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EFFECTS OF REGULATION ON DRUG LAUNCH AND PRICING IN INTERDEPENDENT MARKETS

Patricia M. Danzon Andrew J. Epstein

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ABSTRACT

This study examines the effect of price regulation and competition on launch timing and pricing of new drugs. Our data cover launch experience in 15 countries for drugs in 12 therapeutic classes that experienced significant innovation over the decade 1992-2003. We use prices of established products as a measure of the direct effect of a country's own regulatory system, and find that launch timing and prices of innovative drugs are influenced by prices of established products. Thus, if price regulation reduces drug prices, it contributes to launch delay in the home country. New drug launch hazards and launch prices in low-price countries are also affected by referencing by other, high-price countries, especially within the EU, as expected if manufacturers delay launch in low-price markets to avoid undermining higher prices in other countries. Thus, referencing policies adopted in high-price countries can impose welfare loss on low-price countries. Prices of new drugs are influenced mainly by prices of other drugs within the same subclasses, however, dynamic competition from new subclasses undermines new drug launch in older subclasses. Association with a local firm accelerates launch only in certain regulated markets. These findings have implications for US proposals to constrain pharmaceutical prices in the US through external referencing and drug importation.

Patricia M. Danzon
Health Care Systems Department
The Wharton School
University of Pennsylvania
204 Colonial Penn Center
3641 Locust Walk
Philadelphia, PA 19104
and NBER
danzon@wharton.upenn.edu

Andrew J. Epstein School of Public Health Yale University 60 College St, Room 301 New Haven, CT 06520 andrew.epstein@yale.edu

I. Introduction

New drugs are potentially global products and can contribute importantly to health outcomes for consumers and expenditures for payers. Prompt launch is also critical for drug manufacturers, given the fixed patent life over which to recoup the high costs of R&D.¹ In fact, in a study of the 1990s launch experience of 85 drugs in 25 industrialized countries, only roughly half of the potential country-compound launches occurred, and many of the eventual launches involved months or years of delay (Danzon, Wang and Wang, 2005). Earlier studies (e.g., Peltzman, 1973; Grabowski and Vernon, 1978) found that increased regulatory requirements for proof of safety and efficacy ("registration") contributed to launch lags in the US in the 1960s and 1970s. However, in the 1990s the US and the EU harmonized and accelerated new drug review, such that remaining differences in registration requirements cannot fully explain observed launch lags, especially between EU countries.

By contrast, price regulation has become more complex and potentially plays an important role in launch lags, through both direct effects on drug launch in the regulating country and indirect (spillover) effects on launch in other countries. Price regulation may reduce launch incentives directly, by reducing a manufacturer's expected price and NPV. This direct effect of regulation predicts that non-launch is most likely in small countries and for drugs with low expected sales volume, assuming some fixed costs of launch. Price regulation may also lead to bureaucratic delay and strategic delay by firms or regulators to influence the ultimate price.² Even if a country's price regulation leads to launch lags in that country, the welfare effect for its citizens is ambiguous *a priori*, because any foregone health benefit due to fewer/later launches

¹ Pharmaceutical R&D takes on average 8-12 years and costs roughly \$802m. (in 2001 US dollars) per new compound approved in the US (DiMasi, Hansen and Grabowski, 2003).

² Delay could also reflect launch budget constraints faced by firms, in which case a rational strategy may be to launch first in the most profitable markets and use the revenues generated to cover launch in less profitable markets.

must be weighed against the savings from lower drug prices and better pre-launch information about a drug's relative safety and effectiveness.³

More problematic from a social welfare perspective are the indirect effects of price regulation that arise when one country regulates its drug price by reference to the price of the same drug in other countries ("external referencing"). For example, Canada caps the price of innovative new drugs at the median price in seven countries and some EU countries use the mean or minimum price in a group of referenced countries. By undermining market segmentation and price discrimination, external referencing creates incentives for a firm to delay or not launch in relatively low-price countries that are referenced by potentially higher-price countries. The welfare consequences in the referenced low-price countries are clearly negative, since they suffer reduced access to new drugs and possibly higher prices due to external referencing by higher-price countries. Parallel trade (drug importation) may also contribute to non-launch and higher prices in low-price countries that are potential sources of parallel exports, with possibly negative welfare consequences in these countries. Parallel trade is legal between EU members, and is under consideration in the US.

Previous studies of launch experience for new drugs (Danzon, Wang and Wang, 2005; Lanjouw, 2005; Kyle, 2006, 2007) have generally concluded that price regulation contributes to launch delay. However, these studies used proxy measures for regulation that were rough and sometimes inaccurate; they lacked data on prices of competitor products, a critical control variable; they estimated effects on launch lags but not launch prices; and none clearly distinguished between the direct (own-country) and the indirect (spill-over) effects of regulation.

³ Negative external effects may accrue to other countries if the regulated prices result in suboptimal contribution to joint costs of R&D (Danzon and Towse, 2003).

This paper examines the contribution to drug launch lags and pricing of a country's own regulatory system vs. spillover effects from other countries. We also estimate effects of dynamic competition and launch by local corporations. We use quarterly sales data, by drug, from IMS Health⁴ from 1992-2003 for 15 major countries and 12 major therapeutic classes, all of which experienced entry of a new subclass during our study period (for example, in the antiulcerant class, the proton pump inhibitors [PPIs] displaced the H2-antagonists). We refer to new and old subclasses as "superior" and "inferior," but intend no judgment about their relative value.

Because price regulatory systems are complex and multidimensional, categorizing such systems by binary indicators is imprecise and may lead to severe measurement error. We therefore use the average price of established products, by subclass, as a measure of the net effect of regulation on actual prices and hence on the expected price of a new drug. We distinguish competitor prices for superior vs. inferior subclasses to test the importance of static (within-subclass) vs. dynamic (between-subclass) competition, in particular, whether the availability of cheap products from old subclasses constrains launch prices for new drugs in innovative subclasses. We also test for first-mover advantage.

We estimate a two-equation model of launch hazard and launch price for molecules in superior and inferior subclasses separately, to permit comparison between them. The launch hazard equation is estimated using a complementary log-log (hereafter "clog-log") model with time-varying covariates. We also estimate a random effects clog-log model and a split population clog-log model (Schmidt and Witte, 1989) to test for various forms of molecule-level heterogeneity. The launch hazard results are robust to specification, although both the split population and the random effects hypotheses are confirmed. For the launch price equation, we

⁴ IMS Health Inc. is a market research company that collects data on sales of pharmaceuticals in all significant world markets.

report estimates of price conditional on launch from OLS and GLS random effects models. We test for selection bias, using a two-step Heckman selection estimator (Heckman, 1979). Although the inverse Mills' ratio is sometimes significant, the launch price results are also robust to specification.

Our results confirm that launch hazards and prices of new drugs are positively associated with prices of established products. Thus, to the extent that regulation reduces drug prices, it directly contributes to launch delay/non-launch. Regulatory referencing by foreign countries also plays a role, especially in low-price EU countries. We find no evidence that manufacturers use delay as a strategy to obtain a higher price within a given country. Launch by local corporations accelerates launch timing but does not affect launch prices, with larger effects for local originator firms than local licensees. These effects are concentrated in a few, regulated countries. Launch timing and prices are influenced mainly by prices of drugs in the same subclass, with at most minor effects from drugs in older subclasses. Late-launching drugs in the inferior subclasses have more limited launch success than new drugs in new subclasses, confirming the importance of non-price dynamic competition.

Estimating the welfare consequences of regulation is complex and beyond the scope of this paper.⁵ However, the evidence here does suggest that indirect regulatory strategies that rely on referencing to foreign prices or importing from foreign countries can impose significant welfare loss on these foreign countries due to launch lags, non-launch and higher prices. Such

⁵ Philipson et al. (2008) estimate the welfare effects of launch delay due to registration regulation in the US. In their model, registration regulation is assumed to reduce drug risk but delay consumer access to drug benefits, which are measured by actual drug prices. By contrast, our focus is on launch delay due to price regulation. The presumption of price regulation is that unregulated consumer demand is not a valid measure of consumers' marginal benefits from drugs when consumers face minimal or zero out-of-pocket payment, as in most countries. If consumer demand for a drug is highly inelastic due to low/zero co-payment, but consumers ultimately must pay the full price through taxes or insurance premiums, then regulation that constrains prices to reflect the "true" value of drugs can increase consumer welfare in that country. Of course, if regulators set prices below the true value of new drugs to consumers, there may be long-run negative consequences due to reduced R&D.

effects could be particularly severe if the US were to adopt external referencing or parallel trade, given the size of the US market (45 percent of global pharmaceutical sales) relative to other countries.⁶

The remainder of the paper is organized as follows. Section II reviews the relevant literature, Section III outlines regulatory regimes and expected effects, and Section IV describes the theoretical model. Section V details the data and empirical methods, Sections VI-VIII describe the results, and Section IX concludes.

II. Literature

Several recent studies have examined effects of price regulation on lags in new drug launch. Danzon, Wang and Wang (2005) studied the launch experience of 85 new drugs in the 25 leading markets in the 1990s, focusing on drugs that had met registration requirements of one of the two strictest agencies (the US Food and Drug Administration [FDA] and the UK Medicines Agency) and hence could potentially meet registration requirements in other countries. Their analysis used the average price of all competitor products at global launch as a proxy for the direct regulatory effect on expected price. It concluded that price regulation that leads to low prices deters launch and that the potential for price spillovers to higher-price markets exacerbates launch lags in low price markets. Launch by a local firm on average increased launch probability, but country-specific differentials were not examined.

Kyle (2007) used a larger, more heterogeneous sample of compounds and countries, a longer time period (1980-1999), and a vector of dummy variables for types of regulation and

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⁶ As of May 2008, regulating US prices based on external referencing to foreign prices has been proposed but so far not enacted. Drug importation has been approved but so far is not implemented, because the Secretary of Health and Human Services has been unwilling to certify that safety and cost reduction requirements have been met.

price rank.⁷ She concluded that price controls reduced launch probability in countries that impose them. However, the estimated effects may be biased because the regulatory indicators from 2000/2002 sources incorrectly classify some countries, do not reflect changes over the 20-year analysis period, and are missing for several countries.⁸ More fundamentally, indicators for regulatory type imperfectly capture the multidimensional and heterogeneous detail of price control regimes across countries and over time. Country-specific price data were not available, except for a 2002 indicator for a country's price rank. This was negatively related to launch probability prior to 1995 and insignificant after 1995, which seems inconsistent with the conclusion that price controls delay or reduce launch probabilities. Indicators for prior launch in a high-price or low-price country were positively associated with launch probabilities in other countries, consistent with price spillovers. However, differential effects across countries were not estimated; thus, spillovers related to regulatory regime cannot be distinguished from other unobserved factors that could lead to closely sequenced launches, such as coordinated regulatory filings.⁹ Kyle (2006) focuses on launch in the G7 countries. She finds that a binary indicator for

⁷The large and heterogeneous sample probably includes some drugs that could not meet requirements of the strictest agencies, such as the US FDA, the UK Medicines Agency and the EMEA. A count of Medline citations is included as a control for product quality; however, this US-centric measure may be a biased proxy for quality as perceived by non-US countries, especially for drugs not launched in the US.

Several examples can illustrate the measurement error that results from applying the 2000/2002 listing of regulatory indicators to the 1980-1999 data. In the paper, "price controls refer to a cap on either the ex-manufacturer price or the amount a national health service pays for a pharmaceutical product." By this definition, therapeutic reference pricing should count as a form of price control, because the reference price is a cap on reimbursement for all drugs in a class. In fact, therapeutic reference pricing is included as a separate dummy variable and has a coefficient that is positive and sufficiently large (0.854) to dominate the negative price control coefficient (-0.418). But, this effect appears to be spurious due to measurement error: although 7 countries are classified as having therapeutic reference pricing, in fact only the Netherlands had comprehensive therapeutic referencing, and this was only introduced in 1991. Similarly, 6 countries are categorized as "Pharmacoeconomic evidence recommended," but in fact pharmacoeconomic evidence was used informally to support price negotiations in many countries from the early 1990s. In the late 1990s, both the UK and Canada required pharmacoeconomic evidence, but neither of these countries are categorized recommending use of pharmacoeconomic evidence. Given this measurement error, the large negative coefficient on this variable is unlikely to provide an accurate measure of the effect of requiring pharmacoeconomic evidence. Germany is listed as having a Prescribing budget, but in fact it was only in place from 1993-2000, and never enforced.

High (1%) significance levels for most explanatory variables are surprising and may be upward biased by including multiple indications for the same compound. Because follow-on indications are usually not subject to

drug price controls is negatively associated with launch probability, but is significant only in some specifications. This analysis also finds significant effects of firm characteristics on launch.

Lanjouw (2005) examined first launch of a large, heterogeneous sample of new drugs in 68 countries between 1982 and 2002, using covariates measured at first global launch. She also used binary indicators for price regulation regimes that are invariant across the 20-year study period and lacked data on prices. She concluded that even moderate price controls in high income countries reduced the long-run likelihood of drug launch, while price controls in less wealthy countries reduced launch probability in the short-run but not the long-run. Spillover effects were not addressed.

Our study adds to this literature in several ways. We use product-specific data on prices for competitor products, categorized into new ("superior") and old ("inferior") subclasses. These prices for established, competitor products provide the most accurate available measure of the net effect of regulation on the expected price for a new drug. Separate analysis of launch experience of drugs in new vs. old subclasses provides evidence on the dynamic (between subclasses) vs. static (within subclass) competition in this industry. Our analysis is the first to examine determinants of launch prices as well as launch lags, which provides some validation of the launch hazard estimates. We distinguish carefully between direct effects of regulation vs. indirect effects through cross-national spillovers, since their welfare implications are very different. Furthermore, our detailed analysis shows that launch by a domestic corporation affects launch hazard but not launch price, with larger effects if the local firm is the originator rather than just a licensed marketing partner. We show that these effects are significant only in a few regulated countries and are not general, as suggested by previous studies. Our findings are robust

separate price regulation and simply receive the same price as previous indications, with minimal price-related regulatory delay, including them could also bias coefficient estimates.

to estimation methods, including controls for within-molecule correlation and unobserved heterogeneity.

III. Pharmaceutical Regulation: Registration and Price/Reimbursement

New drugs face two possible regulatory hurdles, registration and price approval.

Registration

In all countries, drug registration and market access require proof of safety, efficacy and manufacturing quality. In the 1990s, the US FDA and counterpart agencies in Europe and Japan harmonized some requirements while retaining autonomy in data evaluation and decision-making. The FDA adopted user fees to hire more reviewers and created fast track and priority review procedures. Since 1995, the newly-created European Medicines Agency has offered both centralized and mutual recognition procedures that can lead to simultaneous registration of new drugs in all EU countries, as an alternative to the traditional country-by-country review through national drug approval agencies. Thus cross-national differences in drug registration regulation cannot explain large systematic differences in launch lags among EU countries or between the US and EU. Japan is an exception in retaining special requirements, including clinical trials on Japanese citizens.

Price/Reimbursement Regulation

Once a new drug clears registration hurdles, most countries with national health insurance systems require price approval as a condition of reimbursement.¹¹ Countries use one or both of two criteria to set launch prices: (a) "internal referencing" to prices of competitor

¹⁰ Under the mutual recognition procedure, a manufacturer selects one rapporteur country to review the application; other countries have 90 days in which to challenge the approval, otherwise it takes effect automatically. The centralized procedure is mandatory for biologics and optional for other innovative drugs.

¹¹ Price approval is generally not required if the drug is launched without reimbursement, but such unreimbursed launch is rare, except for "lifestyle" drugs.

products in the same therapeutic class, with potential for mark-ups for superior efficacy based on pharmacoeconomic data, ¹² or (b) "external referencing" to the minimum, median or mean of prices of the same drug in specified comparator countries. Most price regulatory regimes disallow post-launch price increases, and price cuts are sometimes mandated; hence the launch price is critical to a drug's life-cycle price profile. Internal referencing may entail bureaucratic delay and possibly strategic delay if firms (regulators) hold out to achieve a higher (lower) price. However, regulators should have incentives to weigh any costs associated with launch delay against the benefits in lower prices, with no significant spillover to other countries.

By contrast, because external referencing regimes benchmark their price to the price of the same drug in other countries, they create incentives for firms to delay launch in referenced lower-price countries until prices have been established in other potentially higher-price countries. Thus countries that use external referencing can impose launch delay and associated costs on the countries that they reference.¹³

Identifying the contribution of these regulatory regimes to drug launch experience in specific countries is complex because most countries use multiple forms of regulation, including both internal and external referencing, and the details of each regulation type differ across

¹² Internal referencing may involve informal negotiations between the manufacturer and the regulator, as in France, or a more mechanistic reference price (RP) reimbursement system as in the Netherlands. Under therapeutic RP, drugs are classified based on mechanism of action and indication; generic RP groups drugs only with the same molecule, hence mainly off-patent drugs with generic equivalents. All drugs in a group receive the same reimbursement or reference price. A manufacturer may in theory charge a higher price, but the patient must pay any excess over the RP.

¹³ For example, suppose that in the absence of referencing, drug prices would be roughly proportional to GDP per capita. If high-income countries regulate their prices by referencing the mean price in a group of countries that includes some high- and some low-income countries, the low-income countries are likely to experience spillover launch delay and welfare loss. On the other hand, if a low-income country caps its prices at the minimum price in a group of low-income countries, any launch delay and welfare loss experienced by the referring low-income country would be internalized. Assuming the firm has less chance of getting a high price in the referring country than in the other referenced countries, it would rationally delay launch in the referring country until launch has occurred in the potentially higher-priced, referenced countries. Thus, referencing is predicted to lead to delays in lower-price countries, regardless of who does the referencing, but effects are external to the regulator's calculus when higher-price countries reference lower-price countries.

countries. The effects of internal referencing depend on the details of implementation, in particular, whether drugs from older subclasses are included in the set of comparator products; the size of mark-ups, if any, whether the new drug can show superior efficacy or convenience; and any mark-ups for local production. Moreover, external referencing webs are complex, sometimes informal and extend transitively to unrelated countries. ¹⁴ The complexity of these relationships makes it impossible to derive a firm's optimal launch sequence and minimum reservation price for each drug-country pair. However, a clear prediction is that referencing of low-price countries by high-price countries will exacerbate launch delay and possibly raise prices in the low-price countries. ¹⁵

Of the countries in our data during our study period, France and Japan used both internal and external referencing. Canada used external referencing for "innovative" (first in class) drugs and internal referencing for "me-too" (late entrant) drugs. The Netherlands adopted a comprehensive but largely ineffective internal reference price reimbursement system in 1991; in 1996 it added price controls based on external referencing. Italy used a cost plus system until 1993; since then it has used variants of external referencing. Greece and Portugal also used externally referenced price controls for most of the period. Italy used a cost plus system until 1993; since then it has used variants of external referencing.

Free Pricing

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¹⁴ For example, if country A references country B, the effects in B depend on a variety of factors, including: the other countries in A's reference set; whether A references the mean, median or minimum of prices; whether an initial regulated price in A is updated if launch in B occurs after launch in A; and whether other countries reference B.

¹⁵ If regulators in low-price countries negotiate with a fixed drug budget constraint, they may offer a higher price in return for restricted distribution of a drug, e.g., available only from specialists or only for certain patient categories. ¹⁶ Many other countries, including most US health plans, the UK, Sweden, Italy, Germany and Spain used RP reimbursement for generically equivalent, off-patent compounds for at least part of our period. However, because these generic RP systems apply to off-patent drugs only, they are unlikely to affect launch decisions for new drugs, unless inter-brand effects are significant due to either competition or informal referencing.

¹⁷ The EU countries that used external reference pricing include Denmark (since April 1997, up to 10 EU countries excluding Greece and Italy), Greece (lowest in the EU), Ireland (lower of UK or the average in Denmark, France, Germany, the Netherlands, and the UK), Italy (average of up to 12 EU countries, must be on market for 4 countries and at least 2 with direct price controls), the Netherlands (since June 1996, average price in Belgium, France, Germany, and the UK), and Portugal (lowest in France, Italy, and Spain) (Burstall, 1998).

In the US, Germany and the UK, a new drug could be launched and reimbursed without regulatory approval of price, although other control mechanisms applied. In the US, multiple private health plans negotiate discounts in return for preferred formulary status, and Medicaid requires discounts off the price charged to private payers. These mechanisms may delay drug diffusion but not launch. In Germany, the reference price reimbursement system adopted in 1989 excluded new on-patent drugs, which could be launched and reimbursed without price approval, until 2005. The UK permits free pricing of individual drugs, subject to a rate of return constraint on each firm's portfolio of drugs. The renegotiation of this profit control scheme every five years usually involves international price comparisons and required cuts in UK prices. Since 1999 the National Institute of Clinical Excellence (NICE) has reviewed cost-effectiveness as a condition of reimbursement for most new drugs. This could slow new drug launch or lead to non-launch if the review were negative for drugs launched after 1999.

Parallel Trade

Parallel trade between EU states is legal under the Treaty of Rome. In our data, parallel imports are reported only in the high-price EU countries (Germany, the Netherlands, Sweden, and the UK) that also have incentives for pharmacists to substitute parallel imports for higher-priced, locally-sourced products. Parallel exporting countries are mainly the low-price EU countries (France, Spain, Italy, Greece, and Portugal) (Burstall, 1998). Parallel export risk may raise a firm's reservation price for launch in low-price countries within the EU, and hence contribute to non-launch if the low-price countries are unwilling to pay this reservation price. In practice, this may manifest as launch delay.

¹⁸ From 1993-2000 Germany had a national drug budget with physicians nominally at risk for budget overruns, although paybacks were not enforced. In 2005 on-patent drugs were added to the reference pricing system for reimbursement.

Given the heterogeneity of each regulatory type and each country's price regulatory system, categorizing countries by one or more regulatory indicator variables is unlikely to measure accurately even the direct effect of the regulation on prices and hence expected NPV. For example, although France, Japan and Canada all use both internal and external referencing, weighted price indexes for 1999 show Canadian and French prices roughly 30 percent lower than the US, whereas Japan's prices were over 20 percent higher than the US (Danzon and Furukawa, 2003). Moreover, an indicator that country A uses external referencing cannot capture the external effects that accrue as launch delay in the referenced, lower-price countries. Rather than use binary indicators, we therefore use average prices of competitor products as the most accurate measure of the net direct effect of each country's regulatory system on expected prices for new drugs. In countries where prices are not regulated, average prices of established products test for effects of price competition. We measure parallel trade risk and the indirect, spillover effects of regulation using a set of variables described below.

IV. Theoretical Model

If markets were separable and prices were unregulated, profit-maximizing firms would set prices independently for each country and would launch promptly after registration in all markets where the expected net present value of revenues exceeds country-specific fixed launch costs. With price regulation and potential spillovers, a necessary condition for launch of drug s in country j is that expected net present value of revenues exceeds country-specific costs plus any revenue loss in other countries due to spillovers:

$$E\{\int_{t=1}^{T} \left\{ \left[P_{sjt} \left(P_{bjt} \left(R \right), P_{gjt} \left(N_{gjt} \right), Y_{j}; P_{skt} \right) - C_{sjt} \right] \times Q_{sjt} \left(Q_{jt}, N_{bjt}, N_{gjt} \right) - \sum_{k \neq j} X_{jkt} \left(P_{sjt}, R_{kt}, I_{kt} \right) \right\} e^{-rt} dt > F_{jt} (H) \quad (1)$$

where E is the expectations operator. The expected price of product s, P_{sjt} , is assumed to depend on: P_{bjt} , the average price of competitor brand products, which depends on regulatory regime R; the average price of competitor generics P_{gjt} which depends on number of generic competitors N_{gjt} ; per capita income, Y, which may affect demand and regulatory stringency; and P_{skt} , the price of drug s in countries $k...K \ge 0$ that are referenced by j. Q_{sjt} is expected sales volume, which depends on aggregate sales in the class Q_{jt} , and on the number of brand and generic competitors N_{bjt} and N_{gjt} . Expected net revenue from launch also depends on $X_{jk} = \partial(P_kQ_k)/\partial P_{sj}$, the spillover effect of P_{sj} on revenues in country k, which either references to j or derives parallel imports I from j. $F_j = F_R + F_P$ is total fixed cost of drug launch, including registration cost F_R and price approval cost F_P , which may be lower if the launching corporation is homebased H; T is the duration of the economic life of the drug indexed by t; and r is the discount factor.

Equation 1 can be rewritten to derive the firm's reservation price for launch in country j. It is increasing in X_{jk} , the potential revenue loss in country k due to spillovers from country j $(\partial P_{sj}^{Ask}/\partial X_{jk}>0)$. Thus, external referencing and parallel importing that result in negative spillovers from P_j to higher-price countries can undermine incentives to launch in country j, even if the NPV of within-country sales is positive. Reservation price and launch probability are decreasing in expected market size $(\partial P_{sj}^{Ask}/\partial Q_{sj}<0)$, assuming some fixed costs of launch, F. If

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¹⁹ A potential concern is whether the average price of branded competitor products P_{bjt} is endogenous and depends on number of competitors (N_{bjt} and N_{gjt}), which may also be endogenous. We model P_{bjt} as independent of N_{bjt} because in regulated regimes price is set by regulation and existing firms do not generally lower their prices in response to new entry. Similarly in unregulated pharmaceutical markets, there is little evidence that entry leads to lower prices of existing products, plausibly because insured patients are not sensitive to full price. In the UK and Germany, patient co-payments are fixed per script, independent of price. In the US, entry may lead to increased discounting by firms to pharmaceutical benefit managers (PBMs), but our price data do not reflect these off-invoice discounts. Number of brand competitors is plausibly exogenous because of the 6-12 year lead time required for R&D and regulatory approval of new compounds; it was insignificant and dropped from the estimates reported here. Entry delay is shorter for generics, but is still several years from starting compound formulation to regulatory approval.

F is invariant across countries and compounds, firms are less likely to seek launch in country-compound pairs with low expected sales, due to low prices or small volume. Price regulation can reduce launch probabilities directly by reducing comparator prices P_{bjt} and hence expected prices for new drugs, or possibly by raising F due to costs of negotiation.

The regulator's reservation or maximum offer price depends on the regulatory regime. Under internal referencing, the regulator's offer depends directly on prices of substitute products $(\partial P_{sj}^{Offer}/\partial P_{bj} > 0)$. Under external referencing, although the regulatory formula is based on prices in comparator countries, prices of existing products provide an empirical proxy for achievable price levels under the referencing formula. Regulatory offers may be related to per capita income $(\partial P_{sj}^{Offer}/\partial Y_j \ge 0)$, given the political pressures on regulators to constrain health spending within budgets that are related to per capita income levels. Budget impact may lead regulators to offer lower prices for drugs with relatively large potential volume, Q_{sj} , other things equal $(\partial P_{sj}^{Offer}/\partial Q_{sj} \le 0)$. If local corporations are favored, $\partial P_{sj}^{Offer}/\partial H_j \ge 0$.

If P_{sj}^{Offer} - $P_{sj}^{Ask} \ge 0$ and a launch price can be agreed within this range, launch occurs. If not, negotiations may continue and launch may ultimately occur if either P_{sj}^{Offer} increases, P_{sj}^{Ask} falls, or some mechanism can be negotiated to bridge the difference, such as a price-volume offset. 20

In our data, we observe only the launch date and launch price conditional on launch, not the dates of registration approval or negotiation details. We therefore estimate reduced form equations for the launch hazard and launch price as a function of the determinants of the firm's ask price and the regulator's offer price. The reduced form launch hazard equation is:

$$h_{sit} = h\{P_{bit}, P_{git}, Q_{it}, X_{it}, N_{bit}, N_{git}, I_{it}, Y_{i}; P_{sKt}; H\}$$

$$(4)$$

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²⁰ In addition to internal and external referencing, France applies company-specific and therapeutic class spending limits that result in price cuts if volume sold exceeds target levels.

Measurement and predicted signs of variables are discussed below.

Because the bargaining range P^{Offer} - $P^{Ask} > 0$ also defines the range of launch price, the reduced form launch price equation includes the same variables as the launch hazard equation:

$$P_{sj} = f\{P_{bj}, P_{gj}, Q_j, X_j, I_j, Y_j; P_{sKt}; H\}$$
(5)

In theory, expected price and market size at global launch t_0 influence the decision to seek registration and hence the launch hazard, and hence could identify the launch hazard equation, whereas launch price depends on competitor price values only at launch. In practice, both prelaunch values and change over time of the expected price and quantity variables were insignificant in the launch equation, after controlling for contemporaneous values. Identification in the two-stage selection models therefore relies mainly on functional form, as discussed below.

We estimate separate models for new drugs launching in superior vs. inferior subclasses. Coefficient differences between these equations are expected due to dynamic competition and related factors. For example, a drug that is a late entrant in an old subclass with declining sales may not be worth launching unless it offers significant advantages over established products in the subclass or expects favorable treatment by local regulators or markets.

V. Data and Methods

Data

We use data from IMS Health's Midas database on drugs in 15 countries for 12 therapeutic classes, all of which experienced the launch of a new subclass shortly before or during our study period, 1992-2003. The data for each molecule include active ingredient, originator corporation(s) and marketing companies, pack description, launch date, therapeutic class, and quarterly data on outpatient sales at manufacturer prices (revenue in local currency)

and unit volume (IMS standard units)²¹ from 1Q 1992 through 4Q 2003. After the data were screened for internal consistency, revenue was adjusted for inflation using country-quarter-specific Producer Price Indexes available from the International Monetary Fund, with 2003 as the base year, and converted to US dollars using the average 2003 country-specific exchange rate. Brazil and Mexico sales were reported in our IMS data only in US dollars. Price per dose for each drug was calculated on a quarterly basis as the ratio of total revenues to standard units sold.²²

Of the 375 molecules in the dataset, 116 are classified as superior and 221 of their potential 1,740 drug-country launches had occurred prior to our study period; 259 are classified as inferior and 1,276 of their potential 3,885 drug-country launches had occurred prior to our period.²³ During our 12-year study period, we observe 885 of the 1,519 potential superior drug-country launches and 390 of the 2,609 potential inferior drug-country launches. For 91 country-molecule pairs, two distinct formulations (form 2-level, such as an oral solid and a liquid) of a molecule launched simultaneously, and in four country-molecule pairs three distinct formulations launched simultaneously, resulting in 1,367 country-molecule-product launches (946 superior and 421 inferior) in our study period.

Launch Estimation

²¹ The IMS standard unit is a proxy for a dose for each formulation e.g. one tablet or capsule, 5ml. for liquids. The IMS price data for the US do not reflect off-invoice discounts given by manufacturers to pharmacy benefit managers and HMOs; hence the US prices are upward biased for manufacturer net revenues.

²² We combined multiple form-3 level formulations (e.g. tablets and capsules, possibly of different strength) in a given country and quarter into a single observation and defined the price as the volume-weighted average price per unit. Identical forms that were launched by different co-marketing companies were also averaged.

²³ Our analysis includes only the first launch of each compound. We exclude follow-on indications or formulations because they typically face fewer registration requirements and pricing would be based on the price of existing formulations. We also focus on launch by an originator or licensee corporation, excluding the few cases of prior launch by copycat products. Five superior molecules and 20 inferior molecules were diffused to all our countries prior to our period. These are included as competitor products but are not in the sample of potential launches.

Because our data consist of quarterly observations, the launch hazard equation was modeled using a maximum likelihood discrete time implementation of a proportional hazards model based on complementary log-log regression, 24 which readily accommodates right censoring, late entry into the risk set, estimation of a flexible baseline hazard, and time-varying covariates. In the clog-log analysis, each drug was considered eligible for launch in all countries starting from its quarter of first launch in any country in our sample ("global launch"), and remained eligible until it launched. Thus each drug s in each country j contributes t_{sj} observations, the number of quarters from product s's global launch through either first launch in country j or 4Q 2003, the end of our study period. To account for the intra-molecule clustering associated with having multiple observations per drug resulting from potential launch in multiple countries, we used robust standard errors or molecule random effects.

The hazard of launch is $h_{sjt} = \Pr[\tau = t \mid \tau \ge t]$, i.e., the probability that drug s launches in country j in period t conditional on not having previously launched, where τ indicates the quarter of launch. Using a clog-log specification implies that

$$\log\left\{-\log(1-h_{sit})\right\} = \lambda(t) + \beta\Gamma_{sit} \tag{6}$$

where Γ_{sjt} is a vector of explanatory variables as outlined above. To facilitate tests of duration dependence, we specified $\lambda(t)$ to be quadratic in the number of quarters since global launch. The clog-log specification can be rearranged to yield an expression for the launch hazard:

$$h_{sit} = 1 - \exp\{-\exp(\lambda(t) + \beta \Gamma_{sit})\}$$
(7)

This in turn leads to a log likelihood function:

$$\ln L = \sum_{s=1}^{n} \sum_{i=1}^{m} \sum_{k=1}^{t_{sj}} \left\{ d_{sjk} \log(h_{sjk}) + \left(1 - d_{sjk}\right) \log(1 - h_{sjk}) \right\}$$
(8)

²⁴ A clog-log specification is preferred to logit in this context because of the clog-log's underlying assumption that the launch decision process is continuous (Allison, 1995).

where d is an indicator for whether launch occurs. To facilitate interpretation, we present marginal effects calculated at the regressors' mean values. The standard errors of the marginal effects were calculated using the delta method; see Bartus (2005) for more details.²⁵

There are two limitations of this basic specification that we address with alternate estimators. First, like other standard duration models, it assumes that the probability of failure goes to unity as time goes to infinity; however, some of the molecules in our sample might not meet drug approval requirements or would have limited market potential in some countries. We therefore also estimate a discrete-time split-population model with time-varying covariates (Schmidt and Witte, 1989; Jenkins, 1995). This specification allows for some empirically-estimated sample-wide proportion of drugs, c, never to launch.²⁶ The log likelihood function for this model is

$$\ln L = \sum_{s=1}^{n} \sum_{j=1}^{m} \sum_{k=1}^{t_{sj}} \left\{ d_{sjk} \log[(1-c)h_{sjk}] + (1-d_{sjk}) \log[c+(1-c)(1-h_{sjk})] \right\}$$
(9)

The second term of equation (9) accounts for the possibilities that drugs not observed to launch in the data either never launch, or launch after the end of the study period. Note that when c is zero, the split-population model log likelihood reverts to the standard clog-log model in equation (8).

A second limitation of standard duration models is that they do not account for unobserved heterogeneity. We suspect that there are time-invariant unobserved characteristics common to a molecule across countries that influence the probability of launch. Failure to

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 $^{^{25}}$ We calculated marginal effects for interactions of dummy variables X_1 and X_2 as $\{[F(X\beta|X_1=1,X_2=1) - F(X\beta|X_1=0,X_2=1)] - [F(X\beta|X_1=1,X_2=0) - F(X\beta|X_1=0,X_2=0)]\}$, where F is the inverse of the clog-log function, $1-\exp[-\exp(X\beta)]$. Other regressors were held at their means or at zero if they were related to X_1 or X_2 . Standard errors were calculated using the delta method.

²⁶ This feature is especially relevant for analysis of molecules in inferior subclasses, in which 116 of the 259 molecules did not experience a new launch during the study period (compared with two of the 116 molecules in superior subclasses).

account for unobserved heterogeneity in the estimation may lead to coefficient attenuation, and may overstate the degree of negative duration dependence and understate the degree of positive duration dependence (Heckman and Singer, 1984; Lancaster, 1990). The most straightforward way to address this issue is to augment the hazard specification with a term for the drug-level heterogeneity v_s :

$$h_{sit} = 1 - \exp\left\{-\exp\left(\lambda(t) + \beta\Gamma_{sit} + \nu_{s}\right)\right\}$$
(10)

We estimated v_s assuming a Normal distribution, as well as an empirically-derived finite discrete distribution. As results were robust and there are no *ex ante* reasons to prefer one, we report results for the Normal distribution.

Launch Price

We use ordinary least squares to model the log of launch price of drug *s* in country *j*, conditional on launching. We use molecule-clustered standard errors in the OLS estimation, and to account for unobserved molecule characteristics we also report results from a GLS random effects estimator. To account for the potential selection bias produced by the correlation between the propensity to launch and the launch price, we also estimate a Heckman selection model with a first-stage clog-log regression, which, as described above, is equivalent to a proportional hazards model of new drug launch.²⁷

Variable Definitions

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$$M_{sjt} = \frac{\phi \left[\Phi^{-1}(\hat{p}_{sjt})\right]}{\Phi[\hat{p}_{sit}]}$$

where $\phi[\cdot]$ is the standard Normal density function and $\Phi[\cdot]$ is the standard Normal distribution function. As a check of sensitivity we also estimated a full information maximum likelihood Heckman model based on a probit in the first stage that offers robust, clustered standard errors.

²⁷ The clog-log-based Heckman model is estimated following a two-step procedure that ensures consistent standard errors (Heckman, 1979). Following Lee (1983) and Greene (1992), the inverse Mills' ratio for drug s in country j and time t, M_{sjt} , is calculated using the predicted probability of launch \hat{p}_{sjt} from a clog-log regression as

Below we discuss the variables in our launch and price model specifications, grouped into: regulation/expected price, expected sales volume, spillovers, first mover advantage and timing, country of domicile, country and year effects, and product characteristics. We also explain where the price model specification is different from the launch model specification.

Regulation/Expected Price: We use the (log lagged) average price of competitor brand (originator and licensed) products in the same therapeutic class as a comprehensive measure of the direct effect of price regulation on expected price for a new drug. Average price of competitor products is the explicit regulatory benchmark in internal referencing regimes; it should also be a rough proxy for the net effect of external referencing formulae. In free-pricing countries, the average price of established products provides an estimate of the expected price of a new drug, assuming competition constrains similar products to have similar prices. Average prices for superior and inferior subclasses are distinguished, to test for differential effects within vs. between subclasses, as proxies for static vs. dynamic price competition in free pricing countries; in regulated markets, analogous effects operate via the regulator's choice of comparator products.

Expected Sales Volume: The (log lagged) total number of doses sold in the same therapeutic class as the new drug is included as a measure of expected volume.²⁹ The expected effect on launch hazard and price is uncertain *a priori*, depending on whether the firm's opportunity cost of delay dominates the regulator's concern over budget impact. It was insignificant and was dropped from the price equation to conserve degrees of freedom.

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²⁸ In theory, we could identify the effects of regulation on launch experience using difference-in-difference analysis applied to countries that adopted specific changes to their price regulatory regimes relative to those that did not. In practice, most countries experienced gradual evolution of regulatory regimes over the period, with too few clear-cut changes and too much between-country heterogeneity to permit difference-in-difference estimation for specific regulatory types.

²⁹ We tried distinguishing volume by subclass and including number of competitor firms, but these were not significant and were dropped.

Spillovers: To test for indirect effects in low-price countries of regulatory referencing by high-price countries, we included three count variables that measure the number of countries a molecule has already launched in, categorized by low-price EU countries (France, Italy, Spain, Portugal, Greece), high-price EU countries (Germany, the Netherlands, Sweden and the UK) and high-priced non-EU countries (the US, Japan and Canada). Our categorization of low- and highprice EU countries is supported by actual average prices (see below). These variables are also interacted with indicators for whether the potential launch is in a low- vs. high-price EU country. These interactions test whether spillover effects are greater for launches in low-price EU countries, which are referenced by and are the main sources of parallel trade to higher-price EU countries, and whether spillovers are greater within the EU than from non-EU to EU countries, as expected.

In the price equation, we include similar interactions, except that we use the Minimum Own Price in high-price EU, low-price EU and high-price non-EU countries, defined as the lowest price received for the molecule in any country where launch has already occurred, for each country group, rather than simple count variables for number of prior launches. Estimates using Maximum Own Price were similar to those reported here using Minimum Own Price. Both variables could not be included together due to collinearity.³⁰

We also include a dummy variable Any PI Share in Subclass, to test whether risk of competition from parallel import reduces the propensity to launch or reduces launch prices.³¹ Because the IMS data do not identify the country from which PIs originate, we cannot test directly whether propensity to be a parallel exporting country reduces the launch hazards in the

³⁰ We estimated specifications of the launch and price models that measured spillovers with either the number of countries previously launched in or minimum/maximum own prices. Number of prior launches was selected for the final launch model specification and minimum own price for the final price model specification because each yielded the better model fit and was more closely related to the dependent variable.

31 We estimated a model specification that included the average price of parallel imports, and it was not significant.

exporting country. Rather, the propensity to parallel export is subsumed in the country fixed effects.

<u>First Mover Advantage and Timing</u>: To test for first mover incentives for launch, the launch equation includes an indicator variable for quarters in which there are no molecules in the country-subclass. To test for first mover advantage on price, we include indicator variables for whether a launch was the first, second or third entrant in country-subclass.³²

A quadratic in years since global launch is included to control for the decline in incentives for launch with time lapsed since global launch, because patents run regardless of launch and compounds may undergo obsolescence due to entry of newer compounds. An indicator for molecules launched before 1990 controls for their relatively old age. An indicator for molecules launched since 1996 tests for effects of the EMEA regime. It is expected to be positive if the cost-reducing effects of the EMEA coordinated registration process outweighed the increased risk of spillovers. Molecules launched during 1990-1995 are the referent category.

Country of Domicile: Previous studies have found that new drugs launch more quickly in the home country of the originator firm, attributed to greater experience or favor with domestic regulatory agencies (Danzon, Wang and Wang, 2005; Kyle, 2006, 2007). Effects on launch prices have not been examined. We include dummy variables to test for differential effects of three categories of local corporations: Local Originator identifies a molecule's originator corporation launching in its country of domicile; Solo Licensee identifies a locally-domiciled, licensed corporation that launched the molecule in at least one country by itself; and Local Comarketer identifies a locally-domiciled, licensee corporation that launched together with another firm in its home country and did not launch alone in any country. The Solo Licensees are firms that have demonstrated ability to obtain drug registration on their own; thus their local expertise

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 $^{^{32}}$ The first mover advantage variables are based on the number of branded competitors by subclass, N_{bjt} .

should be positively associated with launch hazard, and possibly also with price. The Local Comarketers have not demonstrated registration expertise, hence they are predicted to have less effect on launch hazard than for either Originator or Solo Licensee firms; however, Local Comarketers may affect price. These categories are not mutually exclusive; for example, a molecule with a small Local Originator could also have a Solo Licensee or a Local Comarketer as a marketing partner, all from the same country.

<u>Country and Year Effects</u>: We include country fixed effects to capture other country-specific factors that may affect launch delay and launch prices (controlling for expected price, volume and per capita income), in particular, pure bureaucratic delay and parallel export risk. Germany is excluded as the referent country.

The dollar-euro/ECU exchange rate and the PPI are included in the price equation to control for exchange rate and indexing trends that could bias our dollar-denominated estimates of competitor prices and launch prices. Year effects were included in some specifications, but were generally insignificant and are not reported here.

<u>Product Characteristics</u>: The price equation includes product characteristics that affect price per dose, including pack size, pill strength (grams per unit), and indicator variables for specialized formulations (oral delayed and non-oral solids), with oral solids (basic tablets and capsules) as the referent formulation.

VI. Descriptives

Table 1 reports the total number of molecules ever launched, the number launched in our time period, and mean and median launch lags, by country and subclass. For the superior subclasses, Germany and the US (two free pricing countries) have the most molecules ever

launched (88 and 86) and launched during our period (72 each). Sweden, the US, the Netherlands, Germany, and the UK (all higher price, less-strictly regulated) also have the shortest median launch delay (17.4-18.7 months). Japan, Portugal and France (all price-regulated countries) have the fewest superior molecules (53, 62 and 69 respectively) and the longest mean launch lags (41, 31, and 37).

For the inferior subclasses, Japan leads in number of inferior molecules (158) ever launched, followed by Germany (131), and even Portugal (113) has more than the US (97). The number of inferior molecule launches during our period is highest in three regulated markets (Japan [43], Brazil [40] and Greece [47]) whereas most other countries have fewer than 27. Mean launch lags are generally much longer for inferior than for superior molecules. These differences in launch experience in the superior vs. inferior subclasses confirms that the older subclasses may be more heterogeneous, including some molecules that could not meet strict regulatory requirements and/or have limited sales potential in some markets.

Table 2 reports the mean number of manufacturers per molecule in 2003 by country, subclass and license type, to illustrate differences in market structure that may influence outcomes across countries. The expected number of originator/licensee firms per molecule is 1-2, assuming that an originator's profit-maximizing strategy is usually to launch alone or with at most one co-marketing partner. Consistent with this, the mean number of originator/licensee firms per superior molecule is 1.0 in the US, the UK, and the Netherlands, and only slightly higher in most other countries. However, licensees are more common in Italy (1.8), Spain (1.7) and Japan (1.3), suggesting that having a local co-marketing partner may be particularly valuable in these markets.

Parallel imports are found only in the four higher priced EU countries—Germany, the Netherlands, Sweden and the UK—and the majority of molecules in these countries have some PI presence by 2003. This concentration of PIs in a few countries may provide insufficient variation to estimate PI effects accurately, as noted below.

Unbranded generics are more numerous for molecules in older subclasses, which is unsurprising because these are generics that enter after patent expiry and compete on price.

Unbranded generics are most common in the US. By contrast, Other Brands, which includes branded generics and copy products that compete mainly on brand, are most numerous in Germany, Japan, and Brazil.

Table 3 reports mean prices by country-subclass-license type. Each mean price is the unweighted mean of prices for all products in that country-subclass-license type. These means thus reflect differences in molecules and formulations, in addition to price differences for similar products, and hence are not valid indexes of cross-national price differences for a standardized basket of drugs. However, these unweighted mean prices provide a rough measure of benchmark competitor prices used by regulators and firms in forming price expectations, except that in practice regulators and firms may focus on a narrower subset of close substitute products within these broad subclass averages.

These unweighted mean prices show that, for originator/licensee superior products in the EU, the price-regulated regimes (France, Spain, Portugal, Greece, Italy) have relative low prices. Thus, we classify them as "low-price EU markets." The countries with freer pricing, reimbursement regulation and/or late adoption of price regulation (Germany, the UK, the Netherlands, Sweden) have higher mean prices, and we classify them as "high-price EU

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³³ Danzon and Furukawa (2003, 2008) report weighted price indexes, based on standardized market baskets, for originator and generic products in 1999 and 2005

markets." The US has the highest prices, followed closely by Canada; we also classify
Switzerland and Japan as "non-EU high-price markets" based on other price index comparisons
(Danzon and Furukawa, 2003), although Japan's unweighted mean prices are quite low in Table
3. Originator prices are lower in inferior than superior subclasses, and country rankings are
similar but with smaller differentials. Other Brand prices are generally higher than for
Unbranded Generics, as expected.³⁴ For unbranded generics, the relatively high mean US price
is surprising and reflects its larger number of superior products, whereas volume-weighted price
indexes for standardized products show generic prices relatively low in the US (Danzon and
Furukawa, 2003). PI prices generally fall between generic and originator prices, as expected;
again, these differentials are not based on a standardized product mix and are not intended to
provide an accurate measure of originator/PI price differentials or the impact of PIs on firms or
consumers/payers.

Table 4 shows, for each country, the total number of molecules ever launched, and the numbers that were launched by a Local Originator, a Solo Licensee or a Local Co-marketer corporation. Also reported is the average number of countries in which each country's originated molecules were launched. Using launch by a Local Originator as a proxy to identify molecules originated in each country, for superior drugs, the US had the largest number (30) of launches, followed by Japan (17), and the UK, France and Switzerland (11 each). All other countries had 0 to 4 launches. Drugs originating in Japan and Spain diffused to dramatically fewer countries on average (4) than drugs originating in other countries (10.6-12.8). By contrast, Japan had the largest number of inferior molecules (45), and the US had relatively fewer (29), similar to Germany and Switzerland (25-26). All countries other than the US had a larger number of

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³⁴ Some of the PI, Unbranded Generic and Other Brand means include very few products. Because most molecules have at least one originator/licensee product, these means are less robust than the originator means.

inferior molecules than superior molecules. Analyzing the features of these other countries' systems that favor older drugs is an important issue, but it is beyond the scope of this paper. In general, inferior molecules diffuse to fewer countries than superior molecules.

Means and standard deviations for variables in the launch and price equations are reported in Appendix Tables A1 and A2.

VII. Determinants of Launch

Table 5 reports coefficients and marginal effects from our basic launch specification, using clog-log estimates with either robust, clustered standard errors or molecule-level Normal random effects to control for unobserved molecule characteristics.³⁵ Our discussion focuses on the random effects estimates for the superior subclasses, noting differences for the OLS estimates and for the inferior subclasses where relevant.

Domestic Prices: Direct Effect of Price Regulation

Launch hazards of superior products are significantly related to mean prices of competitor brand products in the superior subclass: a 10 percent increase in competitor prices is associated with a 0.054 percentage point increase in the launch hazard, which seems reasonable given the 3.8% average launch hazard for superiors molecules per quarter.³⁶ Thus to the extent that regulation reduces prices, it reduces incentives to launch. These estimates may underestimate the magnitude of regulatory effects, if our measure of mean price, based on all originator and licensed products in the subclass, includes some that are not in comparator group

³⁵ Parallel specifications using a split population clog-log model and a discrete-distribution random effects clog-log model with were also estimated. Results were generally consistent with those reported here.

³⁶ Because average prices of competitor products vary mainly by class and country, we estimated specifications without controls for country and class fixed effects, to show maximum potential impact of competitor prices. In fact, results for the price variables were either insignificant or negative, suggesting uncontrolled heterogeneity. We therefore report here the specifications that include country and class fixed effects, such that identification is based on variation within country-class over time. Results for other variables are robust to excluding class fixed effects.

of close substitutes used by regulators. For the inferior subclass, launch hazards are positively related to prices in the own subclass but the effect is not significant. Cross-class price effects are insignificant, indicating that regulatory benchmarking and/or competitive constraints operate primarily within rather than between subclasses, and that dynamic competition is driven by product characteristics other than price. Similarly, the effects of number of generic competitors are negative but statistically insignificant, providing further evidence that availability of older, cheaper generic substitutes is not a significant deterrent to the launch of new brand products, even in older subclasses where generics are more numerous. This may reflect the fact that late launches of inferior products occur mainly in countries where most generics are branded generics that do not compete aggressively price, unlike in the US.

Unit volume for the therapeutic class³⁷—whether measured contemporaneously, at global launch or as a growth rate —was not significantly positive for superior or inferior products, even when country and class effects were omitted. Thus, on average potential market size is not a significant launch determinant for this sample of drugs-countries, possibly because regulators constrain prices more stringently for compounds with large potential sales, which offsets incentives of firms to launch these drugs more promptly.³⁸

Launch Timing and Sequence

For both superior and inferior products, launch hazards appear first to decrease then increase with time since global launch, reaching a minimum at 13.5 yrs from global launch for

³⁷ Volume is measured here as total volume in the relevant inferior and superior subclasses; separate subclass measures were also insignificant.

³⁸ The insignificant effects of volume on launch of new products found here contrasts with significant positive effects in Danzon, Wang and Wang (2005). These different findings may reflect differences in sample countries and drugs, in addition to our use here of more detailed measures of country-class prices and other characteristics. Our finding here of no significant volume effects on average does not rule out the possibility that small expected sales volume is an issue for specific drugs in specific countries.

superior drugs and 49.0 years for inferiors.³⁹ These average quadratic specifications reflect the diverse launch patterns in Table 1, which shows a fairly clear pattern for superior drugs, with median launch lags of less than one year in the high-price EU countries and the US, followed by launch within the second year for all other countries except France, Portugal and Japan, where launch typically occurs later. For inferior drugs, median launch lags are much longer, which the other evidence suggests is only partly explained by the fact that we observe them later in their life-cycle.

Inferior drugs launched before 1990 are more likely to launch than later entrants in the same subclass, possibly due to accumulated global brand capital of these earlier entrants and despite their presumably shorter remaining patent life.⁴⁰ The poor launch performance of late entrants in older subclasses is consistent with the hypothesis that dynamic competition from the newer subclass disadvantages late entrants to an inferior subclass.

The coefficient on the post-1996 global launch indicator is negative but insignificant for superior drugs. Taken at face value this suggests that on average the EMEA process has not affected speed of diffusion, possibly because price approval is the rate-limiting regulatory hurdle and any cost-reducing effect of accelerated approval is offset by increased risk of spillovers. Such inferences are tentative, however, because our analysis period is too early to observe the full effects of the EMEA, which focused initially on biologics and truly innovative drugs. *Indirect Regulatory Effects: Cross-National Spillovers*

The evidence is strongly consistent with the hypothesis that launch in low-priced EU countries is adversely affected by the risk of spillover to higher-price EU countries through external referencing and possibly PI risk. For superior drugs, the coefficients on number of

³⁹ The minimums from the split population estimates are 10.9 years for superior and 50.3 years for inferiors.

⁴⁰ Patent expiry is less critical to expected sales in countries with few generics or primarily branded generics that do not compete aggressively on price. This includes all the low-price EU countries during our time period.

countries in which launch has already occurred are all positive and significant, with the exception of prior launch in the three lowest price EU countries, Spain, Portugal and Greece. The marginal effect is largest for prior launch in the UK or Germany, at 0.030, and is 0.025 for prior launch in Sweden or the Netherlands. By contrast the marginal effect of prior launch in Italy or France is 0.013, and for high price non-EU countries the marginal effect is only 0.009. ⁴¹ This pattern confirms that prior launch in high-price EU countries is associated with a larger increase in current launch hazard than is prior launch in other countries, as expected, if, to avoid negative spillovers, firms delay launch in low-price EU countries until launch has occurred in higher-price EU countries. Moreover, because Spain, Portugal and Greece reference the lowest prices in a group of relatively low-price countries, including France and Italy, a firm's optimal launch strategy plausibly leads to launching last in these three countries, after higher prices have been established in the countries that they reference. For inferior drugs, marginal effects are much smaller and generally insignificant, consistent with other evidence that these late launching drugs in older subclasses reflect either delayed launches of previously launched drugs or atypical drugs.

To explore further the spillover effects for superior drugs, we estimated a specification that includes counts of prior launches in high-price EU, low-price EU and other high-price countries (Canada, Japan, Switzerland and the US), together with interactions between these launch counts and indicators for whether the current observation is a low- or a high-price EU country. Marginal effects of these interactions are reported in Table 6. The marginal effect of a prior launch (from zero to one) in a high-price EU country on launch in a low-price EU country is 0.0018, whereas the effect of a prior launch in another low-price EU country is only 0.0005,

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⁴¹ The p-values for Wald tests comparing these marginal effects are as follows: UK/Germany vs. Italy/France, p=0.019; UK/Germany vs. high-price non-EU countries, p=0.009; Sweden/Netherlands vs. Italy/France, p=0.062; Sweden/Netherlands vs. high-price non-EU countries, p=0.003.

and the difference is statistically significant. Similarly, the marginal effect on launch in a low-price EU country is greater from a prior launch in a high-price EU country than from a launch in a high-price non-EU country, as expected, because referencing and parallel trade within the EU is only to EU countries. This evidence is thus consistent with the hypothesis that the observed pattern reflects spillovers, not simply some unobserved factor that leads to correlation in launch timing across countries.

Parallel import presence in the class is not associated with launch hazard in the importing country, after controlling for country fixed effects. This may simply reflect the high correlation between the PI indicator and the country indicators for the four countries with PI presence—Germany, Sweden, the Netherlands and the UK—and the high PIs presence across classes in those countries. It is also more likely that parallel trade risk leads primarily to non-launch or launch delay in the parallel export countries, not in the importing countries. We cannot measure this effect in the exporting countries because our data do not report PI country of origin; hence it is subsumed in country effects. 42

Launch by a Local Corporation

Launch is more likely for both superior and inferior molecules from firms that are domiciled domestically. For superior drugs, the marginal effect of launch by a Local Originator is 0.13-0.18, compared with 0.03-0.04 for Solo Licensees and 0.02-0.03 for Local Co-marketers. This ranking confirms that compounds that are originated by local firms have significantly greater local advantage than compounds that are simply represented by local licensee firms, ⁴³

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⁴² We also tried including the average price of PIs, but this was not significant after controlling for country fixed effects. It is possible that parallel trade has modest incremental effects compared to external referencing, because external referencing reduces prices on all units whereas parallel trade affects only the fraction of sales that are imported.

⁴³ P<0.05 for comparisons of marginal effects between Local Originators and either Solo Licensees or Local Comarketers.

although even the latter accelerate launch somewhat relative to having no local connection. Firms that are local originators by definition have local R&D activities and hence are likely to be larger, have greater regulatory expertise and be viewed as more valuable to the local economy than firms that are just co-marketers, with Solo Licensees in the middle. To test whether these local corporation effects differed across countries, we combined the three indicators into a single Any Local Corporation indicator and estimated a specification with interactions between Any Local Corporation and country fixed effects. 44 Marginal effects of these interactions are reported in Table 7. The effect of the Any Local Corporation indicator is insignificant for the referent, Germany, and for most other countries. The exceptions are France, Italy, Spain, Switzerland and Japan, where launch by a local corporation has a significant positive effect for superior drugs. Overall, this evidence indicates that the large positive mean effect of involvement by a local corporation on launch hazard, observed in our baseline specifications and in previous studies, is confined to a few countries, with no significant domestic advantage in most countries. This suggests that the local corporation advantage reflects primarily industrial policy to support local firms in certain countries, not that local firms in general have greater expertise in dealing with local regulatory systems in all countries.

To shed further light on cross-national differences, Appendix Table 3 reports the number of launches and mean and median launch lags for launches by Local Originators, Solo Licensees and Local Co-Marketers compared with launches by non-local corporations, by country. For Japan, the mean launch delay for superior drugs is 1.8 months with a Local Originator, compared with 53.0 months with no local firm, and 59 months with a Solo Licensee and 49 months with a Local Co-marketer. The Local Originator advantage could partially reflect the fact that clinical

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⁴⁴ We tried to estimate a specification with interactions between each of the three local corporation indicators and country fixed effects. The clog-log coefficients could not be estimated due to small sample sizes for several countries.

trials for these drugs were probably done on Japanese subjects, as required for registration in Japan. However, the fact that these Japan-originated drugs on average diffused to only 4 countries besides Japan indicates that their home advantage is not just speed due to local trials; rather, it reflects a tendency for Japan to approve locally-originated drugs that have limited potential in other countries. In France, Italy and Spain, there is some evidence that launch lags are shorter if a Local Co-marketer is present than if the drug is launched by a Solo Licensee. This is consistent with the hypothesis that Local Co-marketers are added only where they are expected to bring advantage to foreign originators in dealing with regulators; it is also consistent with anecdotal evidence that regulators in these countries traditionally encouraged such co-marketing deals as a form of industrial policy to support local firms that lacked strong R&D capabilities. *Country Fixed Effects*

For superior drugs, compared to Germany, the referent country, other country effects are all negative. Marginal effects are smallest for the UK (-0.011) and other relatively free pricing countries; marginal effects are largest for Japan (-0.044), reflecting its unique registration requirements; and the major EU parallel export countries (Spain, France, Greece, Portugal and Italy) are all significantly negative, as are several other countries. For inferior drugs, none of the country marginal effects is significant in the basic clog-log estimation, possibly due to within-class heterogeneity. However, for the US, the clog-log coefficient, the RE marginal effect and the split population estimates are all negative, implying that late-launching inferior drugs are less likely to launch in the US than in other countries.

Split Population and Random Effects Estimators

The split population and random effects estimates are generally consistent with the cloglog estimates. However, the split population estimates (shown in Appendix Table A4) imply that the probability of never launching is highly significant for 27.7 percent of inferior molecule-country pairs, compared with only 4.9 percent of the superior molecules. This provides further evidence that certain molecules, especially late entrants in inferior classes, are not marketable in certain countries. Whether this reflects inability to meet regulatory or market requirements cannot be determined with our data.

VIII. Determinants of Launch Price

Table 8 reports the determinants of (log) launch price, as estimated using OLS with robust, clustered standard errors, and GLS random effects to control for unobserved molecule heterogeneity. The model specifications reported here include country fixed effects. Year fixed effects were also included, to control for any bias in our inflation and exchange rate adjusters, but these coefficients were generally insignificant and are not reported. Therapeutic class effects were omitted because they are highly collinear with the variables measuring competitor prices and order of entry within class. Our discussion focuses on estimates that include GDP per capita; excluding GDP changes primarily the country fixed effects, as reported below. 46

Competitor Prices

For both superior and inferior products, launch prices are significantly positively related to prices of competitor products in the same subclass (elasticity of 0.12 for superiors and 0.17 for

⁴⁵ The estimated percent never launching is significantly higher once we control for Local Originator Corporation, consistent with the hypothesis that some countries approve locally-originated drugs that have limited diffusion potential outside their home country.

⁴⁶ Because our observed launch prices are conditional on launch, we estimated both a two-stage Heckman selection

⁴⁶ Because our observed launch prices are conditional on launch, we estimated both a two-stage Heckman selection correction model that includes as a regressor the inverse Mills' ratio from a clog-log first-stage hazard equation and a traditional FIML Heckman model based on a probit first-stage equation. As shown in Appendix Table A5, the coefficients on the inverse Mills' ratios are larger for the inferior drugs, but are significant only for the superior drugs (and only in the two-stage models); otherwise results are generally similar to the conditional estimates. These results are broadly consistent with the finding from the split population estimates that at least some molecules do not have global launch potential. Because the first-stage launch equation is identified primarily off functional form, we focus our discussion on the conditional estimates in Table 8 and do not attempt to draw inferences about differences between conditional and unconditional estimates.

inferiors). The cross-subclass elasticity of inferior drug prices on launch price of new superior drugs is also positive but smaller (0.08). This confirms the earlier evidence, that launch probabilities are influenced mainly by prices of established products within subclass, implying that dynamic competition between subclasses is based on non-price product attributes.⁴⁷ Generic prices have no significant effect on launch prices of new superior brands, suggesting weak price competition between new brands and old generics. Launch prices of new inferior products are affected negatively by generic prices in the class (elasticity -0.09), which may reflect a selection effect: late entrants in inferior subclasses launch only if they expect to receive high prices relative to competing generics.

Launch Timing and Sequence

Controlling for inflation and exchange rates, launch prices for superior products decline with time elapsed since global launch. The estimate of the net impact of time since global launch from the random effects models is -3.8% (p=0.05). This contradicts the hypothesis that manufacturers would rationally delay launch in the expectation of receiving a higher price purely due to delay.

For superior products, there is no evidence of first-mover advantage in prices, although second and third entrants do receive higher prices relative to later entrants in a class. For inferior products, the first or second entrants in the subclass appear to receive a price premium relative to other inferior drugs. This conclusion is tentative, however, because it is based on a very small number of inferior subclasses for which first and second launches occur in our time period.

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⁴⁷ When therapeutic class indicators are included, they are significant for superior drugs in all 11 classes, compared to anti-hypertensives, the referent category. For inferior drugs, class effects are significant for only 4 classes. After including the class FEs, such that identification is within country-class-year, the competitor price variable for superiors becomes insignificant, and the coefficient for inferiors is significant, consistent with greater within-class heterogeneity for this subclass.

⁴⁸ The net impact of time since global launch (a quadratic) was calculated as $\beta_{tsgl} + 2\beta_{tsgl^2}\bar{t}$, where the mean value of time is 2.16 years.

Cross-national Spillovers

For both superior and inferior products, launch prices are positively related to the lowest price previously received in other high-price EU countries, whereas effects of launch in low-price EU countries is insignificant. The minimum own price received in non-EU countries is significantly positive for superior molecules, but insignificant for inferior molecules, possibly because they are less likely to launch in high-price non-EU countries such as the US and Canada.

We also estimated equations with interactions to test the hypothesis that spillovers to low-price EU countries are largest from high-price EU countries. Marginal effects of the interactions are reported in Table 9, which parallels Table 6 for the launch model in structure and results. We find that the effect on launch price in a low-price EU country is larger for a 10% increase in a drug's minimum own price in high-price EU countries than from a 10% increase in minimum own price in other low-price EU countries. Specifically, the difference in the marginal effects in the minimum own price elasticity based on the OLS model is 0.39, and based on the RE model is 0.26. Similarly, the effect on launch price in a low-price EU country is substantially larger for a 10% increase in a drug's minimum own price in high-price EU countries than it is for a 10% increase in minimum own price in high-price non-EU countries. The differences in the own price elasticity are 0.37 from the OLS model and 0.24 for the RE model, consistent with the hypothesis that launching first in high-price EU markets can influence prices in low-price EU markets. This evidence from launch prices thus validates that launch delay in low-price markets may ultimately yield higher prices in these markets through spillovers from higher-price markets, in addition to avoiding contamination of those high-price markets.

The indicator for PI presence is insignificant for superior drugs, but significant and negative for inferior drugs, indicating that PI presence reduces launch prices mainly for late entrants in older subclasses.

Local Corporations

For superior drugs, launch by a Local Originator is positively associated with launch price, but the effect is not significant. Estimated effects of both types of licensees are negative but insignificant. For inferior drugs, the average effect of launch by a Local Originator is significantly negative, which may reflect the fact that these late-launching inferior drugs have relatively weak characteristics, relative to established drugs in the subclass. The fact that the RE effects are smaller than the OLS effects is consistent with this. Launch by a Local Licensee is negative but insignificant. Tests for country-specific differences in local corporation effects (results not shown) are generally insignificant. This lack of evidence of a price premium for drugs launched by local firms, despite a significant advantage in the launch equation for a subset of countries, suggests that the launch advantage reflects favoring by the registration authorities rather than by pricing authorities. However, conclusions are tentative due to small sample sizes.

The US dollar per euro/ECU exchange rate, which declined from a high of 1.38 in 1992 to a low of 0.83 in 2000, is insignificant for superior products but large and significant for inferior products. This suggests that cross-national differences in launch prices for the more broadly diffused superior products were constrained by exchange rates, whereas launch prices for inferior products, which were less likely to launch in the US, were priced independently of the USD/Euro exchange rate. ⁴⁹ For superior drugs, the country-specific Producer Price Index

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⁴⁹ If manufacturers set launch prices for superior drugs to reflect changes in the USD-euro/ECU exchange rate, presumably because they were sensitive to referencing and PIs, the exchange rate variable would be insignificant.

(our price deflator) is significantly negative in the OLS specification, suggesting that launch prices of drugs on average have not kept pace with economy-wide inflation, but this conclusion is tentative because the RE estimate is insignificant.

Product Characteristics

Product characteristics have expected effects. Price per unit is significantly negatively related to pack size, particularly for pack size over 100 units, possibly indicating economies of scale in packaging and/or the competitive use of large packs to give discounts to pharmacies in countries such as the US, that permit pharmacists to dispense from large packs. Price is unrelated to strength (grams of active ingredient per unit) in the RE estimates, suggesting that the positive coefficient in the OLS results for superiors reflects between- rather than within-molecule effects. Compared to the oral solid formulations (the omitted category), price per dose is significantly higher for injectable and non-oral forms (liquids, creams etc).

Country Fixed Effects

The country fixed effects in Table 8 are dramatically different depending on whether GDP per capita is included. Based on the RE estimates, without GDP controls, prices for superior molecules in the US and Japan are significantly higher than in Germany; prices in Switzerland and Canada are higher but insignificant, whereas prices in Brazil, Mexico and all other EU countries are lower, except the Netherlands and Sweden which are similar to Germany. However, after controlling for GDP per capita, Brazil, Mexico, Spain, Portugal and Greece have prices significantly higher, and Switzerland, the UK and Sweden have prices significantly lower than Germany.

These international spillovers were less significant for inferior drugs, however. If manufacturers priced inferior

drugs based on local prices that were flat or falling in ECUs, the prices would be rising over time in USD, due to USD devaluation, thus explaining the positive and significant effect of exchange rate.

Although these are not pure hedonic country effects, taken at face value they imply that the prevailing spread in drug prices across EU countries is compressed relative to the counterfactual of differentials based on per capita income. This may explain at least in part why Spain, Portugal and Greece, the lowest-price EU countries in our sample, adopt more stringent price controls than the higher-price northern countries. However, these low-price EU countries appear to be constrained in their ability to keep price differentials in line with income differentials, in part due to spillovers that result from external referencing by and parallel importing to higher-income countries. The evidence here confirms that the resulting interconnectedness across countries contributes to delay or non-launch of new drugs in these low-price EU countries, as firms seek to avoid spillovers to prices in higher-price EU countries.

IX. Conclusions

Our findings demonstrate that launch timing and launch prices of new drugs are related to prices of established products in the same subclass, with smaller cross-subclass effects. This implies that, to the extent that price regulation reduces price levels, it contributes directly to the longer average launch delays observed in low-price countries. Our estimates suggest that the magnitude of these direct effects is quite small, although downward bias due to measurement error is possible. Welfare conclusions are ambiguous, assuming that regulators weigh the benefits of lower prices against any welfare loss from reduced access to new drugs for their citizens.

We also find significant evidence suggesting that regulatory referencing to lower foreign prices and parallel trade create incentives for manufacturers to delay launch in low-price countries until higher prices have been established in other countries. Consistent with such

strategies, launch in higher-price EU countries is associated with increased launch hazard in lower-price EU countries, and launch prices in low-price EU countries are directly related to prior launch prices in high-price EU markets. This evidence, that spillover effects are greatest from high-price EU to low-price EU countries, in both launch and price models, supports the hypothesis that they are attributable to referencing and parallel trade, not to other unmeasured factors that may lead to closely sequenced launches across countries. To the extent that referencing or parallel importation by higher-price countries leads to longer launch delay or higher prices in lower-price countries than would otherwise occur, a welfare loss is imposed on low-priced countries by the higher-price countries that adopt these regulatory strategies.

Although these low-price countries regulate their drug prices, their low drug prices also reflect their relatively low per capita income. In fact, despite price regulation, Spain, Portugal and Greece had relatively high drug prices given their income, whereas high-price EU countries had lower drug prices relative to their per capita GDP. How much external referencing and parallel trade have contributed to this convergence of pharmaceutical prices relative to GDP among EU countries is beyond the scope of this paper. Both theory and the evidence here suggest that parallel trade is less important than external referencing. We do find that the parallel import threat reduces launch prices of late-launching inferior products, and that country fixed effects are negative for countries that are significant parallel exporters. This rather weak evidence on effects of parallel trade may simply reflect measurement challenges, in particular, the concentration of parallel imports in four of our countries and lack of data on country of export for parallel imports.

Although prices of new drugs are constrained by prices of established drugs, either by regulation or competition, such effects occur primarily within subclasses, with weaker

constraints from older, cheaper drugs. Late entrants in older subclasses diffuse less broadly than new drugs in newer subclasses, and these late inferior launches are at lower prices, linked to the lower prices in these older subclasses. This evidence suggests significant dynamic competition on non-price product attributes in pharmaceutical markets, even though price competition and regulatory referencing are mainly within subclasses.

Our analysis indicates that the positive effects of launch by a local corporation found in previous studies are confined in our sample to a few countries—France, Italy, Spain, Switzerland and Japan—and appear to reflect primarily local registration advantage for drugs originated by local companies. We find weaker evidence of benefits from use of local licensees as comarketing partners, particularly in France, Italy and Spain. The fact that these local corporation advantages are confined to specific countries suggests that they reflect industrial policy strategies favoring local firms in these countries, rather than a more general local firm advantage based on their having expertise or familiarity in dealing with local regulators.

This evidence, that policies of external referencing (and, with weaker evidence, parallel importing) impose an external cost on the referenced or exporting countries, is based on the EU, where such policies already exist. However it has important implications for the US debate over drug price controls through external referencing and drug importation. Theory suggests that the launch lag externality will be greater if referenced countries are small and low price, compared to a much larger, higher-price referencing or importing country. Because the US has both higher brand prices and much higher total volume than most potential reference or exporting countries, the impact on these countries if the US were to adopt referencing or importing would potentially be much larger than the EU effects documented here.

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Table 1. Launch and Molecule Count and Mean and Median Launch Delay by Country and Subclass

		Superior Su	bclasses		Inferior Subclasses					
0 - 11	Mala a la a	Landon	Mean Launch	Median Launch	Mala a la a	Landon	Mean Launch	Median Launch		
Country	Molecules	Launches	Delay	Delay	Molecules	Launches	Delay	Delay		
High-Price E	ı									
Germany	88	72	18.5	9.5	131	18	30.4	17.5		
UK	80	58	18.7	6.5	116	24	69.2	41.5		
Netherlands	73	56	18.1	10	112	21	43.7	15		
Sweden	77	62	17.4	7	82	19	49.9	18		
Low-Price El	J Countries									
France	69	53	30.9	29	105	19	87.6	59		
Greece	72	55	30.1	22	107	37	116.8	54		
Italy	76	61	24.8	21	127	26	74.7	48.5		
Portugal	62	48	37.0	33.5	113	26	85.8	67		
Spain	76	62	28.1	21	112	23	43.9	31		
High-Price no	on-EU Coun	tries								
Canada	73	62	25.6	16	98	22	91.9	66.5		
Japan	53	42	41.0	40	158	43	63.3	28		
Switzerland	78	63	23.9	18	111	21	55.5	47		
USA	86	72	17.9	8	97	23	86.4	62		
Low-Price no	n-EU Count	ries								
Brazil	71	60	31.2	20.5	92	40	107.1	90.5		
Mexico	72	59	28.6	17	105	28	84.0	55		

Note: Launch delays measured in months

Note: Sample includes all molecules present and new launches occurring in our data during 1992-2003

Note: Launch delays are calculated only for country launches that occurred during 1992-2003 (regardless of when the global launch occurred)

Table 2. Mean Number of Manufacturers per Molecule by Country, Subclass and License Type in 2003

Superior Subclasses Inferior Subclasses Originator Originator / Unbranded Parallel Other Unbranded Parallel Other Licensee Molecules Country Licensee Generic **Brand** Molecules Generic **Brand** Import **Import High-Price EU Countries** 1.6 4.2 1.3 1.3 3.0 3.3 114 Germany 1.2 86 3.0 UK 1.0 0.3 0.1 0.0 79 1.1 1.0 0.3 1.1 103 Netherlands 1.0 5.5 0.0 1.1 0.1 97 0.0 68 0.0 6.1 Sweden 1.1 0.6 1.2 0.1 1.1 1.4 0.2 74 73 0.5 **Low-Price EU Countries** N/A 0.0 1.2 2.0 N/A 0.4 98 France 1.2 0.6 66 Greece 1.1 0.1 N/A 1.4 70 1.0 0.1 N/A 2.1 93 1.8 0.5 0.6 1.4 N/A 2.0 105 Italy N/A 74 1.0 Portugal 1.3 N/A 1.1 61 1.1 N/A 8.0 101 1.1 0.4 Spain 1.7 2.2 1.4 N/A 1.1 73 0.9 N/A 1.3 100 **High-Price non-EU Countries** Canada 1.1 0.4 N/A 8.0 69 1.1 2.0 N/A 1.7 85 1.3 0.4 N/A 1.5 52 1.5 0.3 N/A 3.3 153 Japan 1.1 N/A 0.2 77 1.1 0.2 N/A 0.5 93 Switzerland 0.1 1.0 0.0 USA 1.4 N/A 85 1.2 7.0 N/A 0.3 91 **Low-Price non-EU Countries** 2.1 3.0 Brazil 1.2 1.2 N/A 68 1.1 1.4 N/A 84 1.1 Mexico 1.1 0.3 N/A 0.9 70 0.7 N/A 1.9 90

Note: Sample includes all molecules present in IMS dataset in 2003

Table 3. Mean Price per Molecule by Country, Subclass and License Type in 2003

		Superior Su		Inferior Subclasses				
	Originator				Originator			
	/	Unbranded	Parallel	Other	/	Unbranded	Parallel	Other
Country	Licensee	Generic	Import	Brand	Licensee	Generic	Import	Brand
High-Price El	U Countries							
Germany	30.31	0.49	9.23	0.94	2.56	0.32	0.33	0.75
UK	13.26	0.64	1.36	0.61	6.18	0.93	0.46	1.27
Netherlands	35.53	0.05	2.22	7.75	10.79	0.45	0.61	0.72
Sweden	24.41	0.34	1.79	0.45	13.08	0.26	0.42	0.98
Low-Price EU	J Countries							
France	6.52	0.41	N/A	0.89	0.28	0.39	N/A	0.75
Greece	12.84	1.11	N/A	0.56	6.26	0.45	N/A	0.80
Italy	2.24	0.41	N/A	0.77	0.30	0.31	N/A	0.61
Portugal	1.83	0.44	N/A	0.78	0.33	0.20	N/A	0.34
Spain	6.13	0.33	N/A	1.06	0.28	0.63	N/A	0.69
High-Price no	on-EU Coun	tries						
Canada	49.30	0.57	N/A	0.60	1.03	0.93	N/A	0.57
Japan	12.27	0.44	N/A	0.73	2.62	1.26	N/A	0.59
Switzerland	34.64	0.81	N/A	0.76	5.81	0.37	N/A	1.43
USA	52.70	0.90	N/A	14.22	11.05	1.43	N/A	12.72
Low-Price no	n-EU Coun	tries						
Brazil	2.34	0.39	N/A	0.95	4.21	0.21	N/A	0.35
Mexico	18.34	0.69	N/A	1.65	5.84	0.26	N/A	0.53

Note: All prices are ex manufacturer prices per standard unit in 2003 US Dollars

Note: Sample includes all molecules present in our dataset in 2003

Table 4. Number of Launches and Mean Number of Other Country Launches by Country, Subclass and Domestic Status of Corporations

Molecules in Superior Subclasses

Molecules in Inferior Subclasses

			riginator rations	Solo Licensees	Local Co- marketers	ī	Local Originator Corporations		Solo Licensees	Local Co- marketers
			Mean # Other					Mean # Other		
Country	Total Launches	Launches	Country Launches	Launches	Launches	Total Launches	Launches	Country Launches	Launches	Launches
High-Price EU		Ladiioiioo	Ladiioiioo	2441101100	Ladiioiioo	Ladiioiioo	Ladiioiioo	2001101100	244.161.166	<u> </u>
Germany	88	4	12.8	10	3	131	26	7.7	22	4
UK	80	11	12.7	7	0	116	18	11.1	11	1
Netherlands	73	0	N/A	0	0	112	2	12.5	3	0
Sweden	77	3	13.3	1	0	82	0	N/A	3	0
Low-Price EU	Countries									
France	69	11	10.5	5	3	105	19	7.3	16	2
Greece	72	0	N/A	0	0	107	0	N/A	2	0
Italy	76	1	0.0	13	14	127	11	5.0	25	13
Portugal	62	0	N/A	2	2	113	0	N/A	5	2
Spain	76	2	4.0	5	7	112	8	5.1	11	4
High-Price no	n-EU Count	ries								
Canada	73	0	N/A	3	0	98	1	1.0	10	0
Japan	53	17	4.0	6	11	158	45	1.9	36	24
Switzerland	78	11	10.6	4	0	111	25	11.2	10	0
USA	86	30	10.6	20	0	97	29	9.6	33	1
Low-Price nor	-EU Countr	ies								
Brazil	71	0	N/A	2	0	92	3	0.7	4	1
Mexico	72	0	N/A	0	0	105	1	0.0	2	0

Note: Sample includes all molecules present and all country launches occurring in IMS dataset through 2003

Note: Molecules with multiple local corporations in a country were assigned to a single one in the order of Local Originator, Solo Licensee or Local Co-marketer

Table 5. Coefficients and Marginal Effects for Launch Model (Standard Errors in Brackets)

	Coefficients			Marginal Effects				
		rith Robust red SEs	Clog-log w RI		Clog-log w Cluster		Clog-log w Ri	rith Normal Es
		Subo	class			Sub	class	
Variables	Superior	Inferior	Superior	Inferior	Superior	Inferior	Superior	Inferior
Log Avg Price of Superior Brands (Lag	0.1138**	-0.1089	0.1442***	-0.1202*	0.0053*	-0.0001	0.0086***	-0.0003
1Q)	[0.0552]	[0.0683]	[0.0476]	[0.0616]	[0.0027]	[0.0001]	[0.0032]	[0.0002]
Log Avg Price of Inferior Brands (Lag	0.0658	0.1247*	0.0894*	0.0928	0.0031	0.0001	0.0053*	0.0002
1Q)	[0.0612]	[0.0697]	[0.0506]	[0.0609]	[0.0029]	[0.0001]	[0.0031]	[0.0002]
Log Total Volume of All Drugs in Class	-0.0738	0.0829	-0.0202	0.1386	-0.0034	0.0001	-0.0012	0.0004
(Lag 1Q)	[0.0527]	[0.1058]	[0.0581]	[0.0889]	[0.0027]	[0.0001]	[0.0035]	[0.0002]
Num Generic Manufs per Molc in	-0.0040	0.0035	-0.0079	0.0103	-0.0002	0.0000	-0.0005	0.0000
Superior Subclass (Lag 1Q)	[0.0045]	[0.0081]	[0.0059]	[0.0083]	[0.0002]	[0.0000]	[0.0004]	[0.0000]
Num Generic Manufs per Molc in	-0.0016	-0.0030	-0.0014	-0.0023	-0.0001	0.0000	-0.0001	0.0000
Inferior Subclass (Lag 1Q)	[0.0021]	[0.0031]	[0.0017]	[0.0025]	[0.0001]	[0.0000]	[0.0001]	[0.0000]
No Molecules in Superior Subclass	0.2585	-0.2298	0.1416	-0.2018	0.0135	-0.0002	0.0089	-0.0005
D.V.	[0.1848]	[0.2583]	[0.1947]	[0.2353]	[0.0105]	[0.0002]	[0.0129]	[0.0006]
No Molecules in Inferior Subclass D.V.	-0.5880***	-0.3552	-0.5529	-0.3155	-0.0210***	-0.0002	-0.0264	-0.0007
	[0.2181]	[0.8341]	[0.4677]	[0.5482]	[0.0074]	[0.0005]	[0.0181]	[0.0011]
Time Since Global Launch (Yrs)	-0.6240***	-0.2767***	-0.4540***	-0.2772***	-0.0291***	-0.0002**	-0.0271***	-0.0007***
	[0.0578]	[0.0323]	[0.0536]	[0.0288]	[0.0046]	[0.0001]	[0.0049]	[0.0002]
Time Since Global Launch Squared	0.0231***	0.0028***	0.0158***	0.0029***	0.0011***	0.0000**	0.0009***	0.0000***
(Yrs)	[0.0027]	[0.0004]	[0.0030]	[0.0004]	[0.0002]	[0.0000]	[0.0002]	[0.0000]
First Global Launch Before 1990 D.V.	-0.0034	0.5749**	-0.2426	0.7042**	-0.0002	0.0006*	-0.0131	0.0023**
	[0.1931]	[0.2743]	[0.2860]	[0.3089]	[0.0090]	[0.0003]	[0.0149]	[0.0011]
First Global Launch in [1996-end] D.V.	-0.0497	-0.0329	-0.1018	-0.0383	-0.0023	0.0000	-0.0058	-0.0001
	[0.1479]	[0.2194]	[0.1907]	[0.3420]	[0.0066]	[0.0002]	[0.0108]	[0.0009]
Num Already Launched (UK, Germany)	0.5935***	0.5632***	0.4902***	0.4131***	0.0290***	0.0004	0.0301***	0.0011**
	[0.0920]	[0.1848]	[0.0785]	[0.1466]	[0.0071]	[0.0003]	[0.0072]	[0.0005]
Num Already Launched (Sweden,	0.5079***	0.6167***	0.3935***	0.3492**	0.0245***	0.0005*	0.0239***	0.0009*
Netherlands)	[0.0705]	[0.1548]	[0.0749]	[0.1436]	[0.0052]	[0.0003]	[0.0059]	[0.0005]
Num Already Launched (Italy, France)	0.2688***	0.0334	0.3057***	0.0945	0.0126**	0.0000	0.0185***	0.0002
	[0.0986]	[0.1192]	[0.0840]	[0.1364]	[0.0055]	[0.0001]	[0.0058]	[0.0004]
Num Already Launched (Spain,	0.0670	0.2820***	-0.0409	0.1621*	0.0031	0.0002**	-0.0024	0.0004

Portugal, Greece)	[0.0661]	[0.1038]	[0.0654]	[0.0950]	[0.0030]	[0.0001]	[0.0040]	[0.0003]
Num Already Launched (Canada,	0.1907***	0.0034	0.1321**	-0.0844	0.0089***	0.0000	0.0079**	-0.0002
Japan, Switzerland, USA)	[0.0637]	[0.1011]	[0.0560]	[0.1048]	[0.0030]	[0.0001]	[0.0035]	[0.0003]
Any PI Share in Subclass D.V.	0.0161	-0.1603	0.0670	-0.1269	0.0008	-0.0001	0.0041	-0.0003
•	[0.1536]	[0.4589]	[0.1510]	[0.3596]	[0.0072]	[0.0003]	[0.0092]	[0.0009]
Launch by Local Originator Corporation	1.3954***	2.5515***	1.5681***	3.0584***	0.1286***	0.0080*	0.1822***	0.0274***
D.V.	[0.2676]	[0.2759]	[0.1626]	[0.2261]	[0.0458]	[0.0042]	[0.0420]	[0.0104]
Launch by Solo Licensee Corporation	0.5481***	1.3352***	0.5646***	1.1030***	0.0328**	0.0020*	0.0424***	0.0042**
D.V.	[0.1765]	[0.2071]	[0.1551]	[0.1928]	[0.0130]	[0.0011]	[0.0157]	[0.0018]
Launch by Local Co-marketer	0.5592***	1.6151***	0.3463*	1.1864***	0.0340**	0.0029	0.0239	0.0048*
Corporation D.V.	[0.1786]	[0.5425]	[0.1984]	[0.3468]	[0.0145]	[0.0026]	[0.0160]	[0.0028]
USD to (ECU or Euro) Exchange Rate	0.0945	-0.1796	-0.1920	-0.9665*	0.0044	-0.0001	-0.0115	-0.0025
	[0.4523]	[0.5755]	[0.4216]	[0.5723]	[0.0209]	[0.0004]	[0.0255]	[0.0017]
UK D.V.	-0.2430	0.1925	-0.1616	0.2704	-0.0101	0.0002	-0.0090	0.0008
	[0.2096]	[0.3408]	[0.2000]	[0.3605]	[0.0085]	[0.0003]	[0.0111]	[0.0010]
Netherlands D.V.	-0.8790***	0.6443*	-0.7894***	0.7236*	-0.0276***	0.0007	-0.0343***	0.0024*
	[0.2465]	[0.3618]	[0.2194]	[0.3934]	[0.0082]	[0.0005]	[0.0106]	[0.0015]
Sweden D.V.	-0.7520**	-0.0208	-0.5688**	0.1684	-0.0249**	0.0000	-0.0269**	0.0005
	[0.2968]	[0.4805]	[0.2307]	[0.4521]	[0.0099]	[0.0003]	[0.0116]	[0.0012]
France D.V.	-1.2645***	-0.7394	-1.2202***	-0.5880	-0.0340***	-0.0004	-0.0452***	-0.0012
	[0.2103]	[0.5553]	[0.2304]	[0.5148]	[0.0078]	[0.0004]	[0.0110]	[0.0012]
Greece D.V.	-1.2725***	0.8260	-1.0673***	1.0116*	-0.0341***	0.0009	-0.0418***	0.0038*
	[0.2637]	[0.5778]	[0.2496]	[0.5260]	[0.0086]	[0.0006]	[0.0116]	[0.0020]
Italy D.V.	-0.9829***	0.0104	-0.8500***	0.1479	-0.0296***	0.0000	-0.0361***	0.0004
	[0.2439]	[0.5519]	[0.2173]	[0.4900]	[0.0084]	[0.0004]	[0.0109]	[0.0013]
Portugal D.V.	-1.9128***	0.0829	-1.7246***	0.2123	-0.0405***	0.0001	-0.0536***	0.0006
	[0.2319]	[0.6012]	[0.2522]	[0.5390]	[0.0084]	[0.0004]	[0.0115]	[0.0015]
Spain D.V.	-0.8052***	0.1745	-0.6327***	0.2515	-0.0261***	0.0001	-0.0292***	0.0007
	[0.2007]	[0.5481]	[0.2181]	[0.5054]	[0.0079]	[0.0004]	[0.0112]	[0.0014]
Canada D.V.	-1.0640***	-0.5232	-0.9515***	-0.3179	-0.0310***	-0.0003	-0.0389***	-0.0007
. 51/	[0.2341]	[0.5649]	[0.2103]	[0.4931]	[0.0084]	[0.0004]	[0.0108]	[0.0012]
Japan D.V.	-2.4793***	-0.0753	-2.5477***	-0.0787	-0.0436***	-0.0001	-0.0613***	-0.0002
0 % 1 151/	[0.2353]	[0.6375]	[0.2339]	[0.4660]	[0.0083]	[0.0004]	[0.0114]	[0.0012]
Switzerland D.V.	-1.0800***	-0.3521	-0.9100***	-0.2888	-0.0313***	-0.0002	-0.0378***	-0.0007
HOA B.V	[0.2644]	[0.5853]	[0.2540]	[0.5726]	[0.0089]	[0.0004]	[0.0119]	[0.0014]
USA D.V.	-0.9119***	-1.2651**	-0.9440***	-1.3975***	-0.0283***	-0.0005	-0.0387***	-0.0022**

	[0.2876]	[0.6150]	[0.2336]	[0.4889]	[0.0085]	[0.0003]	[0.0104]	[0.0011]
Brazil D.V.	-1.1626***	0.3617	-0.9768***	0.5353	-0.0326***	0.0003	-0.0395***	0.0017
	[0.2411]	[0.5740]	[0.2211]	[0.4874]	[0.0084]	[0.0005]	[0.0111]	[0.0014]
Mexico D.V.	-1.3041***	0.1336	-1.1008***	0.4037	-0.0345***	0.0001	-0.0426***	0.0012
	[0.2765]	[0.6370]	[0.2448]	[0.5370]	[0.0086]	[0.0005]	[0.0115]	[0.0015]
Anti-asthma D.V.	-0.0373	0.3802	-0.1868	0.0520	-0.0017	0.0003	-0.0104	0.0001
	[0.3098]	[0.2919]	[0.3652]	[0.3537]	[0.0140]	[0.0003]	[0.0191]	[0.0009]
Anti-clotting D.V.	-0.9282***	-0.3463	-0.9634**	-0.1215	-0.0304***	-0.0002	-0.0415***	-0.0003
	[0.3196]	[0.4787]	[0.4011]	[0.5280]	[0.0098]	[0.0003]	[0.0143]	[0.0013]
Anti-depressants D.V.	0.0637	0.2604	0.1975	0.1251	0.0030	0.0002	0.0127	0.0003
	[0.2100]	[0.4250]	[0.2943]	[0.4409]	[0.0101]	[0.0004]	[0.0200]	[0.0012]
Epileptics D.V.	-0.1567	-0.1624	-0.1626	-0.4113	-0.0069	-0.0001	-0.0093	-0.0009
	[0.1965]	[0.8146]	[0.2912]	[0.9170]	[0.0085]	[0.0005]	[0.0161]	[0.0018]
Anti-nauseants D.V.	-0.8991***	-0.5673	-0.8997**	-0.7966	-0.0299***	-0.0003	-0.0397***	-0.0016
	[0.2905]	[0.4947]	[0.3895]	[0.8121]	[0.0092]	[0.0003]	[0.0147]	[0.0014]
Parkinsons D.V.	-0.5001	0.5686	-0.2244	0.9530	-0.0189*	0.0005	-0.0123	0.0035
	[0.3046]	[0.4592]	[0.3891]	[0.6221]	[0.0104]	[0.0006]	[0.0199]	[0.0033]
Anti-psychotics D.V.	-0.2864	0.4003	-0.0411	1.9889***	-0.0118	0.0004	-0.0024	0.0111
	[0.3312]	[0.6463]	[0.4012]	[0.7153]	[0.0125]	[0.0007]	[0.0233]	[0.0077]
Anti-ulcerants D.V.	-0.3775	-0.4143	-0.5970*	-0.7731	-0.0154*	-0.0003	-0.0296**	-0.0016
	[0.2491]	[0.4924]	[0.3152]	[0.5888]	[0.0092]	[0.0003]	[0.0140]	[0.0011]
Lipid lowering D.V.	0.0097	0.2351	0.2194	0.5170	0.0005	0.0002	0.0143	0.0016
	[0.2384]	[0.5583]	[0.3349]	[0.5803]	[0.0112]	[0.0005]	[0.0237]	[0.0021]
Migraine D.V.	-0.6203*	-0.4250	-0.2167	-0.7202	-0.0222**	-0.0003	-0.0119	-0.0015
	[0.3335]	[0.6389]	[0.4338]	[0.9795]	[0.0104]	[0.0004]	[0.0222]	[0.0017]
Osteoporosis D.V.	-0.2208	0.5820	-0.3252	0.3763	-0.0094	0.0005	-0.0173	0.0010
	[0.3251]	[0.3609]	[0.3237]	[0.3919]	[0.0128]	[0.0004]	[0.0156]	[0.0012]
Constant	-1.1015	-5.6436***	-1.6217	-5.7109***				
	[0.9652]	[1.7327]	[1.0217]	[1.6644]				
Num Observations	23,400	96,041	23,400	96,041	23,400	96,041	23,400	96,041
Number of Molecule-level Clusters	111	239	111	239	111	239	111	239
Model Log-Likelihood	-3071.2	-2007.6	-3045.7	-1963.4				
Mean of Dependent Variable	0.0378	0.0041	0.0378	0.0041	0.0378	0.0041	0.0378	0.0041

Standard errors in brackets

* significant at 10%; ** significant at 5%; *** significant at 1% Note: All prices are ex manufacturer prices per standard unit in 2003 US Dollars

Table 6. Marginal Effects of Prior Foreign Launch on Launch Hazard in Low-Price EU Countries for Superior Subclasses (Standard Errors in Brackets)

Marginal Effects of Prior Foreign Launch in Low-Price EU Countries

Clog-log with Robust Clustered SEs

Net effect of a Single Prior Launch in a High-Price EU Country Net effect of a Single Prior Launch in a Low-Price EU Country Difference	0.0018*** [0.0004] 0.0005*** [0.0002] 0.0013*** [0.0003]
Net effect of a Single Prior Launch in a High-Price EU Country Net effect of a Single Prior Launch in a High-Price non-EU Country	0.0018*** [0.0004] 0.0006*** [0.0002] 0.0012***
Difference	[0.0004]

Standard errors in brackets

^{*} significant at 10%; ** significant at 5%; *** significant at 1%

Table 7. Marginal Effects of Launch by a Local Corporation (Standard Errors in Brackets)

Superior Subclasses Inferior Subclasses # Launches # Launches Country-Countryby Any Mean by Any Mean specific Local Quarterly specific Local Quarterly Country Net Effect Corporation Hazard Net Effect Corporation Hazard Germany 0.0121 6.3% 0.0030 0.3% 11 6 [0.0162] [0.0024]UK 0.0446 10 4.4% 0.0029* 7 0.4% [0.0346] [0.0017]0 1 Netherlands N/A 4.4% 0.0140 0.2% [0.0222]Sweden -0.0107 3 3.7% 0.0054 1 0.3% [0.0070][0.0042]0.0316*** 6 France 11 3.2% 0.0009 0.3% [0.0111] [8000.0]Greece N/A 0 3.4% 0.0031 1 0.5% [0.0057]Italy 0.0145** 21 4.5% 0.0040* 9 0.5% [0.0069] [0.0023] 0.0110** 4 5 0.4% Spain 4.3% 0.0026 [0.0056][0.0023]Portugal -0.0001 13 2.4% N/A 0 0.4% [0.0010]2 Canada 0.0058 3.9% 8000.0 3 0.3% [0.0078] [0.0006]0.0092*** 1.8% 31 1.0% Japan 25 0.0096* [0.0029][0.0057]Switzerland 0.0448*** 5 12 4.4% 0.0013 0.3% [0.0155] [0.0013]**USA** 0.0097 41 5.4% 0.0008 12 0.3% [0.0091] [0.0005]Brazil 0.0042 2 3.7% 0.0046 6 0.5% [0.0049] [0.0036]3 0 Mexico N/A 3.5% 0.0176 0.4% [0.0137]

Standard errors in brackets

Note: Effects for some countries were not estimable due to an absence of launches by local corporations in those countries

^{*} significant at 10%; ** significant at 5%; *** significant at 1%

Table 8. Determinants of Launch Prices, OLS and Normal Random Effects Regressions (Standard Errors in Brackets)

	0	LS w/ Robust	Clustered SI	Es	Normal Random Effects			
	Superior	Inferior	Superior	Inferior	Superior	Inferior	Superior	Inferior
Variables	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses
Superior Brands' Price Missing D.V.	-0.0615	0.0172	-0.0636	0.0252	-0.0644	0.0324	-0.0685	0.0362
	[0.1879]	[0.1199]	[0.1886]	[0.1205]	[0.1134]	[0.1114]	[0.1138]	[0.1125]
Log Avg Price of Superior Brands (Lag	0.1574***	-0.004	0.1586***	0.0018	0.1244***	-0.0085	0.1283***	0.0032
1Q)	[0.0394]	[0.0467]	[0.0395]	[0.0480]	[0.0201]	[0.0336]	[0.0201]	[0.0336]
Inferior Brands' Price Missing D.V.	0.1093	-0.7439	0.114	-0.8025	0.004	-1.2424**	0.0157	-1.2936***
	[0.1523]	[0.7108]	[0.1541]	[0.7423]	[0.1118]	[0.4849]	[0.1121]	[0.4878]
Log Avg Price of Inferior Brands (Lag 1Q)	0.0393	0.2154***	0.0387	0.2134***	0.0844***	0.1681***	0.0830***	0.1669***
	[0.0398]	[0.0490]	[0.0397]	[0.0488]	[0.0206]	[0.0294]	[0.0206]	[0.0297]
Generics' Price Missing D.V.	0.0674	0.3731*	0.0689	0.3779*	0.0095	0.2412*	0.0115	0.2416*
	[0.1168]	[0.2001]	[0.1165]	[0.2011]	[0.0758]	[0.1236]	[0.0761]	[0.1251]
Log Avg Price of Generics in Class (Lag	0.0292	-0.1194***	0.0294	-0.1206***	0.0169	-0.0857**	0.0162	-0.0834**
1Q)	[0.0295]	[0.0441]	[0.0296]	[0.0439]	[0.0194]	[0.0340]	[0.0195]	[0.0345]
Time Since Global Launch (Yrs)	-0.0427	-0.0245	-0.0413	-0.0264	-0.0391	-0.0077	-0.0367	-0.0114
	[0.0265]	[0.0214]	[0.0265]	[0.0215]	[0.0260]	[0.0212]	[0.0261]	[0.0212]
Time Since Global Launch Squared (Yrs)	0.0028	-0.0006	0.0027	-0.0005	0.0005	-0.0012*	0.0003	-0.0011
	[0.0018]	[0.0006]	[0.0018]	[0.0006]	[0.0018]	[0.0007]	[0.0018]	[0.0007]
First Brand Launch in Ctry-Subclass D.V.	0.1998	0.8668**	0.2034	0.8990**	0.2132	1.1729***	0.2164*	1.2199***
	[0.1656]	[0.4080]	[0.1656]	[0.4390]	[0.1307]	[0.3525]	[0.1312]	[0.3562]
Second Brand Launch in Ctry-Subclass	0.3496***	0.6166*	0.3486***	0.6076*	0.2819***	0.6197***	0.2770***	0.6406***
D.V.	[0.0824]	[0.3115]	[0.0824]	[0.3244]	[0.0749]	[0.2309]	[0.0751]	[0.2321]
Third or Fourth Brand Launch in Ctry-	0.2412***	0.3085*	0.2399***	0.3039*	0.1859***	0.2007	0.1816***	0.2066
Subclass D.V.	[0.0611]	[0.1615]	[0.0613]	[0.1631]	[0.0555]	[0.1465]	[0.0557]	[0.1471]
High-price EU Min Own Price Missing	0.2170***	-0.0523	0.2138**	-0.043	0.0649	-0.0451	0.0631	-0.0526
D.V.	[0.0821]	[0.1372]	[0.0821]	[0.1379]	[0.0575]	[0.0956]	[0.0578]	[0.0966]
Log Min Own Price in Hi-Price EU (Lag	0.2179***	0.3905***	0.2174***	0.3960***	0.1000***	0.2746***	0.1012***	0.2847***
1Q)	[0.0623]	[0.0885]	[0.0622]	[0.0890]	[0.0261]	[0.0586]	[0.0262]	[0.0590]
Low-price EU Min Own Price Missing	-0.0185	-0.0755	-0.0161	-0.0733	0.0251	0.0545	0.03	0.051
D.V.	[0.0524]	[0.1262]	[0.0527]	[0.1259]	[0.0483]	[0.1028]	[0.0485]	[0.1039]
Log Min Own Price in Low-Price EU (Lag	-0.0243	-0.1086	-0.0234	-0.101	-0.0221	-0.1095	-0.0188	-0.1113
1Q)	[0.0394]	[0.1196]	[0.0390]	[0.1184]	[0.0279]	[0.0790]	[0.0280]	[0.0797]

High-price non-EU Min Own Price	0.1054	-0.3101***	0.1076	-0.3195***	0.1019*	-0.1126	0.1049*	-0.1244
Missing D.V.	[0.0649]	[0.1100]	[0.0651]	[0.1112]	[0.0554]	[0.0926]	[0.0556]	[0.0936]
Log Min Own Price in Hi-Price non-EU	0.2682***	0.0482	0.2677***	0.0532	0.1433***	-0.0278	0.1425***	-0.0197
(Lag 1Q)	[0.0522]	[0.0661]	[0.0519]	[0.0640]	[0.0251]	[0.0564]	[0.0252]	[0.0570]
Any PI Share in Subclass D.V.	0.022	-0.5218**	0.0232	-0.4815**	0.0207	-0.4798***	0.0236	-0.4662***
	[0.0795]	[0.2029]	[0.0786]	[0.2134]	[0.0650]	[0.1710]	[0.0653]	[0.1732]
Log GDP per Capita	0.862	2.8591*			1.8350**	3.1564***		
	[0.9232]	[1.5889]			[0.7426]	[1.1628]		
Launch by Local Originator Corporation	0.0311	-0.2572*	0.0332	-0.2823*	0.0693	-0.1825	0.0739	-0.2207*
D.V.	[0.1009]	[0.1541]	[0.1011]	[0.1551]	[0.0694]	[0.1146]	[0.0696]	[0.1148]
Launch by Solo Licensee Corporation	-0.0742	-0.1455	-0.0708	-0.1377	-0.0543	-0.0178	-0.0452	-0.0126
D.V.	[0.0770]	[0.1246]	[0.0768]	[0.1238]	[0.0623]	[0.0940]	[0.0625]	[0.0951]
Launch by Local Co-marketer	0.0031	-0.1025	0.0038	-0.1282	0.0265	-0.1417	0.0288	-0.1679
Corporation D.V.	[0.0971]	[0.1908]	[0.0980]	[0.1937]	[0.0782]	[0.1457]	[0.0785]	[0.1470]
USD to (ECU or Euro) Exchange Rate	-0.1475	2.4535**	-0.1373	2.2873**	-0.0265	1.7490**	0.0042	1.5484*
	[0.6339]	[1.0738]	[0.6234]	[1.0742]	[0.4524]	[0.8151]	[0.4542]	[0.8220]
Country-Specific Quarterly Producer	-0.0089**	0.002	-0.0065	0.0101	-0.0088**	-0.0122*	-0.0038	-0.0021
Price Index	[0.0044]	[0.0083]	[0.0039]	[0.0064]	[0.0044]	[0.0067]	[0.0039]	[0.0056]
Avg Pack Size (Up to 100)	-0.0118***	-0.0102***	-0.0118***	-0.0100***	-0.0094***	-0.0093***	-0.0093***	-0.0093***
	[0.0017]	[0.0025]	[0.0017]	[0.0025]	[0.0010]	[0.0017]	[0.0010]	[0.0017]
Pack Size > 100 D.V.	-1.1996***	-1.6199***	-1.2014***	-1.5868***	-0.9891***	-1.4086***	-0.9930***	-1.4137***
	[0.1880]	[0.2118]	[0.1888]	[0.2062]	[0.1114]	[0.1754]	[0.1118]	[0.1773]
Avg Pill Strength (g)	0.4791*	0.0507	0.4928**	0.045	0.0577	0.1877	0.0766	0.1844
	[0.2452]	[0.0701]	[0.2457]	[0.0704]	[0.2737]	[0.1440]	[0.2738]	[0.1437]
Form: Oral Solid Delayed D.V.	-0.1014	-0.191	-0.1092	-0.1573	0.1059	0.0933	0.0926	0.0949
	[0.1994]	[0.1444]	[0.1988]	[0.1452]	[0.1823]	[0.1567]	[0.1829]	[0.1582]
Form: Injectable D.V.	2.0793***	1.7450***	2.0865***	1.7304***	1.7654***	1.9669***	1.7827***	1.9652***
	[0.3009]	[0.3579]	[0.3025]	[0.3570]	[0.0885]	[0.2165]	[0.0887]	[0.2185]
Form: Other	-0.0523	0.3387***	-0.0551	0.3414***	0.0793	0.0856	0.0734	0.0754
	[0.1683]	[0.1087]	[0.1663]	[0.1101]	[0.1702]	[0.1270]	[0.1708]	[0.1278]
UK D.V.	-0.3218***	-0.1577	-0.2751***	-0.0316	-0.2829***	-0.1225	-0.1861**	0.0103
	[0.0957]	[0.1238]	[0.0827]	[0.1202]	[0.0875]	[0.1532]	[0.0784]	[0.1472]
Netherlands D.V.	-0.0707	0.0041	-0.0734	-0.0454	-0.0669	-0.0277	-0.0711	-0.0781
	[0.1034]	[0.1372]	[0.1022]	[0.1291]	[0.0798]	[0.1557]	[0.0802]	[0.1568]
Sweden D.V.	-0.1746	-0.1626	-0.0541	0.2208	-0.3307**	-0.3624	-0.0743	0.0489
	[0.1684]	[0.2808]	[0.0859]	[0.2142]	[0.1335]	[0.2211]	[0.0844]	[0.1645]

France D.V.	-0.2055*	-0.7264***	-0.2382**	-0.8051***	-0.1272	-0.6820***	-0.1956**	-0.8015***
	[0.1117]	[0.2689]	[0.1098]	[0.2668]	[0.1003]	[0.2275]	[0.0969]	[0.2264]
Greece D.V.	0.2805	1.2931	-0.3932***	-0.8704***	1.1945**	1.6279*	-0.2393**	-0.7916***
	[0.7187]	[1.2267]	[0.1198]	[0.2541]	[0.5893]	[0.9173]	[0.1050]	[0.2231]
Italy D.V.	-0.19	-0.286	-0.3485***	-0.7525***	0.0835	-0.0639	-0.2521***	-0.6007***
	[0.1922]	[0.3790]	[0.1031]	[0.2579]	[0.1664]	[0.2958]	[0.0970]	[0.2237]
Portugal D.V.	0.373	1.4824	-0.3098***	-0.7693***	1.2153**	1.8910**	-0.2341**	-0.6161***
	[0.7279]	[1.2936]	[0.1107]	[0.2566]	[0.5961]	[0.9494]	[0.1077]	[0.2260]
Spain D.V.	0.1941	0.8328	-0.2498**	-0.6269***	0.7733**	0.9693	-0.1710*	-0.6661***
	[0.4840]	[0.8717]	[0.0992]	[0.2389]	[0.3931]	[0.6412]	[0.0928]	[0.2222]
Canada D.V.	0.0439	-0.2773	0.0363	-0.2971	0.0338	-0.2256	0.0197	-0.2684
	[0.1078]	[0.2963]	[0.1054]	[0.3000]	[0.0923]	[0.2247]	[0.0925]	[0.2273]
Japan D.V.	0.1482	-0.965	0.5852***	0.5110*	-0.3752	-1.0523	0.5573***	0.5585**
	[0.5017]	[0.8135]	[0.2021]	[0.2901]	[0.3967]	[0.6419]	[0.1221]	[0.2475]
Switzerland D.V.	-0.1417	-1.6358**	0.2091**	-0.4370*	-0.6109*	-1.5100***	0.1389	-0.2329
	[0.4039]	[0.6852]	[0.0911]	[0.2507]	[0.3162]	[0.5209]	[0.0888]	[0.2234]
United States D.V.	0.1838	-0.8207	0.5272***	0.3319	-0.35	-1.0079**	0.3833***	0.2312
	[0.4014]	[0.6421]	[0.1347]	[0.2545]	[0.3125]	[0.5095]	[0.0987]	[0.2295]
Brazil D.V.	1.2852	4.733	-0.3136***	-0.5128**	3.1815**	5.3703**	-0.2182**	-0.4591**
	[1.6940]	[2.9427]	[0.1111]	[0.2464]	[1.3787]	[2.1576]	[0.0940]	[0.2198]
Mexico D.V.	0.9276	2.9574	-0.2477**	-0.9032***	2.3375**	3.4723**	-0.1610*	-0.8161***
	[1.2631]	[2.1732]	[0.1215]	[0.2317]	[1.0153]	[1.5948]	[0.0960]	[0.2294]
Constant	-7.0696	-30.4859*	1.2244	-2.7327*	-16.8192**	31.7366***	0.8265	-1.0996
	[8.7738]	[15.4844]	[0.8798]	[1.5754]	[7.1791]	[11.3552]	[0.7247]	[1.1956]
Log GDP per Capita Included?	Yes	Yes	No	No	Yes	Yes	No	No
Year Fixed Effects Included?	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Observations	950	423	950	423	950	423	950	423
Number of Molecule-level Clusters	109	123	109	123	109	123	109	123
R-squared	0.89	0.79	0.89	0.79	0.87	0.76	0.87	0.76
Mean of Dependent Variable	0.74	-0.39	0.74	-0.39	0.74	-0.39	0.74	-0.39
	1	0.00	· · ·	0.00		0.00		0.00

Standard errors in brackets

Note: All prices are ex manufacturer prices per standard unit in 2003 US Dollars

^{*} significant at 10%; ** significant at 5%; *** significant at 1%

Table 9. Spillover Effects of Own Price on Launch Price in Low-Price EU Countries for Superior Subclasses

Marginal Effects of Log Min Own Price on Launch Price in Low-Price EU Countries

	OLS w/ Robust Clustered SEs	Normal Random Effects
Net effect of Log Min Own Price in High-	0.4236***	0.2508***
Price EU Countries (Lag 1Q)	[0.0861]	[0.0845]
Net effect of Log Min Own Price in Low-	0.0366	-0.0096
Price EU Countries (Lag 1Q)	[0.0591]	[0.0432]
Difference	0.3870*** [0.1103]	0.2604** [0.1055]
Net effect of Log Min Own Price in High-	0.4236***	0.2508***
Price EU Countries (Lag 1Q)	[0.0861]	[0.0845]
Net effect of Log Min Own Price in High-	0.0551	0.0096
Price non-EU Countries (Lag 1Q)	[0.0837]	[0.0778]
Difference	0.3684** [0.1641]	0.2411 [0.1580]

Standard errors in brackets

^{*} significant at 10%; ** significant at 5%; *** significant at 1% Note: All prices are ex manufacturer prices per standard unit in 2003 US dollars

Appendix Table A1. Descriptive Statistics for Launch Model Variables

	Sı	uperior		
	Sub	classes	Inferior	Subclasses
Variable Description	Mean	Std. Dev.	Mean	Std. Dev.
New Launch D.V.	0.04	0.19	0.00	0.06
Log Avg Price of Superior Brand Drugs (Lag 1Q)	0.52	1.33	0.68	1.47
Log Avg Price of Inferior Brand Drugs (Lag 1Q)	-0.54	1.29	-0.48	1.46
Log Total Volume of All Drugs In (Lag 1Q)	12.68	1.75	12.94	1.69
Num Generic Manufs per Molc in Superior Sub (Lag				
1Q)	5.08	10.30	4.41	9.27
Num Generic Manufs per Molc in Inferior Sub (Lag				
1Q)	28.95	37.17	35.80	41.52
No Molecules in Superior Sub D.V.	0.05	0.22	0.05	0.23
No Molecules in Inferior Sub D.V.	0.01	0.07	0.01	0.07
Time Since Molecule Global Launch (Years)	4.41	3.97	14.70	11.88
Time Since Molecule Global Launch Squared	35.20	56.96	357.07	739.83
First Global Launch Before 1990 D.V.	0.21	0.40	0.72	0.45
First Global Launch In [1990-1995] D.V.	0.38	0.48	0.20	0.40
First Global Launch In [1996-2002] D.V.	0.42	0.49	0.08	0.26
Num Already Launched (UK, Germany)	0.69	0.82	0.51	0.73
Num Already Launched (Italy, France)	0.51	0.72	0.48	0.70
Num Already Launched (Sweden, Netherlands)	0.56	0.78	0.30	0.61
Num Already Launched (Spain, Portugal, Greece)	0.58	0.94	0.58	0.91
Num Already Launched (Canada, Japan,				
Switzerland, USA)	1.18	0.94	1.01	0.95
Any PI Share In Subclass- D.V.	0.16	0.36	0.22	0.41
Launch by Local Originator Corporation D.V.	0.02	0.13	0.02	0.15
Launch by Solo Licensee Corporation D.V.	0.04	0.19	0.03	0.16
Launch by Local Co-marketer Corporation D.V.	0.01	0.11	0.00	0.04
USD to (ECU or Euro) Exchange Rate	1.09	0.13	1.11	0.14
Germany D.V.	0.05	0.22	0.05	0.23
UK D.V.	0.06	0.23	0.06	0.25
Netherlands D.V.	0.06	0.25	0.06	0.25
Sweden D.V.	0.06	0.24	0.08	0.27
France D.V.	0.07	0.26	0.07	0.25
Greece D.V.	0.07	0.25	0.07	0.26
Italy D.V.	0.06	0.23	0.06	0.23
Portugal D.V.	0.09	0.28	0.07	0.25
Spain D.V.	0.06	0.24	0.06	0.25
Canada D.V.	0.07	0.25	0.07	0.26
Japan D.V.	0.10	0.30	0.05	0.21
Switzerland D.V.	0.06	0.24	0.07	0.25
US D.V.	0.06	0.23	0.07	0.26
Brazil D.V.	0.07	0.25	0.08	0.27
Mexico D.V.	0.07	0.26	0.07	0.26
Anti-asthma D.V.	0.05	0.21	0.16	0.37
Anti-clotting D.V.	0.07	0.25	0.10	0.29
Anti-depressants D.V.	0.09	0.29	0.09	0.28
Epileptics D.V.	0.15	0.36	0.03	0.16
Anti-hypertensives D.V.	0.21	0.41	0.17	0.38

Anti-nausea D.V.	0.07	0.26	0.04	0.21
Parkinsons D.V.	0.05	0.22	0.01	0.11
Anti-psychotic D.V.	0.05	0.22	0.01	0.12
Anti-ulcerant D.V.	0.11	0.32	0.09	0.28
Lipid-lowering D.V.	0.04	0.20	0.05	0.22
Migraine D.V.	0.03	0.17	0.03	0.17
Osteoporosis D.V.	0.08	0.26	0.22	0.41
Sample Size	23	3,400	96	5,041

Note: All prices are ex manufacturer prices per standard unit in 2003 US Dollars

Appendix Table A2. Descriptive Statistics for Price Model Variables

Variable Description		Si	uperior		
Log of Price in 2003 USD per SU		Sub	oclasses	Inferior	Subclasses
Price in 2003 USD per SU 27.95 150.13 7.10 46.63 Superior Brands Price Missing D.V. 0.08 0.28 0.13 0.33 Log Avg Price of Superior Brands (Lag 1Q) 0.66 1.39 0.29 1.27 Inferior Brands' Price Missing D.V. 0.02 0.14 0.01 0.11 Log Avg Price of Inferior Brands (Lag 1Q) -0.53 1.41 -0.55 1.40 Generics' Price Missing D.V. 0.11 0.32 0.12 0.33 Log Avg Price of Generics In Class (Lag 1Q) -1.23 1.05 -1.06 1.11 Time Since Global Launch (Years) 2.16 2.49 6.54 7.32 Time Since Global Launch in Ctry-Subclass D.V. 0.11 0.31 0.01 0.12 Second Brand Launch in Ctry-Subclass D.V. 0.15 0.36 0.03 0.17 Firist Brand Launch in Ctry-Subclass D.V. 0.15 0.36 0.03 0.17 Firith or Later Brand Launch in Ctry-Subclass D.V. 0.44 0.50 0.85 0.36 Log Min Own Price In Hi-Price Missing D.V. <td< td=""><td>Variable Description</td><td>Mean</td><td>Std. Dev.</td><td>Mean</td><td>Std. Dev.</td></td<>	Variable Description	Mean	Std. Dev.	Mean	Std. Dev.
Superior Brands' Price Missing D.V.	Log of Price in 2003 USD per SU	0.74	1.65	-0.39	1.31
Log Avg Price of Superior Brands (Lag 1Q)	Price in 2003 USD per SU	27.95	150.13	7.10	46.63
Log Avg Price of Superior Brands (Lag 1Q)	Superior Brands' Price Missing D.V.	0.08	0.28	0.13	0.33
Interior Brands' Price Missing D.V. 0.02 0.14 0.01 0.11 1.09 1.40 0.75 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 1.40 0.05 0.12 0.33 1.09 0.07 0.26 0.11 0.32 0.12 0.33 1.09 0.12 0.33 1.09 0.12 0.33 1.09 0.12 0.33 1.00 0.12 0.33 1.00 0.12 0.05	·	0.66	1.39	0.29	1.27
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Portugal D.V. 0.05 0.22 0.07 0.25					
Spain D.V. 0.06 0.23 0.05 0.22					
	Spain D.V.	0.06	0.23	0.05	0.22
Canada D.V. 0.07 0.25 0.05 0.22					
Japan D.V. 0.05 0.21 0.11 0.31	Japan D.V.	0.05	0.21	0.11	0.31

Switzerland D.V.	0.07	0.26	0.06	0.24
USA D.V.	0.08	0.27	0.05	0.23
Brazil D.V.	0.07	0.25	0.05	0.23
Mexico D.V.	0.07	0.25	0.07	0.25
Year 1992 D.V.	0.07	0.25	0.13	0.33
Year 1993 D.V.	0.07	0.25	0.14	0.34
Year 1994 D.V.	0.07	0.25	0.09	0.29
Year 1995 D.V.	0.08	0.27	0.07	0.26
Year 1996 D.V.	0.08	0.28	0.11	0.32
Year 1997 D.V.	0.13	0.34	0.15	0.35
Year 1998 D.V.	0.15	0.36	0.07	0.25
Year 1999 D.V.	0.10	0.30	0.04	0.21
Year 2000 D.V.	0.09	0.28	0.07	0.25
Year 2001 D.V.	0.07	0.25	0.06	0.23
Year 2002 D.V.	0.05	0.22	0.05	0.23
Year 2003 D.V.	0.05	0.21	0.03	0.16
Sample Size		950	4	423

Note: All prices are ex manufacturer prices per standard unit in 2003 US Dollars

Appendix Table A3. Number of Launches, and Mean and Median In-Country Launch Delay by Domestic Status of Corporation

	Loc	al Originat	or	Local Co-marketer								
	Co	rporation	S	Solo Lice	nsee Corp	orations	Co	orporations	S	Non-loc	al Corpor	ations
		Mean	Median		Mean	Median		Mean	Median		Mean	Median
	Molecules	Launch	Launch	Molecules	Launch	Launch	Molecules	Launch	Launch	Molecules	Launch	Launch
	Launched	Delay in	Delay in	Launched	Delay in	Delay in	Launched	Delay in	Delay in	Launched	Delay in	Delay in
	in	Country	Country	in	Country	Country	in	Country	Country	in	Country	Country
Country	Country	(Mos)	(Mos)	Country	(Mos)	(Mos)	Country	(Mos)	(Mos)	Country	(Mos)	(Mos)
Superior Mo	lecules											
Germany	4	5.3	4	10	19.0	19	3	30.3	15	71	18.2	11
U.K.	11	4.3	1	7	11.1	0	0	N/A	N/A	62	18.2	8.5
Netherlands	0	N/A	N/A	0	N/A	N/A	0	N/A	N/A	73	16.5	10
Sweden	3	22.0	0	1	7.0	7	0	N/A	N/A	73	17.0	8
France	11	11.5	4	5	32.6	30	3	16.0	15	50	29.5	27
Greece	0	N/A	N/A	0	N/A	N/A	0	N/A	N/A	72	28.0	21
Italy	1	0.0	0	13	23.5	21	14	17.1	13	48	25.4	20.5
Portugal	0	N/A	N/A	2	66.5	66.5	2	38.5	38.5	58	32.1	31
Spain	2	0.0	0	5	24.0	21	7	25.7	18	62	29.4	23
Canada	0	N/A	N/A	3	34.0	35	0	N/A	N/A	70	27.2	14.5
Japan	17	1.8	0	6	59.2	53	11	49.2	44	19	53.0	47
Switzerland	11	10.3	5	4	48.8	11	0	N/A	N/A	63	21.5	18
U.S.A.	30	7.8	1.5	20	24.3	9	0	N/A	N/A	36	23.8	9.5
Brazil	0	N/A	N/A	2	17.5	17.5	0	N/A	N/A	68	31.0	21
Mexico	0	N/A	N/A	0	N/A	N/A	0	N/A	N/A	72	27.3	17
Inferior Mole	cules											
Germany	26	11.4	0	22	52.2	30.5	4	23.0	19	79	31.8	16
U.K.	18	6.7	0	11	51.6	22	1	103.0	103	86	47.6	26
Netherlands	2	55.5	55.5	3	19.3	0	0	N/A	N/A	106	38.3	19.5
Sweden	0	N/A	N/A	3	112.3	78	0	N/A	N/A	79	52.8	29
France	19	11.2	0	16	91.8	70	2	40.5	40.5	67	49.5	33
Greece	0	N/A	N/A	2	148.5	148.5	0	N/A	N/A	99	76.5	42
Italy	11	2.0	0	25	60.0	36	13	35.7	23	78	54.3	44
Portugal	0	N/A	N/A	5	49.0	51	2	7.0	7	104	61.6	41
Spain	8	5.8	0	11	47.0	24	4	35.0	29	87	48.4	40

Canada	1	17.0	17	10	99.1	124.5	0	N/A	N/A	87	71.5	52
Japan	45	0.7	0	36	64.6	56	24	68.8	54.5	53	65.5	54
Switzerland	25	6.5	0	10	69.4	43.5	0	N/A	N/A	76	38.2	21
U.S.A.	29	74.4	31	33	109.9	85	1	89.0	89	34	76.9	41
Brazil	3	36.0	0	4	67.5	68	1	40.0	40	82	87.4	54
Mexico	1	0.0	0	2	122.0	122	0	N/A	N/A	98	54.6	34

Note: Sample includes all molecules present and all country launches occurring in IMS dataset through 2003

Note: Molecules with multiple local corporations in a country were assigned to a single one in the order of Local Originator, Solo Licensee or Local Co-marketer

Appendix Table A4. Split Population Coefficients and Marginal Effects for Launch Model (Standard Errors in Brackets)

	Coeffi	cients	Margina	l Effects
	Drugs in	Drugs in	Drugs in	Drugs in
	Superior	Inferior	Superior	Inferior
	Subclasses	Subclasses	Subclasses	Subclasses
Log Lagged Avg Price of Superior Brandeds (per Std Unit)	0.1474***	-0.1617***	0.0079***	-0.0002*
	[0.0459]	[0.0613]	[0.0027]	[0.0001]
Log Lagged Avg Price of Inferior Brandeds (per Std Unit)	0.0806	0.1418**	0.0043	0.0001
	[0.0524]	[0.0640]	[0.0028]	[0.0001]
Log Lagged Total Volume of All Drugs in Class (Std Unit)	-0.0813	0.1835**	-0.0043	0.0002*
	[0.0578]	[0.0824]	[0.0033]	[0.0001]
Num Generic Manufs per Molc in Superior Subclass (Lag 1Q)	-0.0046 [0.0055]	0.0060 [0.0085]	-0.0002 [0.0003]	0.0000
Num Generic Manufs per Molc in Inferior Subclass (Lag 1Q)	0.0000	-0.0052*	0.0000	0.0000
	[0.0017]	[0.0027]	[0.0001]	[0.0000]
No Molecules in Superior Subclass D.V.	0.2400	-0.1446	0.0142	-0.0001
	[0.1711]	[0.2410]	[0.0111]	[0.0002]
No Molecules in Inferior Subclass D.V.	-0.5198	0.2670	-0.0219	0.0003
	[0.4650]	[0.5862]	[0.0157]	[0.0007]
Time Since Global Launch (Yrs)	-0.6591***	-0.2872***	-0.0351***	-0.0003**
	[0.0452]	[0.0220]	[0.0054]	[0.0001]
Time Since Global Launch Squared (Yrs)	0.0302***	0.0029*** [0.0003]	0.0016*** [0.0003]	0.0000**
First Global Launch Before 1990 D.V.	-0.3280 [0.2063]	0.6525*** [0.1923]	-0.0150* [0.0090]	0.0009**
First Global Launch in [1996-end] D.V.	0.0124	0.1758	0.0007	0.0002
	[0.1179]	[0.2002]	[0.0063]	[0.0002]
Num Already Launched (Great Britain, Germany)	0.5890***	0.6281*** [0.1198]	0.0329*** [0.0068]	0.0006* [0.0003]
Num Already Launched (Sweden, Netherlands)	0.5162*** [0.0607]	0.5798***	0.0285***	0.0006** [0.0003]
Num Already Launched (Italy, France)	0.3466***	0.1009	0.0188***	0.0001
	[0.0745]	[0.0992]	[0.0049]	[0.0001]
Num Already Launched (Spain, Portugal, Greece)	0.1030*	0.3959***	0.0055*	0.0004**
	[0.0561]	[0.0823]	[0.0031]	[0.0002]
Num Already Launched (Canada, Japan, Switzerland, USA)	0.2392***	0.0023	0.0128***	0.0000
	[0.0467]	[0.0755]	[0.0032]	[0.0001]
Any PI Share in Subclass D.V.	-0.0601	0.0838	-0.0031	0.0001
	[0.1519]	[0.3651]	[0.0080]	[0.0003]
Launch by Local Originator Corporation D.V.	1.6471***	3.2262***	0.1942***	0.0213**

	[0.1631]	[0.2437]	[0.0441]	[0.0107]
Launch by Solo Licensee Corporation D.V.	0.5159***	0.9606***	0.0347***	0.0015*
	[0.1474]	[0.1998]	[0.0131]	[8000.0]
Launch by Local Co-marketer Corporation D.V.	0.4238**	0.7515**	0.0276*	0.0011
	[0.1906]	[0.3469]	[0.0153]	[8000.0]
USD to (ECU or Euro) Exchange Rate	0.1833	-0.9852*	0.0098	-0.0009
	[0.3788]	[0.5444]	[0.0200]	[0.0007]
Great Britain Country D.V.	-0.0926	-0.0730	-0.0047	-0.0001
	[0.2124]	[0.3874]	[0.0107]	[0.0004]
Netherlands Country D.V.	-0.7043***	0.4893	-0.0273***	0.0006
	[0.2214]	[0.4181]	[0.0091]	[0.0005]
Sweden Country D.V.	-0.6790***	0.0153	-0.0266***	0.0000
	[0.2333]	[0.4829]	[0.0098]	[0.0005]
France Country D.V.	-1.2415***	-0.9253*	-0.0387***	-0.0006
	[0.2358]	[0.5316]	[0.0090]	[0.0004]
Greece Country D.V.	-1.2744***	0.9060*	-0.0392***	0.0014*
	[0.2517]	[0.5224]	[0.0094]	[8000.0]
Italy Country D.V.	-0.8575***	-0.1161	-0.0312***	-0.0001
	[0.2231]	[0.4980]	[0.0092]	[0.0005]
Portugal Country D.V.	-2.0003***	-0.0306	-0.0472***	0.0000
	[0.2534]	[0.5357]	[0.0091]	[0.0005]
Spain Country D.V.	-0.7324***	0.2529	-0.0281***	0.0003
	[0.2221]	[0.5129]	[0.0095]	[0.0005]
Canada Country D.V.	-1.0758***	-0.7114	-0.0358***	-0.0005
	[0.2175]	[0.5003]	[0.0090]	[0.0004]
Japan Country D.V.	-2.6683***	0.2334	-0.0509***	0.0003
	[0.2352]	[0.4820]	[0.0088]	[0.0005]
Switzerland Country D.V.	-1.0720***	-0.4324	-0.0357***	-0.0003
	[0.2529]	[0.5640]	[0.0097]	[0.0005]
United States Country D.V.	-0.7689***	-1.1768**	-0.0290***	-0.0007
	[0.2321]	[0.5145]	[0.0088]	[0.0004]
Brazil Country D.V.	-1.0918***	0.2298	-0.0361***	0.0002
	[0.2259]	[0.4912]	[0.0092]	[0.0005]
Mexico Country D.V.	-1.1786***	0.1323	-0.0376***	0.0001
	[0.2473]	[0.5375]	[0.0094]	[0.0005]
Source: Anti-asthma D.V.	0.0427	0.2750	0.0023	0.0003
On the Artholist a DV	[0.2217]	[0.2157]	[0.0122]	[0.0003]
Source: Anti-clotlius D.V.	-1.0787***	-0.0514	-0.0385***	0.0000
Courses Anti-depressents D.V	[0.2925]	[0.3750]	[0.0098]	[0.0003]
Source: Anti-depressants D.V.	0.0636	0.4549*	0.0035	0.0005
Course Anti enilentie D.V	[0.1773] -0.1856	[0.2618] 0.1299	[0.0098]	[0.0004]
Source: Anti-epileptic D.V.			-0.0093	0.0001
Source: Anti-nauseants D.V.	[0.1832] -1.0344***	[0.6761] -0.9279	[0.0091]	[0.0007] -0.0006
Source. Armi-Hauseams D.V.			-0.0376***	
Source: Anti-parkinsonian D.V.	[0.2803] -0.3043	[0.6493] 0.7702*	[0.0097] -0.0143	[0.0004] 0.0011
Jourge. Anti-parkinsonian D.V.	[0.2534]	[0.4077]	[0.0112]	[0.0009]
Source: Anti-psychotics D.V.	-0.5169*	2.2627***	-0.0223*	0.0009
Oddice. Anti-payonolica D.V.	[0.2964]	[0.5165]	[0.0117]	[0.0049]
Source: Anti-ulcerants D.V.	-0.4654**	-0.2850	-0.0211**	-0.0002
Ource. Anti-ulcerants D.V.	-0.4054	-0.2000	-U.UZII	-0.0002

	[0.1896]	[0.3830]	[0.0084]	[0.0003]
Source: Lipid lowering D.V.	-0.0910	0.6765*	-0.0047	0.0009
	[0.2037]	[0.3561]	[0.0102]	[0.0007]
Source: Migraine D.V.	-0.7104**	-0.1054	-0.0281**	-0.0001
	[0.3355]	[0.8103]	[0.0115]	[0.0007]
Source: Osteoporosis D.V.	-0.3858*	0.8537***	-0.0177*	0.0011
	[0.2186]	[0.2539]	[0.0091]	[0.0007]
Constant	-1.1655	-5.8524***		
	[0.9921]	[1.5402]		
Ancillary Split-population Parameter	-2.9734***	-1.0464***		
	[0.2709]	[0.1999]		
Observations	23.400	96.041	23.400	96,041
Number of Molecule-level Clusters	111	239	111	239
Model Log-Likelihood	-3047.3	-1994.69		
Mean of Dependent Variable	0.0378	0.0041	0.0378	0.0041

Standard errors in brackets
* significant at 10%; ** significant at 5%; *** significant at 1%

Appendix Table 5. Determinants of Launch Prices, Heckit Regressions (Standard Errors in Brackets)

	ML Heckr	man Probit w/	Molecule Clus	2-Step I	2-Step Heckman Cloglog w/ Consistent S			
	Drugs from Superior	Drugs from Inferior	Drugs from Superior	Drugs from Inferior	Drugs from Superior	Drugs from Inferior	Drugs from Superior	Drugs Infe
Variables	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses	Subclasses	Subcl
Superior Brandeds Price Missing D.V.	-0.0439	0.0020	-0.0460	0.0083	-0.0408	0.0025	-0.0429	0.0
	[0.1820]	[0.1154]	[0.1829]	[0.1163]	[0.1277]	[0.1119]	[0.1277]	[0.1
Log Lagged Avg Price of Superior Brandeds (per Std								
Unit)	0.1540***	-0.0006	0.1552***	0.0051	0.1538***	0.0014	0.1551***	0.00
	[0.0390]	[0.0437]	[0.0391]	[0.0449]	[0.0187]	[0.0317]	[0.0186]	[0.0
Inferior Brandeds Price Missing D.V.	0.1015	-0.7081	0.1062	-0.7599	0.1037	-0.7009	0.1084	-0.7
	[0.1526]	[0.5937]	[0.1539]	[0.6152]	[0.1374]	[0.4338]	[0.1374]	[0.43
Log Lagged Avg Price of Inferior Brandeds (per Std								
Unit)	0.0420	0.2176***	0.0413	0.2159***	0.0415**	0.2148***	0.0408**	0.212
	[0.0394]	[0.0447]	[0.0392]	[0.0446]	[0.0180]	[0.0302]	[0.0180]	[0.0
Generics Price Missing D.V.	0.0500	0.3676**	0.0515	0.3715**	0.0543	0.3748***	0.0558	0.379
	[0.1089]	[0.1833]	[0.1087]	[0.1846]	[0.0746]	[0.1314]	[0.0746]	[0.13
Log Lagged Avg Price of Generics in Class (per Std								
Unit)	0.0296	-0.1143***	0.0297	-0.1150***	0.0301	-0.1149***	0.0303	-0.11
	[0.0289]	[0.0398]	[0.0289]	[0.0398]	[0.0224]	[0.0372]	[0.0224]	[0.0
Time Since Global Launch (Yrs)	-0.0131	-0.0081	-0.0117	-0.0084	-0.0136	-0.0074	-0.0123	-0.0
	[0.0354]	[0.0215]	[0.0356]	[0.0217]	[0.0268]	[0.0171]	[0.0268]	[0.0
Time Since Global Launch Squared (Yrs)	0.0018	-0.0008	0.0018	-0.0007	0.0019	-0.0008*	0.0018	-0.0
	[0.0019]	[0.0006]	[0.0019]	[0.0006]	[0.0017]	[0.0005]	[0.0017]	[0.00
First Branded Launch in Ctry-Subclass D.V.	0.1568	0.8555**	0.1604	0.8849**	0.1545	0.8468**	0.1582	0.87
	[0.1544]	[0.3447]	[0.1546]	[0.3673]	[0.1302]	[0.3904]	[0.1302]	[0.39
Second Branded Launch in Ctry-Subclass D.V.	0.3461***	0.5935**	0.3450***	0.5841**	0.3494***	0.5930***	0.3483***	0.582
	[0.0814]	[0.2820]	[0.0814]	[0.2935]	[0.0613]	[0.2021]	[0.0613]	[0.20
Third or Fourth Branded Launch in Ctry-Subclass D.V.	0.2329***	0.3158**	0.2316***	0.3125**	0.2355***	0.3142***	0.2342***	0.310
Third of Fourth Branded Laurich in City-Subclass D.V.								
High price ELLMin Own Price Missing D.V	[0.0611] 0.2807***	[0.1526]	[0.0613] 0.2774***	[0.1544]	[0.0490] 0.2746***	[0.1125]	[0.0490] 0.2713***	[0.1]
High-price EU Min Own Price Missing D.V.		0.0478		0.0654		0.0458		0.00
Log(Min Own Brigg) in Hi Brigg ELL (HSD/CLL)	[0.0962]	[0.1468]	[0.0959]	[0.1504]	[0.0687]	[0.1042]	[0.0686]	[0.10
Log(Min Own Price) in Hi-Price EU (USD/SU)	0.2221***	0.3925***	0.2216***	0.3976***	0.2227***	0.3940***	0.2222***	0.399

Low-price EU Min Own Price Missing D.V.	[0.0615]	[0.0781]	[0.0614]	[0.0782]	[0.0287]	[0.0497]	[0.0287]	[0.0
	0.0248	-0.0269	0.0271	-0.0207	0.0242	-0.0262	0.0266	-0.0
	[0.0648]	[0.1185]	[0.0653]	[0.1195]	[0.0579]	[0.1065]	[0.0579]	[0.1
Log(Min Own Price) in Low-Price EU (USD/SU)	-0.0247	-0.1081	-0.0238	-0.1010	-0.0254	-0.1121	-0.0245	-0.1
	[0.0387]	[0.1097]	[0.0384]	[0.1088]	[0.0303]	[0.0690]	[0.0303]	[0.0
High-price non-EU Min Own Price Missing D.V.	0.1333*	-0.2960***	0.1355*	-0.3035***	0.1355**	-0.2909***	0.1377**	-0.29
	[0.0713]	[0.1031]	[0.0716]	[0.1042]	[0.0587]	[0.0894]	[0.0587]	[0.0
Log(Min Own Price) in Hi-Price non-EU (USD/SU)	0.2693***	0.0454 [0.0622]	0.2687*** [0.0505]	0.0499 [0.0605]	0.2698*** [0.0255]	0.0453 [0.0512]	0.2692*** [0.0255]	0.0 [0.0]
Any PI Share in Subclass D.V.	0.0238	-0.5268***	0.0251	-0.4897**	0.0252	-0.5263**	0.0265	-0.48
	[0.0771]	[0.1889]	[0.0761]	[0.1965]	[0.0827]	[0.2113]	[0.0827]	[0.2
Log of GDP per Capita	0.8665 [0.9101]	2.6856* [1.4797]	-		0.8658 [0.9173]	2.6565** [1.3535]		-
Launch by Local Originator Corporation D.V.	-0.0649	-0.4217**	-0.0627	-0.4594**	-0.0493	-0.4502***	-0.0471	-0.48
	[0.1264]	[0.2026]	[0.1265]	[0.2084]	[0.0934]	[0.1566]	[0.0934]	[0.1
Launch by Solo Licensee Corporation D.V.	-0.1154	-0.2653*	-0.1120	-0.2686*	-0.1082	-0.2580**	-0.1048	-0.25
	[0.0850]	[0.1483]	[0.0847]	[0.1525]	[0.0799]	[0.1251]	[0.0798]	[0.1
Launch by Local Co-marketer Corporation D.V.	-0.0748	-0.2988	-0.0739	-0.3404	-0.0640	-0.2748	-0.0632	-0.3
	[0.1069]	[0.2293]	[0.1079]	[0.2371]	[0.1007]	[0.1846]	[0.1007]	[0.1
USD to (ECU or Euro) Exchange Rate	-0.1009	2.6998***	-0.0908	2.5672**	-0.1111	2.7060***	-0.1009	2.57
	[0.6026]	[1.0292]	[0.5929]	[1.0368]	[0.5481]	[0.9399]	[0.5483]	[0.9
Country-Specific Quarterly Producer Price Index	-0.0085**	0.0037	-0.0061	0.0114*	-0.0087	0.0033	-0.0063	0.0
	[0.0041]	[0.0080]	[0.0038]	[0.0062]	[0.0054]	[0.0072]	[0.0047]	[0.0
Avg Pack Size (Up to 100)	-0.0117***	-0.0102***	-0.0117***	-0.0100***	-0.0117***	-0.0101***	-0.0117***	-0.00
	[0.0017]	[0.0024]	[0.0017]	[0.0024]	[0.0010]	[0.0018]	[0.0010]	[0.0]
Pack Size > 100 D.V.	-1.1988***	-1.5944***	-1.2007***	-1.5613***	-1.1982***	-1.5919***	-1.2000***	-1.55
	[0.1820]	[0.1965]	[0.1830]	[0.1928]	[0.1291]	[0.1766]	[0.1291]	[0.1
Avg Pill Strength (g)	0.4911**	0.0584	0.5048**	0.0537 [0.0663]	0.4991***	0.0594 [0.1073]	0.5128***	0.0 [0.1
Form: Oral Solid Delayed D.V.	-0.0508	-0.2028	-0.0588	-0.1720	-0.0670	-0.2036	-0.0748	-0.1
	[0.1927]	[0.1390]	[0.1917]	[0.1408]	[0.2057]	[0.1335]	[0.2056]	[0.1
Form: Injectable D.V.	2.0887***	1.7359***	2.0958***	1.7220***	2.0867***	1.7400***	2.0939***	1.72

	[0.2931]	[0.3268]	[0.2949]	[0.3259]	[0.0876]	[0.2007]	[0.0873]	[0.20
Form: Other	-0.0446	0.3067***	-0.0475	0.3063***	-0.0429	0.3054***	-0.0458	0.30
	[0.1739]	[0.1018]	[0.1719]	[0.1027]	[0.1193]	[0.0939]	[0.1194]	[0.09
Britain Country D.V.	-0.3076***	-0.1866	-0.2605***	-0.0713	-0.3140***	-0.1867	-0.2670***	-0.0
	[0.0920]	[0.1277]	[0.0798]	[0.1171]	[0.1127]	[0.2028]	[0.1012]	[0.19
Netherlands Country D.V.	-0.0330	-0.0385	-0.0358	-0.0884	-0.0390	-0.0403	-0.0418	-0.0
	[0.1037]	[0.1476]	[0.1028]	[0.1423]	[0.1044]	[0.2056]	[0.1044]	[0.20
Sweden Country D.V.	-0.1475	-0.1482	-0.0265	0.2114	-0.1465	-0.1485	-0.0256	0.20
	[0.1633]	[0.2709]	[0.0835]	[0.1993]	[0.1678]	[0.2803]	[0.1084]	[0.2
France Country D.V.	-0.1466	-0.7581***	-0.1795	-0.8343***	-0.1514	-0.7618***	-0.1843	-0.83
	[0.1171]	[0.2498]	[0.1136]	[0.2502]	[0.1302]	[0.2892]	[0.1255]	[0.29
Greece Country D.V.	0.3305	1.0752	-0.3469***	-0.9645***	0.3232	1.0454	-0.3535***	-0.97
	[0.7123]	[1.1265]	[0.1165]	[0.2406]	[0.7294]	[1.0664]	[0.1337]	[0.28
Italy Country D.V.	-0.1485	-0.3721	-0.3079***	-0.8155***	-0.1501	-0.3928	-0.3094**	-0.83
	[0.1934]	[0.3383]	[0.0992]	[0.2389]	[0.2093]	[0.3590]	[0.1238]	[0.28
Portugal Country D.V.	0.4509	1.3193	-0.2355**	-0.7982***	0.4461	1.2893	-0.2397*	-0.80
	[0.7261]	[1.1875]	[0.1136]	[0.2327]	[0.7399]	[1.1055]	[0.1390]	[0.28
Spain Country D.V.	0.2250	0.7180	-0.2212**	-0.6554***	0.2188	0.6871	-0.2271*	-0.67
	[0.4806]	[0.7975]	[0.1000]	[0.2184]	[0.4871]	[0.7491]	[0.1187]	[0.28
Canada Country D.V.	0.0718	-0.3104	0.0641	-0.3316	0.0675	-0.3110	0.0598	-0.3
	[0.1093]	[0.2641]	[0.1068]	[0.2662]	[0.1186]	[0.2858]	[0.1183]	[0.28
Japan Country D.V.	0.2672	-0.9519	0.7064***	0.4279*	0.2639	-0.9265	0.7027***	0.43
	[0.4899]	[0.7753]	[0.2141]	[0.2573]	[0.4908]	[0.7530]	[0.1574]	[0.29
Switzerland Country D.V.	-0.1155	-1.5948**	0.2371**	-0.4716**	-0.1200	-1.5708**	0.2324**	-0.4
	[0.3986]	[0.6504]	[0.0930]	[0.2325]	[0.3904]	[0.6319]	[0.1142]	[0.28
United States Country D.V.	0.2117	-0.7338	0.5569***	0.3505	0.2071	-0.7150	0.5520***	0.3
	[0.3981]	[0.6103]	[0.1398]	[0.2414]	[0.3852]	[0.6160]	[0.1218]	[0.28
Brazil Country D.V.	1.3378	4.3471	-0.2695**	-0.5864***	1.3316	4.2883*	-0.2743**	-0.59
	[1.6745]	[2.7229]	[0.1140]	[0.2229]	[1.7056]	[2.5018]	[0.1195]	[0.2]
Mexico Country D.V.	0.9822	2.6749	-0.1994*	-0.9558***	0.9780	2.6347	-0.2027*	-0.95
	[1.2479]	[2.0068]	[0.1196]	[0.2136]	[1.2567]	[1.8511]	[0.1213]	[0.28
Constant	-7.0641	-28.7925**	1.2740	-2.7248*	-7.0475	-28.5116**	1.2834	-2.72
	[8.6436]	[14.3425]	[0.8686]	[1.4717]	[8.8706]	[13.2053]	[0.8744]	[1.3
Inverse Mills Ratio	-0.1612	-0.2277	-0.1610	-0.2476	-0.1503**	-0.2215*	-0.1502**	-0.23
	[0.1197]	[0.1817]	[0.1197]	[0.1887]	[0.0703]	[0.1164]	[0.0703]	[0.1
			•					-

Log GDP per Capita Included?	Yes	Yes	No	No	Yes	Yes	No	N
Year Fixed Effects Included?	Yes	Y						
Observations	23,465	96,074	23,465	96,074	23,465	96,074	23,465	96,

Standard errors in brackets
* significant at 10%; ** significant at 5%; *** significant at 1%