (or ‘bulk’) from the licensor on an exclusive basis and will commercialise the product under its own name, using the marketing authorisation that was licensed to it by the licensor. The Commission has considered a possible narrower segmentation within the out-licensing market for the out-licensing of IPRs for particular APIs or pharmaceutical products, or both.

Other upstream markets
The Commission has identified other separate markets, such as input markets for substances required for the production of APIs, and dosage delivery mechanisms.

Markets downstream from finished pharmaceuticals
Pharmaceutical companies reach distributors and end users through different distribution channels characterised by very different competitive dynamics. The Commission has thus far defined the following separate markets in the pharmaceutical industry distribution value chain:

1. pre-wholesale services;
2. wholesale services;

52 See Ibid., at paragraph 122; Case COMP/M.5865 – Teva/Ratiopharm, at paragraph 408.
54 Ibid, at paragraphs 120 and 121.
55 See Case IV/26.911 – Zoja/CSC-ICI.
56 See Case COMP/M.6044 – Alliance Boots/Andreae-Noris Zahn, at paragraphs 6 ff.
57 See Case COMP/M.4301 – Alliance Boots/Cardinal Health, at paragraphs 10 ff. This activity has been distinguished from the direct distribution of manufacturers to customers (e.g., retail pharmacies, hospitals); see Case COMP/M.5865 – Teva/Ratiopharm, at paragraphs 450–452.
III GEOGRAPHIC MARKET DEFINITION

i Finished pharmaceuticals
The Commission has consistently found geographic markets for finished pharmaceutical products to be national. However, when the Commission has taken into consideration the future presence of a particular pharmaceutical company in a specific market by reference to its pipeline products, R&D or patents, the Commission has tended to find wider geographic markets (EEA-wide or worldwide). This is due to the fact that R&D tends to occur on a multinational and, often, global scale.61

ii Markets upstream of finished pharmaceuticals
Upstream markets have generally been found to be at least EEA-wide or worldwide in scope, regardless of whether they concern the sale of APIs,62 contract manufacturing63 or out-licensing.64

IV COMPETITIVE ANALYSIS IN THE PHARMACEUTICAL SECTOR

i Preliminary considerations
Whether the Commission considers that a transaction in the pharmaceutical industry raises competition concerns is likely to depend on the nature of the business activities and nature companies involved in the concentration. For example, mergers between two originators active in the same markets may raise traditional horizontal concerns. Mergers between originators and research firms might affect competition in the current and future relevant product markets, particularly if both companies have competing late-stage pipeline products or the transaction could result in a decrease in overall R&D.

58 See Case COMP/M.7323 – Nordic Capital/Ghd Verwaltung, at paragraph 39; Case COMP/M.5805 – 3i/Vedici Groupe; Case COMP/M.5548 – Barclays/RBS/Hillary; Case COMP/M. 4367 – APW/APS/Al Nordic Capital/Capio; Case COMP/M.4229 – APHL/L&I/Netcare General Healthcare Group; Case COMP/M.4788 – Rozier/BHS.
59 See Case COMP/M.2432 – Angelini/Phoenix/JV; Case COMP/M.2573 – A&G/Grospharma, at paragraphs 11 and 12; and Case COMP/M.5865 – Teva/Ratiopharm, at paragraph 452.
60 See Case COMP/M.7323 – Nordic Capital/Ghd Verwaltung.
61 For future markets, see Case COMP/M.5865 – Teva/Ratiopharm, at paragraph 422.
62 Ibid, at paragraph 396.
64 See Case COMP/M.6613 – Watson/Actavis, at paragraphs 120 and 121; Case COMP/M.5865 – Teva/Ratiopharm, at paragraph 396; Case COMP/M.5295 – Teva/Barr, at paragraph 190; and COMP/M.6278 – Teva/Ratiopharm, at paragraph 19.
Different competition concerns may arise when a concentration involves one or more generic producers. Generic companies not only compete among each other in the development of biosimilar and molecule generics of originator pharmaceuticals; they also compete with originator pharmaceuticals after the expiration of the relevant patent.

**ii  Key competitive drivers in the pharmaceutical markets**

To date, competition concerns in notified cases have focused on potential restrictions arising from direct overlaps in the relevant market or markets. In addition, the competitive assessment is likely to take into consideration innovation and other aspects of dynamic competition, the effects of regulation and reimbursement schemes on competition and the commercialisation stage of the relevant products. The Commission has focused its analysis on the different competitive drivers of the pharmaceutical markets.

**Innovation and product differentiation**

The Commission tends to consider innovation to be of critical importance for the pharmaceutical sector. In the Commission Communication accompanying the Pharmaceutical Sector Inquiry, the Commission highlighted the significant R&D efforts of originator companies and other stakeholders (e.g., research companies) in order to innovate. Current Competition Commissioner Margrethe Vestager seems to have embraced this by indicating that the Commission’s ‘focus on investment and innovation in merger control is also clear in the pharmaceutical sector’, although when pharmaceutical companies announce a merger the Commission needs to ‘carefully balance the benefits of pooling their resources with the potential negative impact of eliminating an innovator’.

In a number of cases, the Commission considered the potential risks that a concentration entails for the development of pharmaceuticals and innovation, both in relation to originators (e.g., when developing new pharmaceutical products or variants) and to generic manufacturers (e.g., when developing biosimilars). Similarly, the Commission has also taken into account the potential impact of competition exerted by ‘pipeline’ products (i.e., the competition that products might face from other products not yet released on the market) when reviewing certain transactions. This is due to the fact that, if the adequate conditions are met (e.g., those regarding the pharmacological characteristics and therapeutic use), pipeline products might be actual or credible future competitors of existing products. In assessing pipeline competition, the Commission has focused on instances where one merging

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67 See, e.g., Case COMP/M.5661 – Abbott/Solvay Pharmaceuticals, at paragraph 119; Case COMP/M.5502 – Merck/Schering-Plough, at paragraph 38.
69 See Case COMP/M.5502 – Merck/Schering-Plough, at paragraph 38.
party is planning to enter a market with a new product within a period of two years, and the merging other party (or the parties combined) has a market share of 35 per cent or more on any possible market definition where the pipeline products and existing products overlap.\(^70\)

Intellectual property rights (IPRs) are also a key element in the promotion of innovation. The pharmaceutical industry invests heavily in R&D and tends to rely on IPRs to protect innovation, and to manufacture or distribute its products, or both (e.g., through out-licensing). In recent cases, the Commission analysed the potential effects of transactions on R&D, for example by decreasing the merging parties’ incentives to further investigate, or by obstructing the licensing of patents for R&D.\(^71\) Such an exercise presents the potential pitfall, from an analytical perspective, of the different approaches of economists to the relationship between market structure and innovation.\(^72\)

As occurs in other markets, product differentiation can also play an important role in determining the competitiveness or closeness of competition between different pharmaceuticals. For example, originators might enjoy a better position in the market as a result of the publication of clinical trial evidence (e.g., as regards the efficacy or safety profile of a product) or where their products are well known to end users (e.g., as a result of branding advertising).

**Authorisation, price and reimbursement conditions**

The Commission sometimes finds that pharmaceutical markets present rigid pricing and entry conditions.\(^73\)

The sale of prescription medicines is generally regulated in the EU and is often subject to reimbursement conditions from the social security systems of the Member States. As a result, the use of prescription medicines may rely heavily upon the national authorisations and guidelines that doctors and medical staff use for their prescription; and upon the extent to which the financial burden is ultimately borne by the affected Member State (or private health insurances, or both) in question.

As previously indicated, the impact that authorisation procedures, pricing and reimbursement conditions have on consumption levels for certain pharmaceutical products is such that these elements have been used in a number of Commission precedents to exclude products from the relevant market despite therapeutic indications being identical.\(^74\) In addition, price regulation and reimbursement schemes in the Member States are likely to be of crucial importance when analysing the ability of a merged entity to increase prices

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71 See Case COMP/M.7275 – *Novartis/GlaxoSmithKline Oncology Business*, at paragraph 105; Case COMP/M.5476 – *Pfizer/Wyeth*, at paragraphs 91 and 93.


73 See, e.g., Case COMP/M.7275 – *Novartis/GlaxoSmithKline Oncology Business*, at paragraph 62.

74 Ibid, at paragraphs 123–124; Case COMP/M.5865 – *Teva/Rationpharm*, at paragraph 90.
EU Merger Control in the Pharmaceutical Sector

post-transaction in markets for prescription medicines.\(^{75}\) Reimbursement schemes that encourage generic competition may also be taken into account when determining whether a transaction leads to competition concerns.\(^{76}\)

**Generic and biosimilar competition**

The existence of competitive constraints from generic manufacturers can be of great importance in determining whether a concentration will give rise to competition concerns. Broadly speaking, after the expiry of the relevant patent, generic pharmaceutical companies are in competition with one another and with originator companies.\(^{77}\) In general, generic companies focus on price competition and tend to invest less in branding and advertising, given the limited importance attributed by them to differentiating their products from originator products. In addition, in a number of countries, regulatory substitution rules and mechanisms will ensure generic market penetration with limited promotional activities being needed. Finally, in a majority of countries in the EU, the entry of a generic product may trigger price reductions, as a result of which demand tends to shift away from originators.

In assessing competition between originator and generic companies, the Commission considers, *inter alia*, the asymmetries in their respective product offerings or market focuses to assess whether the two products are closely competing.\(^{78}\)

**Competitive dynamics for OTC products**

OTC medicines are subject to different competitive dynamics than those identified for prescription medicines. Indeed, the sale of OTC products is usually significantly less subject to reimbursement regulations or to the prescription guidance of doctors, which shifts the decision-making role to pharmacists and end users. As a result, the success of OTC medicines tends to rely on (consumer-focused) advertising, innovation (often in the form of customisation to suit user preferences) and branding strategies. Generic OTC medicines sometimes exert significant competitive constraints on branded OTC medicines, although the ultimate impact of such constraints will ultimately depend on the brands and products in question.\(^{79}\)

**Non-horizontal effects**

Concentrations between pharmaceutical companies can lead to competition concerns where the parties are active in markets that are upstream, downstream or adjacent of one another.

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\(^{75}\) Ibid, at paragraph 65, where a number of factors were cited in support of the view that prices in all the affected countries for many medicines in mature generic markets were upwardly inflexible or could only be increased with difficulty.

\(^{76}\) See, e.g., Case COMP/M.5295 – Teva/Barr, at paragraphs 186 ff.

\(^{77}\) Ibid, at paragraphs 184–185.

\(^{78}\) See Case COMP/M.5253 – Sanofi-Aventis/Zentiva, at paragraph 484, regarding generic and originator differentiation; Case COMP/M.6258 – Teva/Cephalon, at paragraph 12, regarding generic differentiation.

\(^{79}\) See Case COMP/M.6280 – P&G/Teva OTC Business, at paragraph 20; Case COMP/M.3751 – Novartis/Hexal.
For example, based on Commission’s practice, out-licensing may result in vertically affected markets where: (1) one merging party (company A) is active on a downstream market for the manufacturing of a specific finished dose pharmaceutical; (2) the other merging party (company B) is active upstream as a licensor and contractor of another manufacturer downstream (company C); and (3) the combined market share of the merging parties and of the sub-contracted licensee (companies A, B, and C) on the downstream market exceeds 25 per cent.

As with other non-horizontal mergers, the importance of this type of concerns will depend on the market presence and power of the merging parties in the relevant upstream and downstream markets, and their ability and incentive to leverage this presence into other markets to foreclose competing companies.

iii Remedies and commitments in merger control

The history of EU merger control in the pharmaceutical sector is characterised by the absence of prohibition decisions. However, the Commission has accepted commitments proposed by the merging parties primarily during Phase I investigations. As regards Phase II investigations, cases have been completed with commitments decisions mainly in the related but separate medical devices industry.

Where the Commission has identified concerns that result from the horizontal overlap created by a concentration, it has traditionally required the divestiture of entire product lines or businesses, or both, to eliminate a significant part of the parties’ overlap in a problematic (geographic or product) market. The appropriateness and suitability of the assets, IPRs, licences, supplies, personnel, customer lists or proposed buyer will be key in determining the acceptance of any proposed divestitures, in particular where the divested business risks are not being considered as a stand-alone entity by the Commission but rather as part of a larger business that, when severed from the entity, could have limited viability, competitiveness and ability to innovate post-transaction. Where necessary, the Commission might require

80 For vertical concerns see, e.g., Case COMP/M.6258 – Teva/Cephalon, at paragraphs 133 ff and Case COMP/M.3493 – Yamanouchi/Fujisawa. For conglomerate concerns see, e.g., COMP/M.5999 – Sanofi-Aventis/Genzyme, at Section 4.6.
81 See, e.g., Case COMP/M.5253 – Sanofi-Aventis/Zentiva, at paragraphs 511 ff; Case COMP/M.6258 – Teva/Cephalon, at paragraphs 133 ff.
82 See, e.g., Case COMP/M.3687 – Johnson & Johnson/Guidant; Case COMP/M.6266 – Johnson & Johnson/Synthes; and Case COMP/M.7265 – Zimmer/Biomet.
83 See, e.g., Case COMP/M.3544 – Bayer Healthcare/Roche (OTC Business), at paragraphs 57 ff, regarding divestitures of product lines; Case COMP/M.3751 – Novartis/Hexal, at Section 6, regarding divestitures of sales and marketing rights; Case COMP/M.4314 – Johnson & Johnson/Pfizer Consumer Healthcare, at paragraphs 138 ff, regarding the divestiture of assets (e.g., inventories, clinical data, trademarks); Case COMP/M.4779 – Akzo/ICI, at paragraphs 53 ff, regarding the divestiture of a shareholding in a joint venture. See also Case COMP/M.5253 – Sanofi-Aventis/Zentiva, at paragraphs 550 ff; Case COMP/M.5295 – Teva/Barr, at paragraphs 205 ff; Case COMP/M.7379 – Mylan/Abbott EPD-DM, at Section 5.
84 See, e.g., Case COMP/M.6851 – Baxter International/Gambro, at paragraph 564 and 571 (where the merging parties proposed an up-front buyer). For the need to ensure supplies and
that the notifying party enter into agreements with the divestiture buyer prior to clearance in order to ensure that the latter will be committed to carrying out R&D investments and launches of pipeline products.\textsuperscript{85}

Given the complexity of the pharmaceutical sector, the Commission has sometimes required that the divestiture of product lines or businesses be supported with the provision of technical assistance in the production, sale and marketing of the pharmaceutical product.\textsuperscript{86} Sometimes, to facilitate the market entry and sustained competitiveness of third parties, the Commission has required the divestiture of a product line to be supplemented with the divestiture of a distribution business.\textsuperscript{87}

Insofar as the merger control review of the Commission is limited to the EEA market or markets where it identifies competition concerns, the Commission has accepted where appropriate divestitures of global businesses combined with an exclusive license-back clause to the merging parties for non-EEA markets.\textsuperscript{88} In such cases, where the notifying party also reserves for itself certain decision-making rights related to the output of the divested business (e.g., product development, clinical trials), these should be limited so as to not to give to it joint control on these aspects of the business.\textsuperscript{89}

In other circumstances, the Commission has accepted licensing arrangements as an alternative to divestitures where, for instance, the proposed divestiture would hinder ongoing research or it would be impossible due to the nature of the business.\textsuperscript{90}

In addition, to the extent that the competitive concerns related to the grant of exclusive rights or licences to other competing entities, the Commission has accepted commitments to, dilute or remove altogether minority shareholding relationships and other contractual arrangements (e.g., turn exclusive licensing relationships into non-exclusive relationships, limiting supply agreements).

V CONCLUSIONS

While the substantive analysis of mergers in the pharmaceutical sector might not be fundamentally different from that carried out in other innovation-intensive regulated industries, it presents the following particularities:

\begin{itemize}
  \item[a] the market definition process is characterised by the use of a range of different analytical tools to identify relevant markets. The Commission has a well-established
\end{itemize}

\footnotesize
\begin{itemize}
  \item licences in the medical devices sector, see Case COMP/M.7326 – Medtronic/Covidien and Case COMP/M.7265 – Zimmer/Biomet; Case COMP/M.7379 – Mylan/Abbott EPD-DM, at paragraphs 472 ff. and 485.
  \item See Case COMP/M.7275 – Novartis/GlaxoSmithKline Oncology Business.
  \item See, e.g., Case COMP/M.4314 – Johnson & Johnson/Pfizer Consumer Healthcare, at paragraphs 139 and 140.
  \item See Case COMP/M.5778 – Novartis /Alcon, at paragraphs 291 ff.
  \item See Case COMP/M.7559 – Pfizer/Hospira, at paragraph 297.
  \item See Case COMP/M.7559 – Pfizer/Hospira, at paragraph 310.
\end{itemize}
approach to market definition on the basis of the EPhMRA ATC classification, nuanced by reference to other competitive drivers such as the distinction between OTC and prescription drugs and, where applicable, between originators and generics; actual competition is analysed mainly on the basis of observed overlaps in the relevant market or markets. In addition, the competitive assessment becomes more complex as the Commission takes into consideration innovation and other aspects of dynamic competition, the impact of regulation and reimbursement schemes, and the commercialisation stage of the relevant products; and where concerns arise deriving from horizontal overlaps, the Commission is not reluctant to require divestitures, emphasising the need for the divested business to constitute a viable stand-alone business.
Appendix 1

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