Sales Growth of New Pharmaceuticals Across the Globe: The Role of Regulatory Regimes

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Prior marketing literature has overlooked the role of regulatory regimes in explaining international sales growth of new products. This paper addresses this gap in the context of new pharmaceuticals (15 new molecules in 34 countries) and sheds light on the effects of regulatory regimes on new drug sales across the globe.

Based on a time-varying coefficient model, we find that differences in regulation substantially contribute to cross-country variation in sales. One of the regulatory constraints investigated, i.e., manufacturer price controls, has a positive effect on drug sales. The other forms of regulation such as restrictions of physician prescription budgets and the prohibition of direct-to-consumer advertising (DTCA) tend to hurt sales. The effect of manufacturer price controls is similar for newly launched and mature drugs. By contrast, regulations on physician prescription budgets and DTCA have a differential effect for newly launched and mature drugs. Whereas the former hurts mature drugs more, the latter has a larger effect on newly launched drugs. In addition to these regulatory effects, we find that national culture, economic wealth, and lagged sales also affect drug sales.

Our findings may be used as input by managers for international launch and marketing decisions. They may also be used by public policy administrators to assess the role of regulatory regimes in pharmaceutical sales growth.

Keywords: international new product growth; drug; pharmaceutical; regulation; culture; economics; time-varying effects; penalized splines

Introduction
Over the past decades, marketing research on international new product growth has identified several key drivers of variation across countries. Among them, the most prominently studied are economic and socio-cultural characteristics of nations (Dekimpe et al. 1998, Gatignon et al. 1989, Helsen et al. 1993, Stremersch and Tellis 2004, Talukdar et al. 2002, Tellis et al. 2003, Van den Bulte and Stremersch 2004). Cross-country interrelations have also received special attention, either under the umbrella of cross-national learning, lead-lag, or introductory-lag effects (see Dekimpe et al. 2000 for an extensive review). However, the role of regulation on new product growth has been mostly overlooked.

This lack of research is particularly surprising as the variety of regulatory systems across the globe was among the prime motivations invoked for international new product growth models (Heeler and Hustad 1980, Mahajan and Muller 1994), and recent calls have been made for more research in this area (e.g., Stremersch and Tellis 2004). Government regulation is an interesting and complex topic of study with many societal implications (Shugan 2003). Industries in which regulation affects firms’ marketing strategies and sales include telecommunications, energy, tobacco, liquor, and financial services. Many countries have regulated the marketing of products in these industries by means of advertising regulation for tobacco and financial services, price regulation for telecommunications, and profit regulations for energy.

In this paper, we focus on regulatory regimes in the pharmaceutical industry as there is increased evidence of its great influence, both from academia (Atun and Gurol-Urganci 2006, Berndt et al. 2007, Chintagunta and Desiraju 2005, Danzon and Ketcham 2004, Kyle 2007, Kolsarici and Vakratsas 2008) and practice (European Commission 2004, Urch 2005). Many different regulatory regimes exist and—while the business press routinely refers to their importance (e.g., The Economist 2007)—their effects on sales are not straightforward and largely unexamined. Therefore, pharmaceuticals are an appropriate context...
to examine the role regulation may play in international sales growth.

Our aim is to model market response to regulation in a nonstationary environment, the new product growth context. Therefore, we consider that regulatory regimes may have a differential effect on drugs that were introduced very recently (e.g., a few months ago)—we refer to such drugs as newly launched drugs—than on maturing drugs that have been on the market for a longer period of time. For instance, regulation that caps the number of prescriptions physicians can write may not influence sales of newly launched drugs that much, while it may seriously depress sales when the drug matures. As we will explain, the reasons may be that, in the first months after launch, the volume the physician prescribes is still low and is predominantly used by the patients that benefit most from the new drug, e.g., because alternative treatments for these patients were not effective or showed major side effects. Aside from regulation, prior research on international new product growth also contains some early evidence that other country characteristics, such as national culture and economic wealth, affect new product growth differentially across the life cycle (e.g., Stremersch and Tellis 2004, Tellis et al. 2003). Hence, ignoring potential variation over time in the effects of country characteristics, such as regulation, national culture, or economic wealth on sales may yield misleading or incomplete insights.

To analyze cross-country variation in new drug sales, we obtained monthly sales data for 15 new molecules1 across four therapeutic categories and 34 countries, during their first 84 months (7 years) after launch. To explain the observed international differences, we gather data from many other sources on regulation, culture, and economics. Our data include many developing nations in Eastern Europe, Africa, South America, and the Middle East. This provides us with a strong empirical basis, although constrained to the context of international sales growth of new pharmaceutical drugs.

The remainder of the paper is organized as follows. In the next section, we discuss the effects of countries’ regulatory environments for pharmaceuticals in more detail. In the subsequent sections, we present our data, the modeling framework used, and our results. Finally, we end with a discussion and suggestions for future research.

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1 A molecule is the active substance of a drug. As several competing brands contain the same molecule, we focus on the sales growth pattern of the molecule rather than these brands. This choice is consistent with the literature on new product growth that generally models sales or adoption of a product (e.g., TV), irrespective of the brand (e.g., Sony).

2 QALY is a measure of the beneficial effects of a medical treatment. It is based on the number of life years added as a result of the medical treatment. Each year of perfect health (e.g., because of the loss of a limb, lack of mobility, etc.) is assigned a value between zero and one.
often take the cost of the treatment into account in their joint decision with the patient (e.g., Hart et al. 1997). Therefore, we expect price regulation to positively affect sales.

However, such effect may be limited for newly launched molecules, as compared to maturing molecules. Such expectation comes from prior literature that examined the link between price regulation and pricing patterns over the new drug’s life cycle. This literature has shown that prices fall faster when prices are regulated than when prices are not regulated (Danzon and Chao 2000, Ekelund and Persson 2003).

Interestingly, we also obtained anecdotal evidence for the pricing pattern we hypothesize above for a subset of the countries in our sample. Figure 1 shows the typical pricing pattern, together with 95% confidence bounds of new drugs in three countries with price control (Belgium, Switzerland, and Canada) and three countries without price control (Germany, the United Kingdom, and the United States), over the data window (the Technical Appendix can be found at http://mktsci.pubs.informs.org). The values on the y axis represent the average drug price pattern across all drugs in these six countries in the presence or absence of regulation, expressed in dollars per gram. Figure 1 shows that prices in countries with price regulation decrease faster than prices in countries without price regulation, reaching substantially lower levels for maturing drugs. The upward slope in the nonregulated countries is likely because of inflation.

For all the reasons posited, we may expect that, while manufacturer price regulation may increase sales of maturing drugs, it would in turn have little or no effect on sales of recently introduced drugs.

**Regulation on Physician Prescription Budgets**

The most direct way in which regulators can intervene in pharmaceutical markets for prescription drugs is to limit the total number of prescriptions a physician can write. For instance, Germany in the early 1990s introduced a collective budget for prescription drugs that was later reformed to physician-level prescription budgets. Physicians in Latvia are assigned a fixed budget calculated on the number of patients in their practice. Assigning budgets to physicians for the total number of drugs prescribed may restrict prescription drug sales. Thus, countries that impose such restrictions are likely to show lower drug sales than countries that do not.

Typically, physicians adopt a stepwise approach in prescriptions (Prosser et al. 2003). They try familiar alternatives first, and only prescribe new drugs if the familiar alternatives fail or cause adverse effects. Prosser and Walley (2003) find that physicians typically prescribe newly introduced drugs to patients who do not show significant improvement in their (mostly severe) condition when using prior alternatives or who do not tolerate such alternatives because of side effects. Therefore, the effect of prescription budget regulation on early sales of a newly launched drug may be limited. However, the total market potential for the drug may be smaller in the presence of prescription budget regulation. In regulated markets, physicians will feel more pressure not to prescribe the drug to patients with mostly mild symptoms or who tolerate traditional alternatives well, as compared to nonregulated markets.

**Patient Co-Payment Regulation**

In many countries, patients are required to provide some form of co-payment for prescription drugs. For instance, Belgian patients typically pay for only a small part of their prescription medicine; the rest is paid by a government institution, and the patient contribution is set at the level of the individual drug (again taking into account QALY). Hungary has a similar system (although the co-payment amounts differ), but also adds patient income to the reimbursement level, with higher reimbursements (in many cases 100%) for the poor. In other countries, patients do not co-pay for drugs. For instance, in The Netherlands, the cost of prescription drugs and physician visits is directly settled between physician, pharmacist, and the patient’s insurance company (which can be private or public). In co-payment systems, patients are monetarily sanctioned for drug use and

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3 We obtained anecdotal evidence on pricing and marketing efforts for all drugs in our sample, but for a limited set of countries—Belgium, Canada, Germany, Switzerland, the United Kingdom, and the United States—for which our data supplier had complete information.
thus experience a part of the cost of their treatment directly (Reuveni et al. 2002). Such monetary sanctioning is expected to depress drug sales, given that it makes treatment cost a more salient criterion in the physician-patient joint decision (e.g., Hart et al. 1997).

A policy initiative in Italy provides initial face validity for the above reasoning. Italy abolished its co-payment system at the end of 2000. In the first few months of 2001, pharmaceutical spending in Italy rose by approximately 30% (while the federation of Italian family practitioners assessed the total increase to be around 2%–3%). Soon after, co-payments were reintroduced by the Italian government (Urch 2005). Cross-country variation in drug sales, driven by the absence or presence of a co-payment system, has not been investigated on such a large scale as in the present paper.

There is no reason to expect the presence of a co-payment system to affect newly launched drugs differently than mature drugs, as the level of co-pay in countries may not depend on the drug’s age, or even on its price. For example, in Austria, patients pay a flat fee of €4.45 per prescription. In Portugal, patients pay a fixed percentage of the drug cost according to the essential nature of the drug (specified at the category level).

**Regulation of Marketing Efforts to Physicians**

Marketing activities to physicians represent a large part of the expenditures in the pharmaceutical industry. Mizik and Jacobson (2004) indicate that U.S. companies have spent over $5.8 billion in 2002 on detailing, combined with another $11.5 billion on free sampling. In view of these large costs, scholars have paid a lot of attention to the effects of marketing efforts to physicians on prescription behavior and pharmaceutical sales (Chintagunta and Desiraju 2005; Dekimpe and Hanssens 1999; Ding and Eliashberg 2008; Hahn et al. 1994; Leeflang et al. 2005; Manchanda and Chintagunta 2004; Manchanda and Honka 2005; Manchanda et al. 2004; Mantrala et al. 1994; Narayanan and Manchanda 2008; Narayanan et al. 2004, 2005; Venkataraman and Stremersch 2007). There has been some debate on the direction of the effect of marketing efforts on prescription behavior. Some authors find positive effects (e.g., Gönül et al. 2001, Venkataraman and Stremersch 2007), others find neutral effects (e.g., Rosenthal et al. 2003), and still others find negative effects (e.g., Parsons and Vanden Abeele 1981). Also, the size of these effects has been under debate; some find strong effects (e.g., Gönül et al. 2001), while others find more modest effects (e.g., Mizik and Jacobson 2004). Nonetheless, the overall conclusion is that marketing efforts often have a positive and significant effect on physician decision making. Patients are typically not aware of this influence on the prescription decision.

The marketing efforts that prior literature has considered are as follows: detailing, meetings, and sampling (see Narayanan and Manchanda 2006, Venkataraman and Stremersch 2007). Regulation may constrain the number of detailing visits a manufacturer can make, the number of meetings a manufacturer can organize, or the number of samples a manufacturer can dispense to physicians. For instance, the amount manufacturers spend in the United Kingdom on sales promotions is limited to a proportion of their overall (not drug-specific) profits (the “marketing allowance”). In Spain, promotional costs are capped at a percentage of sales of the firm. Given the expected positive effect of marketing efforts on new drug sales, we expect that countries that limit the marketing efforts of a pharmaceutical firm to physicians would show lower drug sales than countries that do not impose such limits.

One may counter that if the cap is quite high, it may not affect marketing efforts at all and therefore not affect drug sales. However, this argument does not fit the regulatory practices of countries. First, governments implement these caps on marketing efforts to limit prescriptions (see Urch 2005). Second, just from the specific regulations, one can appreciate the low level at which marketing efforts are capped. For instance, in Poland, manufacturers can only leave, at maximum, three samples of the smallest package at a physician’s office over an entire year. This approach contrasts sharply with the United States, where it is common to leave approximately 10 samples at a physician’s office per year (Source: IMS Health, Category for Antiarthritic Treatments (Cox-2)).

Anecdotal evidence we obtained for a subset of the countries confirms the negative influence of such regulation on marketing efforts. Similar to Figure 1, we report the typical pattern of marketing efforts for new drugs, together with 95% confidence bounds, in two countries with restrictions on marketing efforts (Belgium and the United Kingdom) and three countries without such restrictions (Germany, Switzerland, and the United States) in Figure 2. The values on the y axis represent the average pattern in marketing efforts across all drugs in these five countries in the presence or absence of regulation, expressed in thousands of dollars. It shows that marketing efforts are lower in countries with restrictions on marketing efforts than in countries without such restrictions.

In addition, one may expect that the restriction of marketing efforts to physicians has a larger effect on newly launched drugs than on maturing drugs. The literature on advertising has shown that sales-to-advertising elasticity decreases over a product’s life cycle (Chandy et al. 2001, Tellis and Fornell 1988), because consumers are more motivated to gather and analyze information about new products than
older ones (Grunert 1996). The same very likely applies to physicians, as Neslin (2001) and Narayanan et al. (2005) show that detailing is most effective in the introduction phase. Consequently, a regulatory restriction is likely to depress sales of newly launched drugs more than sales of maturing drugs. A potential counterargument may be that firms could increase their marketing efforts to physicians as the drug matures. However, data on the pattern of communication efforts over time generally do not support this argument; rather, they support the contrary (for examples, see Horsky and Simon 1983, Lilien and Little 1976). In the pharmaceutical industry, Narayanan et al. (2005) show that market efforts to physicians for antihistamines increase for Allegra, decrease for Zyrtec, and remain constant for Claritin. Also the anecdotal evidence in Figure 2 shows stable marketing efforts over time in regulated countries.

Regulation of DTCA
While marketing to physicians is a direct way to affect physicians’ prescription behavior, pharmaceutical firms can also attempt to influence patients by advertising the drug directly to consumers. So far, only two countries, New Zealand and the United States, allow pharmaceutical marketers to advertise directly to consumers. The practice is more novel than marketing efforts to physicians. Yet it is quickly becoming very popular among marketers. Spending on DTCA in the United States, for example, has increased from $1.1 billion in 1997 to $5.6 billion in 2006, according to IMS Health.

Prior literature has shown that DTCA triggers patients to request drugs by their brand name from their physicians (Cohen 1988, Mehta and Purvis 2003, Mintzes et al. 2003, Rosenthal et al. 2003, Weissman et al. 2004, West 1999, Wosinska 2002). Such patient requests are often accommodated by physicians mainly because they affect physician visit satisfaction (Kravitz et al. 2003) and show patients that the physician cares (Schwartz et al. 1989). For instance, Bell et al. (1999) found that 46% of patients thought they would be disappointed if a physician did not prescribe an advertised drug they requested, while 24% thought they might attempt to obtain the prescription from a different doctor, and 15% thought they would switch to a new doctor.

Given the expected positive effect of DTCA on drug sales, we expect that countries that forbid DTCA show lower drug sales than countries, such as New Zealand and the United States, that do not impose such a restriction. We expect this effect to become smaller as a drug matures, for reasons similar to those developed on restrictions on marketing efforts to physicians. More precisely, we expect higher DTCA effectiveness for new drugs than for maturing drugs, in line with prior results by Neslin (2001) and consistent with the argument above about consumers’ higher motivation to gather and analyze information about new products than maturing ones. Moreover, DTCA is generally high for newly launched drugs and declines as the drug matures (Rosenthal et al. 2002).

Other Variables
When examining the role of regulation on the growth of new pharmaceuticals, we need to control for other factors that may explain cross-national variation in drug sales. Building our research on existing international new product growth literature, we subsequently consider the roles of national culture (Stremersch and Tellis 2004, Tellis et al. 2003, Van den Bulte and Stremersch 2004), economics (Talukdar et al. 2002, Van den Bulte and Stremersch 2004), introduction lag (Dekimpe et al. 2000, Kumar and Krishnan 2002, Talukdar et al. 2002), lagged sales (Desiraju et al. 2004), and competition (Chintagunta and Desiraju 2005, Desiraju et al. 2004) in the sales growth of new pharmaceuticals.

National Culture
Geert Hofstede⁴ (Hofstede 1980, 2001) identified four main dimensions of national culture that can be related to new drug sales: (1) uncertainty avoidance; (2) individualism; (3) masculinity; and (4) long-term

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⁴ An alternative operationalization is Schwartz’s (1992) model of personal values. Hofstede’s national culture framework is most often adopted by scientists, including in marketing. Obviously, our work can easily be extended to test for the effects of Schwartz’s model of personal values using the same methodology.
orientation.\textsuperscript{5} Hofstede’s national culture dimensions are widely accepted and frequently used for cross-country comparisons (e.g., Dawar and Parker 1994, Lynn et al. 1993, Roth 1995, Steenkamp et al. 1999).

Uncertainty avoidance refers to the extent to which the population of a country feels (intolerable) anxiety when facing uncertain or unknown situations (Hofstede 2001). Patients in uncertainty avoidant cultures show lower subjective health (Hofstede 2001). Lower subjective health perceptions may lead to higher demand for pharmaceutical drugs, considering that physicians typically accommodate patients’ requests for treatment (Bell et al. 1999). However, the positive effect of uncertainty avoidance on drug sales may only materialize for maturing drugs and have little or no effect on newly launched drugs. Indeed, new products are usually perceived as more risky, and their performance more ambiguous than established products and brands (Steenkamp et al. 1999). In cultures of high uncertainty avoidance, consumers are more likely to identify uncertainty in new products, compared to consumers in low uncertainty avoidance cultures (Michaut 2004). Moreover, societies high in uncertainty avoidance consider novel ideas dangerous and are more intolerant toward change than societies low in uncertainty avoidance (Hofstede 1980). This may temper the positive effect of uncertainty avoidance on sales of newly launched drugs.

Individualism describes the relationship between the individual and the collectivity that prevails in a given society and the extent to which people prefer to act as individuals rather than as members of a social group (Hofstede 2001). Individualism may have several effects on drug sales. In general, individualist cultures tend to spend more of their public and private budgets on health care (Hofstede 2001), which fits with the emphasis that such cultures put on individual well-being. By contrast, individual well-being in collectivist cultures is only relevant in relation to the collective well-being. Higher spending on health care will lead to higher drug sales. Patients in individualist societies also seek doctor consultation more actively than patients in collectivist societies, which, in turn, also results in higher drug sales. While collectivist cultures have on average lower drug usage, the difference may be less pronounced for newly launched drugs. Indeed, as already stated, newly launched drugs are typically prescribed to patients with a strong need for new therapeutical treatments, either

Economic Wealth
Economic wealth has often been advanced as the major economic characteristic of nations that affects new product sales growth. The economic wealth of nations, most commonly operationalized by gross domestic product (GDP) per capita, may have several positive effects on sales of pharmaceuticals. On the one hand, the population of more wealthy countries can more easily afford drugs than the population of less wealthy countries (Desiraju et al. 2004,

\textsuperscript{5}We found neither theoretical support nor empirical evidence in our data to include power distance as an explanatory variable. While power distance may be related to the structure of the health system (for instance, one may imagine medical doctors and pharmacists taking a more hierarchical position towards their patients in power distant cultures), there seems to be no argument to relate it to new drug sales.
Stremersch and Tellis 2004, Talukdar et al. 2002, Tellis et al. 2003, Van den Bulte and Stremersch 2004. On the other hand, economic wealth is also associated with better health infrastructure, which in turn may positively affect the sales of drugs (Desiraju et al. 2004). In our view, these arguments are likely to play similarly for newly launched and maturing drugs.

Cross-Country Introductory Lag
Cross-country introduction delays refer to the time elapsed between the first introduction of a new product in any country of the world and its introduction in a focal country (Talukdar et al. 2002). The uptake of new molecules may be faster in laggard countries than in lead countries because patients and physicians learn from experiences in lead countries (Dekimpe et al. 2000, Desiraju et al. 2004, Takada and Jain 1991). These cross-country “learning effects” have been found to generalize to multiple types of decision makers, including consumers, firms, and governments (Dekimpe et al. 2000). However, previous work also suggests that firms are likely to delay entry in countries that they judge “less attractive” than others. Factors influencing such a delay may be regulation (e.g., Kyle 2007) or expected sales. As such, introduction lag is not exogenous to our model. As further explained in the Model section, we solve this issue by using an instrumental variable procedure.

Cross-country learning effects are likely to vary over time. The existence of cross-country learning effects can be explained by the role they play in reducing the uncertainty associated with the adoption of new products. As new products are generally associated with more uncertainty than established products and brands (Steenkamp et al. 1999), we expect that the role of cross-country learning effects will be more pronounced for newly launched than for maturing drugs.

Competition
Although the presence of competitive firms and brands has often been neglected in the new product growth literature, it is now largely recognized as a major influence on new product sales (Chintagunta and Desiraju 2005, Mahajan et al. 1993, Mason 1990; see Chatterjee et al. 2000 for a review). To account for the role of word-of-mouth communication and the substitution dynamics between competing brands, Mahajan et al. (1993) proposed a diffusion modeling approach assessing the impact of competitive entry on market size and the sales of incumbent firms. Recently, Krishnan et al. (2000) found that the entry of a competing brand in the cellular telephone industry can affect the total market potential of the category and/or the speed of diffusion in the category depending on the market studied. In the pharmaceutical industry, early evidence for the antidepressant category (Desiraju et al. 2004) suggests that competition may not have a strong influence. Given these mixed findings, expectations as to the direction of this effect, as well as to its variation over time are difficult to conceive.

Lagged Sales
There are three reasons why lagged sales of a new drug may affect its present sales. First, once a physician starts prescribing a new drug, he may trigger adoption by other physicians, an effect commonly known as contagion. Second, physicians show considerable inertia in prescription behavior (Janakiraman et al. 2008, Venkatakrishnan and Stremersch 2007). Thus, physicians who heavily prescribe a certain drug today are likely to maintain similar behavior in the future. Third, in the case of chronic diseases, patients generally stay on the same drug for a longer time, receiving refill prescriptions repeatedly. All three effects will create duration dependence in sales, either coming from influence across physicians, persistence over time within a physician’s prescription behavior, or persistence over time within a patient’s treatment behavior.

Contagion among physicians and inertia in physicians’ prescription behaviors typically affect already established drugs, rather than newly launched drugs. Thus, we expect the effect of lagged sales to increase over the new drugs’ life cycle.

We provide an overview of all our expectations in Table 1. In columns 2 and 3, we provide the definition of the variables. In columns 4 and 5, we specify the effects expected for each covariate studied. In particular, column 4 specifies the main effect, while column 5 specifies how the effect is likely to change over time as the drug matures. We include our empirical findings in columns 6 and 7, and assess whether our expectations are confirmed in the last column.

Data
We studied the sales growth process of 15 new molecules in four categories that were launched between February 1994 and January 2004. The four categories (Anatomical Therapeutic Chemical (ATC) codes6 between brackets) are lipid modifying agents, more specifically statins (C10aa); urinary antispasmodics (G4bd); erectile dysfunction drugs (G4be); and (other) antihistamines for systematic use (R6ax). We selected these categories for several reasons. First, they represent a substantial portion of the total pharmaceutical market in dollar sales (around 10%).

6 More information on the ATC codes can be found on the official website of the WHO Collaborating Centre for Drug Statistics Methodology: http://www.whocc.no/atcddd/.
Table 1  Summary Table of Variable Operationalization, Expectations, and Empirical Findings

<table>
<thead>
<tr>
<th>Operationalization of the variable</th>
<th>Expected effect on sales in kg per 1,000 inhabitants aged 15 years and older</th>
<th>Estimated effect on sales in kg per 1,000 inhabitants aged 15 years and older</th>
<th>Were expectations confirmed?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Main effect</td>
<td>Time variation</td>
<td>Main effect</td>
</tr>
<tr>
<td>Regulation</td>
<td>Time-varying</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Manufacturer price</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Physician prescription budgets</td>
<td>Time-varying</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Patient co-payment</td>
<td>Time-varying</td>
<td>–</td>
<td>N.S.</td>
</tr>
<tr>
<td>Marketing efforts to physicians</td>
<td>Time-varying</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Direct-to-consumer advertising</td>
<td>Time-varying</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Other variables</td>
<td>Time-invariant</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Uncertainty avoidance</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individualism</td>
<td>Time-invariant</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Masculinity</td>
<td>Time-invariant</td>
<td>–</td>
<td>N.S.</td>
</tr>
<tr>
<td>Long-term orientation</td>
<td>Time-invariant</td>
<td>–</td>
<td>N.S.</td>
</tr>
<tr>
<td>Economic wealth</td>
<td>Time-varying</td>
<td>+</td>
<td>N.S.</td>
</tr>
<tr>
<td>Cross-country introductory lag</td>
<td>Time-invariant</td>
<td>+/-</td>
<td>–</td>
</tr>
<tr>
<td>Competition</td>
<td>Time-varying</td>
<td>+/-</td>
<td>+/−</td>
</tr>
<tr>
<td>Lagged sales</td>
<td>Time-varying</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

Second, they are typically prescribed by general practitioners, rather than administered in hospitals (e.g., oncological drugs). In the drug categories we study, retail sales represent more than 98% of the total volume, the remaining sales being made through in-patient hospital channels and direct government purchases. Third, these drug categories contain a sufficient number of new drugs that entered after the start of the observation window, which is fixed to 02/1994 because of data handling procedures of IMS Health. We consider the sales growth process across 34 countries that span the globe (see Figure 3), including both developed and developing countries.

The country-level sales data consist of quantity sold in grams in retail per 1,000 inhabitants aged 15 years and older. Such normalization corrects for country size, as advocated by Dekimpe et al. (1998), and accounts for the fact that the drugs considered are mainly designed for the adult population. This correction is particularly relevant as we consider together developed and developing countries for which the age distribution may strongly differ. Fifteen years and older
is a typical cut-off in statistics of international organizations, such as the United Nations or Organisation of Economic Cooperation and Development (OECD), and is used by other scholars in the context of international pharmaceutical markets (Lanjouw 2005). We obtained the sales data from IMS Health for each molecule from its market introduction, with a maximum number of 84 months for drug-country combinations. Census data are gathered from the United Nations Statistics Division website (http://unstats.un.org). Table 2 provides an overview of therapeutic categories, the molecules in our sample with their commercial brand names (one molecule can be sold under different brand names and by different companies), and the period in which the molecule was launched for the first time.

The health-care regulatory data include annual data from multiple sources. The first source is URCH Publishing, an independent information provider for the biotechnological, chemical, and pharmaceutical industries. The second source is the OECD, which publishes reports on health regulation among its member states (e.g., Jacobzone 2000). The third source includes local health officials and ministries of health, contacted individually.

We operationalize regulation through five dummy variables. The first (manufacturer price regulation) takes the value 1 if the government directly controls the manufacturer price for drugs, 0 otherwise. The second dummy variable (regulation of marketing efforts to physicians) takes the value 1 if the government has regulated the number of detailing visits a manufacturer can make, the number of meetings a manufacturer can organize, or the number of samples a manufacturer can dispense to physicians, 0 otherwise.

The third dummy variable (regulation of physician prescription budgets) takes the value 1 if regulation restricts the prescriptions a physician can write, 0 otherwise. The fourth dummy variable (patient co-payment regulation) takes the value 1 if patients are required to provide some form of co-payment for prescription drugs, 0 otherwise. The fifth dummy variable (regulation of DTCA) takes the value 1 if DTCA is prohibited, 0 when it is allowed.

While the regulatory environment is intrinsically more complex than the presence or absence of a restriction, it is hard to obtain more detailed information on the national peculiarities for each of the regulatory variables, and to code them in a sound and economical manner. Therefore, other scholars have also resorted to a dummy coding in a manner similar to ours (Kyle 2007). Yet the combination of these five regulatory dimensions still offers 2^5, or 32 possible regulatory frameworks, of which 13 actually occur in practice. We also find quite a significant cross-national diversity in terms of each of the regulatory variables. For instance, about 73% of the countries have manufacturer price regulation; 18% regulate marketing

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Table 2: Our Sample of Pharmaceutical Drugs

<table>
<thead>
<tr>
<th>ATC code</th>
<th>Molecule</th>
<th>Brands</th>
<th>Time of first introduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>C10aa</td>
<td>Atorvastatin</td>
<td>Lipitor</td>
<td>1/1/1997</td>
</tr>
<tr>
<td></td>
<td>Cerivastatin</td>
<td>Baycol, Lipobay</td>
<td>4/1/1997</td>
</tr>
<tr>
<td></td>
<td>Fluvastatin</td>
<td>Lescol</td>
<td>2/1/1994</td>
</tr>
<tr>
<td></td>
<td>Rosuvastatin</td>
<td>Crestor</td>
<td>2/1/2003</td>
</tr>
<tr>
<td>G4bd</td>
<td>Solifenacin</td>
<td>Visicare</td>
<td>8/1/2004</td>
</tr>
<tr>
<td></td>
<td>Tolterodine</td>
<td>Detrol</td>
<td>11/1/1997</td>
</tr>
<tr>
<td>G4be</td>
<td>Alprostadil</td>
<td>Caverject, Muse, Viridal</td>
<td>3/1/1994</td>
</tr>
<tr>
<td></td>
<td>Apomorphine</td>
<td>Uprima, Ixense</td>
<td>6/1/2001</td>
</tr>
<tr>
<td></td>
<td>Sildenafil</td>
<td>Viagra, Revatio, Caverta</td>
<td>1/1/1998</td>
</tr>
<tr>
<td></td>
<td>Tadalafil</td>
<td>Cialis</td>
<td>1/1/2003</td>
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<tr>
<td></td>
<td>Vardenafil</td>
<td>Levitra</td>
<td>3/1/2003</td>
</tr>
<tr>
<td>R6ax</td>
<td>Desloratadine</td>
<td>Clarinex</td>
<td>1/1/2001</td>
</tr>
<tr>
<td></td>
<td>Epinastine</td>
<td>Elestat</td>
<td>6/1/1994</td>
</tr>
<tr>
<td></td>
<td>Fexofenadine</td>
<td>Allegra, Telfast</td>
<td>8/1/1996</td>
</tr>
<tr>
<td></td>
<td>Mizolastine</td>
<td>Mizolene</td>
<td>1/1/1998</td>
</tr>
</tbody>
</table>

1Cerivastatin was withdrawn from the market in 2001. We only include sales data of this molecule until the withdrawal announcement to avoid a structural break in the sales data.
efforts to physicians; 41% limit physicians’ prescription budgets; 85% apply a patient co-payment system; and 97% prohibit DTCA.

As readers may be relatively unfamiliar with countries’ regulation profiles, Figure 3 summarizes all regulation dummies in a composite regulation score (from “no regulation” (on all of the above five aspects), to “fully regulated” (on all of the above five aspects)), on which we position each of the countries. The interior points are determined by the number of regulatory restrictions in effect. This ranking illustrates the intensity of the national regulatory regimes, and, for simplicity, makes no qualitative distinction between the various regulatory aspects.7 The position of countries may change, if they made regulatory changes in the data window. An en dash following the year means “as of that year,” while an en dash preceding the year means “before that year.” Note, however, that changes in regulation are quite rare in our data tries may change, if they made regulatory changes.

For cultural characteristics of countries, we use Hofstede’s (2001) values on four cultural dimensions: uncertainty avoidance, individualism, masculinity, and long-term orientation. These values have been shown to be very robust indicators of national culture (Steenkamp et al. 1999, Tellis et al. 2003). Given the relatively high number of missing values in the long-term orientation dimension, we proceed to the imputation of missing values using the multiple imputation method for continuous data proposed by Schafer (1997, 2002). This method has been recently advocated and used in a similar context by Deleersnyder et al. (2009).

Quarterly GDP per capita is used as a measure of economic wealth, obtained by dividing production-based GDP data from Thomson DataStream at current prices, converted to U.S. dollars, without seasonal adjustment, by United Nations midyear population data for all 34 countries involved. The introductory lag effect for drug i in country j at time t is operationalized as the number of months elapsed since the first introduction of drug i in any country. This variable is time-invariant. The effect of competition on drug i in country j at time t is operationalized as the number of competitive molecules that belong to the same ATC category and that are present/sold in country j at time t. To have a complete picture of the market conditions, we also include the molecules introduced before 1994 and that belong to the same ATC category. This variable is time-varying. Finally, lagged sales for drug i in country j at time t is operationalized as the sales of the same drug i in the same country j in t – 1.

To conclude, note that the tests we conducted (variance inflation factors, condition indices) did not detect severe multicollinearity.

Model Specification
Let $y_{ijt}$ be the monthly grams per thousand inhabitants aged 15 years and above of drug i belonging to the drug category (ATC) c sold in country j at time t, with $t = 1, \ldots, T_{ij}$, where $T_{ij}$ is the number of sample points for drug i from category c in country j. All categories that we consider are available in every country. We specify a nested structure in which each drug i is nested in a particular drug category c. By doing so, we allow for higher correlation among sales of drugs belonging to the same ATC. For all drugs, we have $y_{ijt} = 0$ up to $t = 0$. We model $y_{ijt}$ using the following time-varying coefficient model:

$$y_{ijt} = \beta_{ij1} PRICEREG_{ijt} + \beta_{ij2} PROMREG_{ijt} + \beta_{ij3} BUDGREG_{ijt} + \beta_{ij4} COPAYREG_{ijt} + \beta_{ij5} DT CAREG_{ijt} + \beta_{ij6} G D P_{ijt} + \beta_{ij7} COM P_{ijt} + \beta_{ij8} PRO MREG_{ijt} + \beta_{ij9} COPAYREG_{ijt}$$

$$+ \beta_{ij10} \gamma_{ijt-1} + \beta_{ij11} \bar{P}_{ijt} + \eta_{ijt}$$

$$\text{with } \eta_{ijt} = \alpha_c + \gamma_{ic} + \epsilon_{ijt}, \alpha_c \sim N(0, \sigma_{\alpha_c}^2), \gamma_{ic} \sim N(0, \sigma_{\gamma_{ic}}^2), \text{and } \epsilon_{ijt} \sim N(0, \sigma_{\epsilon_{ijt}}^2),$$

(1)

where the $\beta_p$, $p = 1, \ldots, 13$, are the smoothly time-varying effects of the covariates under investigation. This model allows us to disentangle the effect of each variable on new drug sales from its market introduction until 84 months after introduction. Upon a preliminary visual inspection of the residuals, we allow the error term to be heteroskedastic with $\sigma_{ijt}^2 = \sigma^2 |t|$.8 In the above equation, PRICEREG$_{ijt}$ indicates whether the manufacturer price is regulated (≠1) or not (=0) in country j in month t, PROMREG$_{ijt}$ specifies whether marketing efforts to physicians are regulated (≠1) or not (=0) in country j in month t, BUDGREG$_{ijt}$ denotes whether regulation limits physicians’ prescription budgets (≠1) or not (=0) in country j in month t, COPAYREG$_{ijt}$ indicates whether a co-payment system is applied (≠1) or not (=0) in country j in month t, and DT CAREG$_{ijt}$ stands for

7 While our model in the Results section includes dummies for each of the five different regulations, we are not able to disclose them at the individual country level because these data are, in part, proprietary. For more detail on each country’s regulatory environment, we refer readers to URCH, which publishes many regulatory reports on international pharmaceutical markets at a relatively low price, and the OECD, which publishes reports on the pharmaceutical regulation of its member states.

8 The specification of heteroskedastic errors yields a better fit (LL = −27,002; BIC = 54,304) than the specification of homoskedastic errors (LL = −27,226; BIC = 54,752).
the prohibition of direct-to-consumer advertising (=1) or its allowance (=0). In turn, \( UA_j \), \( IND_j \), \( MAS_j \), and \( LTO_j \) represent, respectively, the time-invariant uncertainty avoidance, individualism score, masculinity, and long-term orientation scores of country \( j \), while \( GDP_j \) is the GDP per capita in country \( j \) in month \( t \). The competition variable \( COMP_{ijt} \) represents the number of competitive molecules that belong to the same ATC category as drug \( i \) and that are sold in country \( j \) at time \( t \). The lagged sales variable \( \tilde{y}_{ijt-1} \) represents the sales of the same drug in the same country in \( t - 1 \). Finally, the introductory lag variable \( IC_{ij} \) indicates the number of months elapsed since the first introduction of drug \( i \) in any country of the world and its introduction in country \( j \). The superscripted bars indicate that variables are standardized (i.e., zero mean, unit variance) across countries and drugs to make the effects comparable to each other.

We also account for heterogeneity across drug categories by including a category-specific intercept \( \alpha_c \) and we specify the nesting structure among drugs within the same category by including \( \gamma_{ic} \). The role of the category- and drug-specific intercepts is to capture a large part of the variation in sales stemming from the molecules’ idiosyncrasies, e.g., drug indication, marketing instruments used, or drug compliance (Bowman et al. 2004). Alternatively, one could have made the time-varying covariate effects \( \beta_{1t}, \ldots, \beta_{13t} \) drug-specific, but such a specification would exhaust the degrees of freedom and lead to an overparameterization of the model.

We model the time-varying coefficients \( \beta_{p1}, \ldots, \beta_{13p} \) in Equation (1) using regression splines, which can be seen as a compromise between linear regression and nonparametric regression models. This class of models is also called conditionally parametric because the time-varying parameters are nonparametric functions, whereas the model is parametric for a specified \( t \). More specifically, we opt for a penalized splines approach, introduced as P-spline smoothing in Eilers and Marx (1996). Penalized splines build on recent developments in semiparametric modeling (for an overview of semiparametric models in marketing, see Van Heerde (2000) and have been increasingly advocated and used in statistics journals and books (see, e.g., Chiang et al. 2001, Huang et al. 2002, Kauermann 2005, Wu and Zhang 2006).

The advantage of splines compared to the specification of a linear, quadratic, or even cubic trend lies in the fact that they do not impose any assumption (linear, quadratic, or cubic) as to the interaction of the explanatory variables with time. They constitute a highly flexible and modular approach by which the time-varying parameters can follow any pattern (Coull et al. 2001, Ruppert et al. 2003), at a low cost in terms of degrees of freedom. Such flexibility is important, given the novelty and the complexity of the research problem, which makes our hypotheses somewhat exploratory. Moreover, some variables may show discontinuities, which suit the semiparametric approach. For instance, the cross-country introduction lag may show discontinuities because of a lack of infrastructure in some lagging countries. We compare our results with the estimates from a fully parametric model in the Discussion section and find them to lead to very similar findings.

Other estimation methods can be used such as penalized least squares (Hastie and Tibshirani 1992) or local plane fitting (Cleveland et al. 1992). Time-varying coefficient models can also be estimated in a Bayesian dynamic linear model framework (Neelamegham and Chintagunta 2004, Van Heerde et al. 2004) and/or using Kalman filter (Naik et al. 1998, West et al. 1985, West and Harrison 1997). To account for the dynamic effect of past values of the explanatory variables on sales, as is the case with dynamic linear models, we include a lagged dependent variable term in Equation (1).

The general idea behind splines is that any smoothly varying function can be seen as a linear combination of basis functions, which usually are polynomial functions of low degree (e.g., linear in case of linear splines, cubic in case of cubic splines). Hence, we can write the time-varying coefficients as

\[
\beta_{pt} = \beta_{p0} + \beta_{p1} t + \sum_{k=1}^{K} u_k \beta_{p(k)} (t - \kappa_k), \quad p = 1, \ldots, 13 \tag{2}
\]

with \( (t - \kappa_k) \), \( k = 1, \ldots, K \), a set of \( K \) linear spline basis functions,\(^{10}\) which in this case are truncated lines with \( \kappa_k \) the truncation point or knot (i.e., the location where the broken lines are tied together). Such a combination of the linear splines basis functions 1, \( t, (t - \kappa_1), \ldots, (t - \kappa_K) \), gives a piecewise linear function, called spline (see Wedel and Lee (1998), for an application of splines in marketing), with \( K \) distinct knots at \( \kappa_1, \ldots, \kappa_K \) chosen in the range of \( t \), in our case between \( t = 1 \) and \( t = 84 \) months. Alternatively, one could specify quadratic or cubic splines (Sloot et al. 2006) using 1, \( t, t^2, (t - \kappa_1)^2, \ldots, (t - \kappa_K)^2 \) for quadratic splines, or 1, \( t, t^2, t^3, (t - \kappa_1)^3, \ldots, (t - \kappa_K)^3 \) for cubic splines.

As we have no prior on the location (on the time axis) where changes in the effect of the explicative

\(^{9}\) Note that the cultural variables are time-invariant. As noted by Steenkamp et al. (1999), a country’s culture is a key environmental characteristic underlying systematic differences in consumer behavior (Triandis 1989). Culture is by nature stable over time and is not supposed to change in the short or medium run.

\(^{10}\) This notation is standard in the statistical literature and indicates that the function equals zero for the values of \( t \) where \( (t - \kappa_k) \) is negative.
variables may occur, we distribute the knots evenly over the complete time period and make sure that we have enough knots to cover the time range. As explained by Sloot et al. (2006), the number of knots should be chosen relative to the number of available observations. While too few knots may result in a model that is too restrictive, too many knots may result in estimation problems. In our application, we use one knot per year that is available, that is \( K = 7 \) knots.\(^{11}\)

The \( u_k^p \) coefficients can be interpreted as weights to each of the basis functions that constrain their relative influence. These weights are penalized; i.e., they are subject to the constraints \( \sum_{k=1}^{K} (u_k^p)^2 < U_p \), for some constants \( U_p \), \( p = 1, \ldots, 13 \). This penalty induces smoothness in the time-varying effects, preventing fits that are too rough or too wavy.\(^{12}\) Previous research has shown that the model defined by Equations (1)–(2), subject to the aforementioned constraints, yields fitted values equivalent to those produced by a linear mixed model estimated (e.g., by Verbeke and Molenberghs 2000), where the intercepts and slopes \( \beta_{p,0} \) and \( \beta_{p,1} \) in Equation (2) are estimated as fixed components and the \( u_k^p \) coefficients are taken as random components with respective variances \( \text{Var}(u_k^p) = \sigma_{2,0}^2, \ p = 1, \ldots, 13 \) (Brumback et al. 1999, Coull et al. 2001, Ruppert et al. 2003). Therefore, the level of smoothing of the time-varying effects is given by \( \sigma_1^2/\sigma_{2,0}^2 \). A higher \( \sigma_{2,0}^2 \) corresponds to a more wavy function, whereas a smaller \( \sigma_1^2 \) corresponds to more smoothness. The level of smoothing is determined by residual maximum likelihood (REML) as demonstrated in Wand (2003), rather than using Akaike’s information criterion or cross-validation methods. As noted by Opsomer et al. (2001), smoothing parameter selection with any data-driven method typically leads to serious overfitting in case of correlated errors. With linear mixed models, however, once the correlation structure is specified, estimation of regression parameters can be carried out without computationally-intensive cross-validation procedures (Kauermann and Komski 2006, Krivobokova et al. 2006). Linear mixed models are very easy to estimate in standard statistical software (e.g., PROC MIXED in SAS or lme in S-PLUS and R). In addition, the mixed model framework permits the use of likelihood ratio tests for model selection and theory testing. More details can be found in Crainiceanu et al. (2005) or Ruppert et al. (2003).

**Instrumenting for Launch Delay**

We deal with the potential endogeneity of the introductory lag variable \( \overline{\Pi}_{ijc} \) in a similar fashion as Desiraju et al. (2004). We find a set of exogenous instruments, denoted by \( Z_{ijc} \), that are correlated with the endogenous variable \( \overline{\Pi}_{ijc} \) but uncorrelated with the residuals \( e_{ijc,t} \) and that have an effect on \( y_{ijc,t} \) only through \( \overline{\Pi}_{ijc} \). As they explained, the underlying idea is to use variation in \( \overline{\Pi}_{ijc} \) that is explained by the exogenous variables for estimation as all other unexplained variation is possibly correlated with the residuals. The approach is similar in spirit to two-stage least squares.

In addition to the exogenous variables depicted in Equation (1), we consider as instrument for \( \overline{\Pi}_{ijc} \) the introductory lag of all other drugs in the same ATC (e.g., drug \( i \)) in country \( j \). This choice is justified by the idea that the introductory lag of the other drugs that belong to the same ATC class \( c \) as drug \( i \) in a particular country \( j \) is a reasonable predictor of the introductory lag of the focal drug in that country. Indeed, we expect that managers’ launch decisions would be affected by similar underlying factors for all drugs in a given ATC, such as the country’s population size or their sales expectations. Moreover, this variable is uncorrelated with the residuals \( e_{ijc,t} \) as confirmed by the high \( p \)-value obtained from the Durbin-Wu-Haussman test for endogeneity (\( p = 0.95 \)).

Formally, we define the introductory lag of other drugs in the same ATC (except drug \( i \)) in country \( j \) as \( \overline{Z}_{i-j,ci} \), the mean value of \( \overline{\Pi}_{ijc} \) for all drugs (except the focal drug \( i \)) in that particular country \( j \). We construct the instrument as the predicted value from a regression of the endogenous variable \( \overline{\Pi}_{ijc} \) on the entire set of exogenous variables in Equation (1) including \( \overline{Z}_{i-j,ci} \), next denoted as \( Z_{ijc} \). Thus, the first stage consists of

\[
\overline{\Pi}_{ijc} = \delta_c + \tau_{ic} + Z_{ijc} \lambda + \omega_{ijc},
\]

with \( \delta_c \) and \( \tau_{ic} \) the category- and drug-specific intercepts and \( \omega_{ijc} \sim N(0, \sigma_{2,0}^2) \). As such, we ensure that \( Z_{ijc} \) and \( e_{ijc,t} \) are uncorrelated by construction. The second stage then consists of using in Equation (1), the predicted value for \( \overline{\Pi}_{ijc} \) given by estimating Equation (3) instead of the endogenous variable \( \overline{\Pi}_{ijc} \). Standard errors are corrected using the results offered by Murphy and Topel (1985).

**Results**

This section presents the estimation results of our model. We first discuss results from the instrumental
For instance, the time-invariant model includes 16 parameters: the effect for each of the 13 explanatory variables, as well as $\alpha_i$, $\sigma_{\epsilon_i}^2$, and $\sigma_r^2$. In turn, the time-varying model includes 26 additional parameters as we now add 2 extra parameters for each of the 13 explanatory variables. The derivation of the number of parameters for the other specifications is pretty straightforward.

The regression of $P_{ij}$ on $Z_{ij}$ in Equation (3) provides an approximate adjusted $R^2 = 0.46$ and the regression of $P_{ij}$ on the instrument $Z_{-i,j}$ alone gives an approximate adjusted $R^2 = 0.40$, indicating that our instrument predicts the endogenous variable reasonably well. To further assess whether the instrumental variable procedure works properly, we compare the estimated effect of the introductory lag when the instruments are used or not. We find that, when using instruments instead of the endogenous variable, the estimated effect of the introductory lag variable is on average 29% higher over the whole time window ($-0.017$ instead of $-0.025$). This result is consistent with our expectations. Recall that managers are expected to make launch decisions to maximize future sales and tend to launch first in countries where they expect higher sales. Thus, an increase in the value of $\epsilon_{ij}$ would lead to an earlier introduction, i.e., a smaller introductory lag, meaning that we should have a negative correlation between $\epsilon_{ij}$ and $P_{ij}$. Therefore, if the endogeneity problem remains uncorrected, we would expect the estimated effect for $P_{ij}$ to be biased downwards, which is consistent with our results.

### Instrumental Variable Procedure

The regression of $P_{ij}$ on $Z_{ij}$ in Equation (3) provides an approximate adjusted $R^2 = 0.46$ and the regression of $P_{ij}$ on the instrument $Z_{-i,j}$ alone gives an approximate adjusted $R^2 = 0.40$, indicating that our instrument predicts the endogenous variable reasonably well. To further assess whether the instrumental variable procedure works properly, we compare the estimated effect of the introductory lag when the instruments are used or not. We find that, when using instruments instead of the endogenous variable, the estimated effect of the introductory lag variable is on average 29% higher over the whole time window ($-0.017$ instead of $-0.025$). This result is consistent with our expectations. Recall that managers are expected to make launch decisions to maximize future sales and tend to launch first in countries where they expect higher sales. Thus, an increase in the value of $\epsilon_{ij}$ would lead to an earlier introduction, i.e., a smaller introductory lag, meaning that we should have a negative correlation between $\epsilon_{ij}$ and $P_{ij}$. Therefore, if the endogeneity problem remains uncorrected, we would expect the estimated effect for $P_{ij}$ to be biased downwards, which is consistent with our results.

### Model Fit Comparisons

In Table 3, we evaluate the model fit with respect to alternate specifications, that is, (1) whether to allow parameters to vary over time (none, all, or some of them); (2) whether to specify linear, quadratic, or cubic spline functions; and (3) whether to include regulatory predictor variables. The fourth column in Table 3 reports the number of parameters for the various specifications.\(^\text{13}\)

We start our estimation procedure by estimating Equations (1)–(2), allowing all effects to vary over time. Our results indicate that this improves model fit. However, when letting all parameters be time-varying, some of the slope parameters $\beta_{ij}$ in Equation (2) turn out to be insignificant. Therefore, we also estimate a variant of the model in Equation (1) with both time-varying and time-invariant effects (i.e., those having a nonsignificant slope above). The latter gets a better Bayesian information criterion (BIC) than a model with time-varying effects only. This result provides evidence that some of the explanatory variables, i.e., physician prescription budgets regulation, DTCA regulation, uncertainty avoidance, individualism, introductory lag, competition, and lagged sales, have a differential impact on the sales of newly launched drugs than on maturing drugs. The other variables appear to have a constant effect over the product life cycle of new drugs. We present the parameters of this model in Table 4 (time-invariant effects) and Figure 4 (time-varying effects).

Our results also indicate that the specification of quadratic or cubic spline functions does not yield a better fit than linear splines when accounting for the additional number of parameters such specifications generate.

Finally, the last line of Table 3 provides the fit measures for this model, excluding the regulatory variables. Such a model yields a substantially lower log-likelihood and a much higher BIC, demonstrating that regulatory differences across countries do considerably improve the model fit. In other words, regulatory regimes provide a valuable and substantive explanation as to the cross-national differences in drug sales.

To evaluate the overall fit of the model, we also calculated an approximate adjusted $R^2$ measure of the best model, which is highly satisfactory (adjusted $R^2 = 0.93$).

\(^{13}\) For instance, the time-invariant model includes 16 parameters: the effect for each of the 13 explanatory variables, as well as $\alpha_i$, $\sigma_{\epsilon_i}^2$, and $\sigma_r^2$. In turn, the time-varying model includes 26 additional parameters as we now add 2 extra parameters for each of the 13 explanatory variables. The derivation of the number of parameters for the other specifications is pretty straightforward.
Parameter Estimates
As our model estimates both time-varying and time-invariant parameters, we depict the time-invariant effects in Table 4 and plot the time-varying parameter estimates, with their 95% confidence bounds in Figure 4 (all results are available in the Technical Appendix at http://mktsci.pubs.informs.org).

We sequentially discuss the effect of (i) regulation, (ii) culture, (iii) economic wealth, (iv) competition, (v) lagged sales, and (vi) introductory lag. Our main findings are in line with the theoretical expectations, summarized in Table 1.

Regarding the role of regulation, each of the regulatory variables, except the regulation of marketing efforts to physicians and patient co-payment, is found to have a significant impact on drug sales. In particular, drug sales tend to be higher in countries with manufacturer price controls than in countries without price controls. While price regulation can cause launch delays (Kyle 2007), it apparently does not constrain availability to patients once a drug is launched. In fact, we find quite the opposite. However, contrary to our expectation, we cannot provide strong evidence of any variation over time in the effect of manufacturer price regulation on drug sales. One possible reason that we do not find strong evidence of time variation in this effect may be that prices change rather smoothly over time and thus may be captured in the diffusion effects (Bass et al. 1994), even if the price in countries with price regulation.
decreases more than in countries without price regulation. Another possible reason may be the collinearity generated by the inclusion of the interaction between price regulation and time, leading to conservative significance levels.

By contrast, the other forms of regulation show a negative effect on drug sales, such as regulation of physician prescription budgets. However, such regulation appears to be more effective for mature drugs while it has little or no impact on newly launched molecules. This result can be explained by the stepwise prescription behavior of physicians.

While regulating marketing efforts of pharmaceutical firms to physicians tends to have a negative, although nonsignificant, effect, the impact of prohibiting DTCA has a highly significant negative effect on new drug sales. This effect appears to be more pronounced for newly launched drugs than for maturing ones. In particular, we find this effect to be the most pronounced around one year after a new drug is introduced.

Co-payment regulation turns out to have a negative and constant effect, although not significant at the 10% probability level, on drug sales. This supports the idea that patients are monetarily penalized for drug usage in co-payment systems. The insignificant effect is likely due to the fact that the presence or absence of such regulation is a rough indicator, compared to the wide variation that exists in co-payment levels.

As to the role of national culture, two of the cultural dimensions, uncertainty avoidance and individualism, tend to have a differential impact on newly launched versus maturing drugs. In particular, we find that drug sales are higher, although nonsignificant at the 10% probability level, in countries with high uncertainty avoidance than in countries with low uncertainty avoidance. This finding can be related to the lower subjective health of inhabitants of uncertainty avoidant nations (Hofstede 2001). This result interestingly complements previous research on international diffusion as uncertainty avoidance has usually been found to temper sales growth of new durables. The nonsignificance of the parameter suggests that the latter effect may still play a role, particularly for recently launched drugs for which we find no effect at all. This may result from the relatively higher perceived risk associated with adopting very new drugs. However, over time this effect is gradually overwhelmed by the effect of uncertainty avoidance on subjective health.

In addition, we find that new drug sales are higher in individualist countries than in collectivist countries, which confirms that the former may spend more of their budgets on drugs (as we control for GDP in the estimation) and may seek doctor consultation more actively. However, we also find that this difference is more pronounced for maturing drugs than for newly launched ones. This result can be explained by the stepwise prescription behavior of physicians.

Furthermore, our results indicate that new drug sales are lower in masculine countries than in feminine countries, which corroborates our expectations that drug intake by males would be considered a weakness in masculine societies. Given that this argument holds for both new and maturing drugs, we find this effect to be constant over time.

Long-term orientation is found to have a negative, significant, and constant effect on drug sales, which supports our hypothesis that patients and physicians in societies high in long-term orientation refrain from classical drug therapy.

Economic wealth, in turn, positively affects new drug sales, which can be related to the fact that wealthier countries have better developed health infrastructure and can more easily afford new drugs than less wealthy countries. As to the effect of competition on new drug sales, we find little negative impact of the number of competitive molecules available in the market. Our results corroborate the findings of Desiraju et al. (2004) for the antidepressant drug category. We do not find evidence that competition among molecules in the same ATC class expands the category. Lagged sales have a significant, positive, and increasing effect on sales, which points to the inertia in the physicians’ prescription behavior and the contagion among physicians and patients. Finally, the effect of the introductory lag (when using the instrument) turns out to be nonsignificant at the 5% probability level, as is also the case in the global diffusion study of Dekimpe et al. (1998). In line with our expectation, this effect, although nonsignificant, is more pronounced for recently introduced drugs. Note that the pattern of variation over time of this effect nicely illustrates the potential of P-splines in modeling discontinuous phenomena.

To assess the robustness of our findings, we reestimate our model with alternate specifications as to the coding of the regulatory variables, the number and location of knots for the time-varying effects, and the inclusion of interaction effects between the regulatory variables and the ATC categories as well as the lagged sales. Our results turn out to be highly robust.

Discussion
This section first explores the implications of our findings for public policy and managers. We then turn to the study’s limitations, and opportunities for future research.
Implications
Of interest for both managers and public policy makers, we find that regulations, in general, matter. Pharmaceutical manufacturers cannot ignore regulatory regimes as they affect the (volume) attractiveness of a country. For instance, drug volumes are ceteris paribus higher in countries with manufacturer price regulation, while they are lower in countries with direct-to-consumer advertising or prescription budget restrictions. These insights may be of value to managers developing international launch plans, and for their expectations of sales growth in markets they enter. In addition, this study confirms that the cultural and economic characteristics of countries affect their attractiveness for pharmaceutical firms. Managers could therefore locate their markets of interest on a map that accounts for regulatory, cultural, and economic dimensions that would guide their strategy.

Our study is also relevant for public policy. First, public policy makers are interested in international variation in new drug sales, as they may wish to compare drug consumption—for health-care system quality or cost reasons—in their own country to others. Second, policy makers want to know the relationship between the regulatory restrictions they enforce and drug consumption. For instance, some regulatory restrictions have a larger effect on newly launched drugs than on maturing drugs, and vice versa. We find that sales of maturing drugs suffer more from prescription budget regulation than sales of newly launched drugs. By contrast, the prohibition of DTCA depresses sales of newly launched drugs more than sales of maturing drugs. Insights such as these provide the regulator with guidelines for the enforcement, monitoring, and adaptation of the regulatory environment. Obviously, these latter recommendations need to be treated with caution, in view of the Lucas critique that may apply to our model (Bronnenberg et al. 2005).

Limitations and Future Research
Similar to most, if not all, studies on international new product growth, we do not study firms’ marketing instruments. Our interest in obtaining insights from a very large set of countries that includes developing countries prohibits the gathering of consistent information on pricing or marketing efforts. Because of the lack of marketing mix information and the intermediation of marketing mix on the effects of regulation on sales, our hypotheses are contingent upon marketing mix effects on sales. The drawback is that our model cannot distinguish between demand (e.g., physician prescription budget regulation may depress drug demand) and supply effects (e.g., restrictions on marketing efforts may restrict the amount of detailing and thereby physician prescriptions). The mere presence of regulations may also “force the hand” of managers and lead to more creative, efficient ways of marketing their products. While our study takes a first step in exploring the role of regulation in the international sales growth of pharmaceutical drugs, further research that disentangles demand and supply would be of great interest to complement our findings.

In addition, we do not model the effects of the national health infrastructure and status, such as health expenditures, number of doctors, or hospital beds per capita. Yet health infrastructure and health status are known to be very much driven by the economic wealth of a country (Desiraju et al. 2004), and we therefore expect it to be partly captured by the effect of GDP per capita in our model.

Third, our dependent variable in this study is sales. We do not discern whether high or low sales are good or bad for society, e.g., in terms of over- or underconsumption of drugs. Some have argued that drug usage is too low. For instance, Hassett (2004) illustrates that many deaths can be prevented by higher statin intake, not only in more restrictive countries (26,000 preventable deaths in Italy over five years), but even in the most intensive drug market (19,000 preventable deaths in the United States over five years). Others—mostly in Western Europe—have argued that drug usage is typically too high. The study of the influence of country characteristics on patient welfare, rather than new drug sales, may yield interesting insights.

Fourth, our sample of countries is probably biased towards more developed health markets because the motivation to cover the market for data intermediaries, such as IMS Health, is higher, while the hurdles to do so are smaller. Moreover, independent variables in models such as ours (e.g., the inventory of the regulatory environment) are also more likely to be missing for low income countries than for developing or developed countries. This sample selection issue should lead to some caution toward our results, even though our sample contains more (developing and low-income) countries than many prior studies in this area.

Fifth, future research on cross-country spillover effects would also be valuable. Such spillover effects may occur in (1) sales—e.g., because physicians affect each other across country borders; (2) marketing instruments—e.g., because manufacturers set their marketing policy on the pan-regional level, rather than the country level; and (3) regulation—e.g., because countries mimic each other in regulatory behavior.

Overall, there is still much research to be done on international growth patterns of new pharmaceutical drugs. Our study is one of the first on the role of regulation in international new product growth processes.
This area is worthy of more academic attention, given its importance.

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