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*Private versus Social Incentives for
Pharmaceutical Innovation*

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Department of Economics

Private versus Social Incentives for Pharmaceutical Innovation*

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Abstract

There is a great deal of debate in society regarding the tendency of pharmaceutical companies to direct their R&D toward marketing products that are “follow-on” drugs of already existing drugs, rather than the development of breakthrough drugs. This paper provides a theoretical framework to study firm incentives for pharmaceutical innovation that disentangle the quest for breakthrough drugs from the firm effort to develop follow-on drugs. We construct a model with a population of patients treated with one of two –horizontally and vertically differentiated– drugs. One of the drugs is the pioneer; the other is the result of an innovative process by a firm that seeks to achieve an improvement over the existing drug. Our results offer theoretical support for the conventional wisdom that pharmaceutical firms devote too many resources to conducting R&D activities that lead to incremental innovations.

Keywords: pharmaceuticals, R&D activities, me-too drugs, breakthrough drugs, incremental innovation, radical innovation.

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1 Introduction

It is widely recognized that of all industrial sectors, the pharmaceutical industry is the sector that traditionally invests most heavily in research and development (R&D). In the United States, R&D investments of pharmaceutical companies have grown consistently over the past 15 years. In 2012, for instance, US biopharmaceutical research companies invested an estimated \$48.5 billion in R&D (PhRMA, 2013). Regarding R&D intensity, and according to a recent report by the European Commission, spending on R&D in 2012 by the pharmaceutical industry amount to 15.3% of its GDP in the US, to 16.3% in Japan and to 14.7% in the European Union (European Commission, 2013).

However, there is a great deal of debate surrounding pharmaceutical R&D activities. Pharmaceutical companies are often accused of devoting too many resources toward the marketing of apparent new products that are “follow-on” drugs of already existing drugs, rather than the development of breakthrough (first-in-class) drugs.¹ In fact, a successful new first-in-class drug will often face competition from a series of follow-on drugs that are therapeutically similar to the pioneering drug. The angiotensin converting enzyme (ACE) inhibitors, a class of drugs used to manage high blood pressure, is illustrative of this. The first ACE inhibitor, captopril, was introduced in the US in 1981. Since then, over 10 ACE inhibitors have been launched (Hernandez and Harrington, 2008).² The development of follow-on drugs is cheaper and less risky than drugs with a novel mechanism of action, but they supposedly do not bring significant therapeutic progress to patients (see, for instance, the discussions by Angell, 2004; Avorn, 2004, and Goozner, 2004). Defenders of incremental innovations argue, however, that medicines based on incremental improvements often represent advances in safety and efficacy, along with providing new formulations and dosing options that increase patient compliance (see diMasi and Paquette, 2004; Wertheimer and Santella, 2009; and the recent article by Doctor Henry I. Miller in the *Wall Street Journal* on January 1, 2014).

This paper aims at contributing to this social debate. We build a theoretical model of innovation to investigate whether there exist arguments that allow us to support the conviction that pharmaceutical firms devote too many resources to market me-too drugs and too few to launch breakthrough drugs. Our model emphasizes the distinction between radical and incremental innovation processes.³ Radical innovation processes may lead to

¹Follow-on drugs are sometimes called “me-too” drugs as they are just copies of existing drugs.

²Another example is omeprazole, the first proton pump inhibitor launched in 1989 to reduce gastric acid production. Proton pump inhibitors have since become the mainstay of treatment for acid-related gastrointestinal disease in adults, and omeprazole was followed by other proton pump inhibitors, with the last launched in 2009.

³The labels “radical” and “incremental” belong mostly to the managerial literature and does not offer a unique description of the difference between the two concepts. In fact, the literature reveals that the

breakthrough drugs, while incremental innovation processes pursue me-too drugs.

In our model there is a continuum of patients in need of medical treatment. Patients can be treated with drugs that are horizontally and vertically differentiated. Vertical differentiation refers to the quality of the drug and includes the health gains experienced by patients. Horizontal differentiation reflects the adequacy of the drug for patients, as different patients in the population will experience different effects of a given medication in terms of tolerability, side effects or interaction with other medicines. In the market, there is a pioneering drug. Moreover, a new drug can be marketed as the result of an innovative process by a pharmaceutical firm that seeks to achieve an improvement over the existing medicine. Finally, there is a physician who makes drug prescription decisions. The physician acts as a perfect agent for the patients and the health authority and, hence, he makes prescription choices based on the price-effectiveness of the drugs.

In this simple set-up we characterize physician prescription choices, given the prices and the characteristics of the two drugs (when the innovation process is successful), and the optimal pricing decision of the innovative firm. The optimal price for the new drug depends on the differences in cost-effectiveness and the horizontal distance between the new drug and the pioneer. When the new drug is much more cost-effective than the pioneer, the innovative firm sets a price such that the new drug will be prescribed to all patients. When the improvements in the cost-effectiveness of the new drug are not substantial, or the two drugs are very horizontally differentiated, then the price set by the innovative firm leads to a drug replacement treatment only for some patients. In all other situations the new drug is not marketed.

We also characterize the incentives of the innovative firm to conduct R&D activities and compare these private incentives with those that would be optimal from a social point of view. The paper distinguishes between radical innovation processes, seeking breakthrough drugs, and incremental innovation processes that aim at launching a me-too drug. In order to differentiate these two kinds of innovations, we follow the approach of measuring the degree of innovativeness of a drug as the size of the differences (either small or large) between the new drug and the pioneer. These differences can emerge either in the horizontal or the vertical characteristics of the drugs. Innovations in the vertical dimension imply a better quality of treatment (or a lower production cost) for all the patients suffering from the disease.⁴ Horizontal innovations would be advances that

definitions of radical and incremental innovations are still puzzling, both at the theoretical and at the empirical level (see García and Calentone, 2002, for a critical review of the innovativeness terminology). In particular, the degree of innovativeness of a product is measured using various dimensions including the level of risk implied in the innovation strategy, the type of knowledge to be processed or the level of investment needed to move onto a new trajectory.

⁴Examples of innovations that would be classified as vertical in our model would include the afore

benefit some but not all patients, because drugs may have lower side effects for a certain group of patients.⁵ Moreover, in order to account for the fact that the level of risk (or uncertainty of the final outcome) is typically larger in the case of radical innovations, we consider that the outcome of a radical innovation process by the innovative firm takes values on a large support and has a greater variance.

The paper provides some interesting findings. We show that for incremental innovation processes pursuing me-too drugs, the social value of the innovation coincides with the private benefits of the firm (as the innovative firm appropriates all the patients' benefits derived from the launching of the me-too drug). If we consider, instead, R&D activities searching for breakthrough drugs, then private and social incentives for conducting research are not aligned. In particular, the incentives for conducting research by the firm are inferior to those socially optimal as there are patients that, despite the larger price of the new drug, benefit from it. These results allow us to show that if a pharmaceutical company can only adopt one of the two types of innovation processes due, for instance, to budget constraints, it may happen that the firm has an incentive to seek a me-too drug as R&D activities oriented to search for a radical innovation are socially superior. At the same time, it never happens that the innovative firm prefers to develop a radical innovation when devoting the resources to incremental innovations is preferable from a social point of view. Our results thus offer theoretical support for the conventional wisdom that pharmaceutical firms devote too many resources to conducting R&D activities that lead to me-too drugs.

The theoretical literature on incentives for pharmaceutical innovations is not abundant, although there is an increasing number of papers that study the interaction between the pricing policy constrained by various forms of regulation and the effort of innovation by pharmaceutical firms. Ganuza, Llobet, and Dominguez (2009) find a bias in the pharmaceutical industry toward small innovations. Their result relies on the low sensitivity mentioned captopril (ACE-inhibitor) and omeprazole (proton pump inhibitor), and also cimetidine (H₂-receptor antagonist), propranolol (β adrenoceptor- antagonist), lovastatin (HMG-CoA-reductase inhibitor), and sumatriptane (5-HT_{1B/1D}-receptor agonist) among others. All these are drugs that, when marketed, met a given need much more effectively than available treatments and were beneficial for all patients in the treatment of their disease. Also, innovations in antibiotics that allow administration once a day, giving patients the possibility of being treated at home, or at least the possibility to reduce hospitalization time are vertical innovations according to our classification. Finally, second-generation antihistamines have some (vertical) improvements over first generation antihistamines like, for instance, less frequent dosing.

⁵For example, in the market for statins, Lovastatin, pravastatin, and fluvastatin represent the class members with the lowest potency to reduce cholesterol levels but are attractive candidates for use in patients who have proven intolerant of more potent statins such as atorvastatin, simvastatin, or rosuvastatin (Kapur and Musunuru, 2008).

of a part of the demand (due to the loyalty of some physicians) to changes in prices. This lack of price-sensitivity provides an excessive reward for small innovations and consequently distorts downwards the incentives of pharmaceutical firms. In our model, the physician acts as a perfect agent for the patients, so that the difference between the social value and the private benefits that the firm obtains from innovation arises from a different source: the ability of the pharmaceutical firm to appropriate or not the patient surplus through the price. The existence of physicians that are loyal to innovative drugs also play an important role in Antoñanzas, Juárez-Castelló, and Rodríguez-Ibeas (2011). They study the incentives of an incumbent pharmaceutical firm to launch an upgraded drug through innovation before it faces generic competition. The paper shows that the equilibrium level of innovation exhibits an inverted U shape, as innovation increases when the proportion of loyal physicians is low and decreases when it is high. Finally, Bardey, Bommier, and Jullien (2010) focus on the long-run impact of reference pricing on pharmaceutical innovation by firms. Their model share some similarities with ours as it makes a clear distinction between incremental and radical innovations in a setting where drugs are horizontally and vertically differentiated. However, the distinction they make between the two types of innovation differs notably from ours. In addition to this, they model a patent race where the innovative process is deterministic and competition in R&D leads to the dissipation of firms' profits. They show that the short-term and long-term effects of price regulation may be antagonists. In their simulation using French data, they find that favoring radical innovation processes at the expense of cost-reducing innovations may generate medium/long-run increases in health expenses, despite potential short-run benefits.

The rest of the paper is organized as follows. The following section presents our model. Section 3 studies prescription decisions by the physician. Section 4 analyzes pricing decisions by the innovative firm and characterizes the market equilibrium. In section 5 we discuss the benefit for patients derived from the launch of the new drug. Section 6 discusses private versus social incentives to innovate. Section 7 proposes a simple model of innovation. Finally, the last section offers some concluding remarks. All the proofs are in the Appendix.

2 The Model

There is a continuum of patients in need of medical treatment. We normalize the size of the population of patients to 1. Patients suffer from the same illness and are identified by a horizontal characteristic x , with x distributed uniformly on the interval $[0, 1]$. The parameter x represents the type of the patient and measures the heterogeneity regarding

the patient genotype or any other characteristic that may induce the disease to have different effects among patients.

Patients can be treated with a drug. We consider that drugs are both horizontally and vertically differentiated. Thus, a drug is defined by a pair of characteristics: (\hat{x}, \hat{h}) . The first characteristic $\hat{x} \in [0, 1]$ captures the horizontal differentiation and reflects the adequacy of the drug for different patients. It is widely recognized that different patients in a population experience different effects of a given medication in terms of tolerability, side effects or interaction with other medicines.⁶ In our model, where patients are distributed along the interval $[0, 1]$, a particular location (type) of a patient reflects the patient's ideal drug. That is, a drug with characteristic \hat{x} is the ideal drug for a patient located at \hat{x} . Those patients who fail to obtain their ideal drug face a cost, and the farther patient type x from \hat{x} is, the lower the benefits he enjoys (or the larger the side effects he suffers) when he is exposed to the drug.⁷

The second characteristic of the drug incorporates the vertical differentiation and it refers to the gross effectiveness \hat{h} of the drug. This is a quality dimension that affects the whole population and it includes the health gains experienced by patients (which may comprise one or both quality and quantity of life). We assume $\hat{h} > 0$, where $\hat{h} = 0$ would mean that the drug has the same effect as no treatment, and the higher \hat{h} the better the drug for all the patients.⁸

In order to determine the health gain of a type- x patient treated with drug (\hat{x}, \hat{h}) , we need to consider both dimensions together. The health gain of a patient of type x when drug (\hat{x}, \hat{h}) is prescribed is $b(\hat{h} - l|\hat{x} - x|)$, where $|\hat{x} - x|$ is the distance between the horizontal characteristic of the drug and the type of the patient, $l > 0$ scales the loss of effectiveness or the extend of side effects and b is the marginal utility of being healthy.

If we denote by \hat{p} the price of drug (\hat{x}, \hat{h}) , the benefit to the health system of treating patient x with drug (\hat{x}, \hat{h}) and paying price \hat{p} for this drug is⁹

$$H(x; \hat{x}, \hat{h}, \hat{p}) = b(\hat{h} - l|\hat{x} - x|) - \hat{p}.$$

⁶For example, in the pharmaceutical market for blood pressure control, the drugs available to treat hypertension may act via the central nervous system, the heart (beta blockers), the kidney (diuretics, saluretics) or the vessels (alpha blockers, ACE inhibitors, AT1 and calcium antagonists). The efficacy and side effects of these medicines differ across patients and, hence, affect physician prescription patterns.

⁷There is no drug that unambiguously dominates another on the horizontal dimension, as patients with a different x react differently to a drug \hat{x} . If two drugs with a different \hat{x} are available, there are patients that benefit more from one of the drugs, while others would be better off when treated with the other.

⁸In Bardey et al. (2010) drugs are also vertically and horizontally differentiated, although their vertical characteristic is binary while ours is a continuous variable (which allows us to consider small and large differences in the vertical dimension).

⁹For the sake of convenience in the exposition, we will refer to the benefit to the health system of treating a patient as the patient benefit.

We assume that there is a pre-existing drug (x^0, h^0) , $x^0 \in (0, 1)$, on the market. The price p^0 of this pioneer drug is exogenously fixed and does not react to the launch of a new medicine. In markets where drug (x^0, h^0) is produced by a number of firms, this assumption can be easily justified on the grounds that the exclusivity of the pioneering drug has already expired and a generic drug has entered the market.¹⁰ Without loss of generality, we assume that $p^0 = c^0$, being c^0 the marginal cost of providing the drug. For simplicity, we assume that all patients benefit from the pioneering drug, that is, the whole market is initially covered by that drug.¹¹

There is a pharmaceutical firm (that we call firm 1), different from the one selling (x^0, h^0) , that can undertake an innovation process.¹² This process is uncertain both because it may or may not lead to a new drug and because the characteristics of the potential new drug are ex-ante random. If firm 1 adopts an innovation process and the process is successful then a new drug is discovered. We denote by (x^1, h^1) the characteristics of the new drug that will be marketed and by c^1 its marginal cost. We assume that the firm producing the new drug freely chooses the price p^1 .

To identify the differences between the new drug and the pioneer, we define

$$\begin{aligned}\Delta_x &\equiv |x^1 - x^0|, \\ \Delta_h &\equiv h^1 - h^0, \\ \Delta_c &\equiv c^1 - c^0.\end{aligned}$$

That is, $\Delta_x \in \mathbb{R}_+$ is the distance between the “types” of the drugs. Similarly, $\Delta_h \in \mathbb{R}$ indicates the difference in quality between the two drugs, and $\Delta_c \in \mathbb{R}$ denotes the difference in marginal costs (the new drug may be more expensive or cheaper to produce than the pioneer).

Finally, there is a physician who makes drug prescription decisions. We assume that the physician acts as a perfect agent for both the patients and the health authority. Hence, the physician assigns the medication to patients based on the price-effectiveness of the two drugs.¹³ Thus, in our model, the patients, the physician, and the health system can

¹⁰There is evidence that once generic drugs enter the market, both the price and sales revenue of pioneering drugs tend to drop by about 80% over the next year (Yin, 2012).

¹¹This hypothesis requires that $b(h^0 - l|x^0 - x|) - c^0 \geq 0$ for all $x \in [0, 1]$, that is, $c^0 \leq \text{Min} \{b(h^0 - l(1 - x^0)), b(h^0 - lx^0)\}$.

¹²For simplicity in the exposition, we have adopted the view of an entrant in the pharmaceutical market launching a new drug. All our analysis would remain valid if, alternatively, we had assumed that there were several firms producing the pioneering drug and that one of them could undertake the innovation process, as in Ganuza et al. (2009).

¹³This is in contrast with Ganuza et al. (2009) where a proportion of physicians are “captured” doctors and prescribe the drug with the highest quality regardless of its price.

be considered as a single agent whose benefits are the patients' health gains and the costs are the price of the drugs.¹⁴

The timing of our game is as follows. In the first stage, firm 1 decides its innovation strategy. In case the innovation process succeeds, the characteristics (x^1, h^1) as well as the marginal cost c^1 of the new drug are known. In the second stage, the innovative firm sets the price p^1 it charges. Finally, in the third stage, provided the new drug is marketed, the physician allocates the drugs to patients (if only the pioneering drug is available, the physician will prescribe it to all the patients). As usual, we solve the game by backward induction.

3 Prescription of Drugs

We now analyze the last stage of the game if a new drug with characteristics (x^1, h^1) has been marketed at a price p^1 . At this stage of the game, the physician decides which patients are prescribed the pioneering drug (x^0, h^0) and which ones the new one (x^1, h^1) , given the prices and the characteristics of the two drugs. The physician takes into account both the expected effectiveness of the drugs and their price. If both drugs provide identical benefits to the health system when treating a patient, we adopt the convention that the physician prescribes the new drug.

Following the physician's decision, the market will be split between the new drug and the pioneer. Depending on both drugs' characteristics, as well as the price p^1 decided by the innovative firm, three different scenarios may arise. We illustrate them in Figure 1.

First, the physician prescribes the pioneering drug to all patients if the price p^1 of the new drug is very high as compared to its health benefits. We denote p^{\max} the price above which no patient is prescribed the new drug (that happens when even patient x^1 is not treated with (x^1, h^1) if $p^1 > p^{\max}$). Formally, p^{\max} is characterized by $H(x^1; x^1, h^1, p^{\max}) = H(x^1; x^0, h^0, c^0)$, or $b(h^1 - l|x^1 - x^0|) - p^{\max} = b(h^0 - l|x^1 - x^0|) - c^0$, which implies

$$p^{\max} \equiv c^0 + b(\Delta_h + l\Delta_x).$$

Note that p^{\max} is increasing in b , l , Δ_x , Δ_h , and c^0 . Therefore, for a given p^1 , it is more likely that the old drug keeps all the market if health has a low marginal value (b low), the disparities in the side effects of the drug for different patients are small (l low), the difference between the two drugs is also small (Δ_h and Δ_x are low), and/or the pioneering drug is very cheap to produce.

¹⁴This assumption allows us to focus on firm incentives to pharmaceutical innovation, leaving aside any distortion caused by the potential strategic behavior of agents as a consequence of the different views they share.

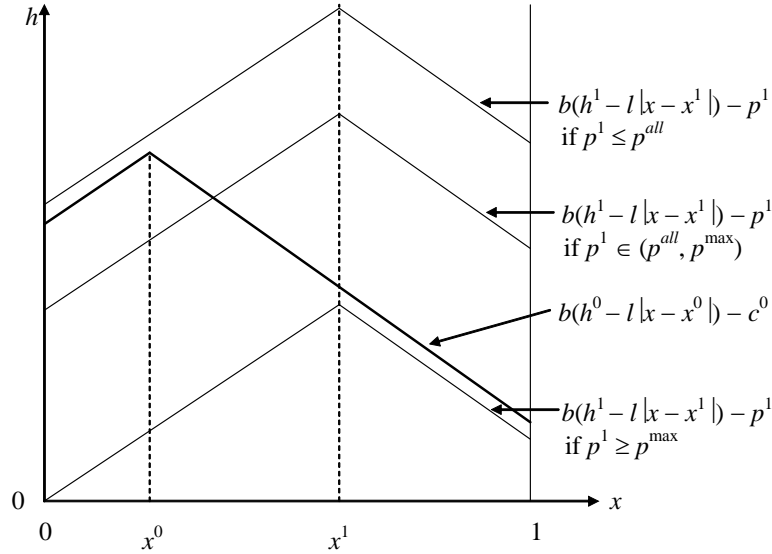


Figure 1: Split of the market as a function of p^1 , c^0 , (x^0, h^0) , and (x^1, h^1) .

If $p^1 = p^{\max}$, the market for the new drug is $1 - x^1$ if $x^1 > x^0$; it is x^1 if $x^1 < x^0$; and it is the whole market if $x^1 = x^0$ (the rest of the patients are prescribed the old drug).

Second, the physician prescribes the new drug to all patients if the price p^1 is low enough. This is the case if p^1 is so low that it is optimal to treat patient x^0 with the new drug. The cut-off value p^{all} below which all patients are treated with (x^1, h^1) is characterized by $b(h^1 - l|x^1 - x^0|) - p^{\text{all}} = b(h^0 - l|x^0 - x^0|) - c^0$, that is,

$$p^{\text{all}} \equiv c^0 + b(\Delta_h - l\Delta_x).$$

Given the expression of p^{all} , a given p^1 is more likely to be lower than p^{all} for large values of c^0 , b , and Δ_h , and for low values of l and Δ_x (i.e., when the difference between the patient who profits more from each drug is low). We note that $p^{\text{all}} = p^{\max}$ if $x^1 = x^0$.

Finally, the physician prescribes each drug to a subset of the patients for $p^1 \in (p^{\text{all}}, p^{\max})$. To identify the market for each drug, consider first the case where $x^1 > x^0$. The two drugs provide the same benefits to a patient of type $\tilde{x} \in (x^0, x^1)$ if $b(h^0 - l(\tilde{x} - x^0)) - c^0 = b(h^1 - l(x^1 - \tilde{x})) - p^1$. That is, the type of the indifferent patient is

$$\tilde{x} \equiv x^0 + \frac{(l\Delta_x - \Delta_h)}{2l} + \frac{(p^1 - c^0)}{2bl}.$$

The physician prescribes treatment (x^0, h^0) at a price c^0 to those patients whose type lies in the interval $[0, \tilde{x})$, whereas patients with type in the interval $[\tilde{x}, 1]$ are treated with (x^1, h^1) at a price p^1 .

Similarly, for $p^1 \in (p^{all}, p^{max})$ and $x^1 < x^0$ the indifferent patient is

$$\tilde{x} \equiv x^0 - \frac{(l\Delta_x - \Delta_h)}{2l} - \frac{(p^1 - c^0)}{2bl}$$

and the markets for the new drug and the pioneer are $[0, \tilde{x}]$ and $(\tilde{x}, 1]$, respectively.

4 The Optimal Pricing Policy

In this section we study the optimal price decision of firm 1 if, as a result of the innovation process, a new drug with characteristics (x^1, h^1) has being achieved.

Firm 1's incentive to price high or low depends on drug differences in terms of vertical characteristics (both in quality, measured by Δ_h , and in marginal cost, measured by Δ_c), and horizontal characteristics (measured by the distance in type Δ_x). The strength of the incentives also depends on the size of the submarket (the subset of patients) in which the innovative firm more directly competes with the pioneering drug. If x^1 is on the right of x^0 (i.e., $x^1 > x^0$), then the competition is fiercer on the right of x^0 than on the left of x^0 . That is, the size of the submarket in which firm 1 directly challenges the pioneering drug is $1 - x^0$. Similarly, if $x^1 < x^0$, the competition is more intense to the left of x^0 , that is, in a submarket of size x^0 . We denote by M the size of this submarket: $M \equiv 1 - x^0$ if $x^1 > x^0$ and $M \equiv x^0$ if $x^1 < x^0$. We also denote $M \equiv 1$ if $x^1 = x^0$ (although M does not play any role when $\Delta_x = 0$).

Proposition 1 summarizes the price decision of the pharmaceutical firm. Except for the region of parameters where there is no room for the firm to make profits, the new drug is always offered at a price in the interval $[p^{all}, p^{max}]$. For convenience, we define

$$p^{int} \equiv \frac{1}{2}(c^0 + c^1) + blM - \frac{1}{2}b(l\Delta_x - \Delta_h),$$

a candidate price for optimum when the optimal price lies in (p^{all}, p^{max}) .

In the proposition, we also use the function $g(\Delta_x)$, which is defined as follows:

$$\begin{aligned} g(\Delta_x) &\equiv \Delta_x + 2 \left(2 - M - 2\sqrt{1 - M} \right) && \text{if } \Delta_x > \sqrt{1 - M} - (1 - M) \\ &\equiv \frac{(1 + M - \Delta_x)}{(1 - M + \Delta_x)} \Delta_x && \text{otherwise.} \end{aligned}$$

The function $g(\Delta_x)$ is increasing, continuously differentiable and fulfils $g(\Delta_x = 0) = 0$.

Finally, as Δ_h and Δ_c have similar effects and they often appear together in the mathematical expressions, we denote the composite effect of these two vertical variables as

$$\Delta_y \equiv \Delta_h - \frac{1}{b}\Delta_c.$$

The parameter Δ_y can be interpreted as a measure of the differences in cost-effectiveness between the new drug and the pioneering one. The larger the value of Δ_y the more cost-effective the new drug is compared to the pioneer.

Proposition 1 *The optimal price decision p^{1*} of firm 1 and its profits $\Pi^1(p^{1*})$ are as follows:*

Region a: *If $\Delta_y \leq -l\Delta_x$ then the new drug is not prescribed: $p^{1*} = c^1$ and $\Pi^1(c^1) = 0$.*

Region b: *If $\Delta_y \in (-l\Delta_x, lg(\Delta_x))$ then the new drug replaces the pioneer for a subset of patients. If in addition*

Region b.i: *$M \geq \frac{1}{2l}(\Delta_y + 3l\Delta_x)$ then $p^{1*} = p^{\max}$ and*

$$\Pi^1(p^{\max}) = b(M - \Delta_x)(\Delta_y + l\Delta_x).$$

Region b.ii: *$M < \frac{1}{2l}(\Delta_y + 3l\Delta_x)$ then $p^{1*} = p^{int}$ and*

$$\Pi^1(p^{int}) = \frac{b}{2l} \left(lM + \frac{1}{2}(\Delta_y - l\Delta_x) \right)^2.$$

Region c: *If $\Delta_y \geq lg(\Delta_x)$, then the new drug takes over the entire market: $p^{1*} = p^{all}$, and*

$$\Pi^1(p^{all}) = b(\Delta_y - l\Delta_x).$$

The optimal profit function Π^1 is continuous in Δ_x, Δ_h and Δ_c .

We now offer some intuitions for Proposition 1. When differences in the cost-effectiveness between the new drug and the pioneer are very negative, that is, for low values of Δ_y , there is no price above the marginal cost under which firm 1 can sell its drug. More precisely, in Region a, where the “aggregate” difference $\Delta_y + l\Delta_x$ between the two drugs is not positive, firm 1 makes zero profits.

On the contrary, in Region c the cost-effectiveness of the new drug is instead larger than that of the pioneering drug. The new drug far outperforms the pioneer and firm 1 decides to set a price for which the new drug is prescribed to all patients. In this case, the price that allows the firm to serve the entire market p^{all} is large enough so that for the firm it is worth setting p^{all} instead of increasing the price further and losing some patients.

Finally, for intermediate values of Δ_y (Region b, which only exists if $\Delta_x > 0$), the optimal price decision by firm 1 depends on the size of submarket M . If M is sufficiently large, then firm 1 sets the maximum price compatible with selling the drug p^{\max} because

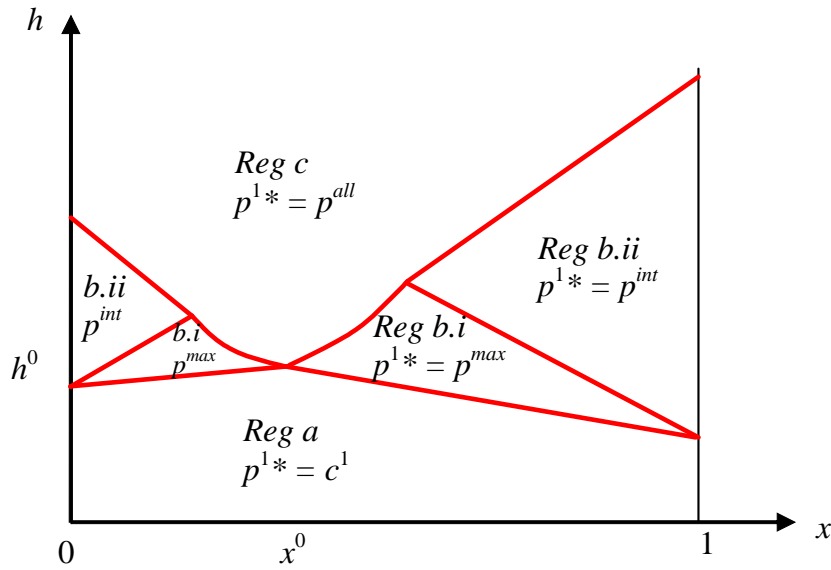


Figure 2: Optimal pricing policy by firm 1.

the number of patients that are treated with the new drug under this price is fairly large. In contrast, for low values of M , firm 1 would sell the drug to too few patients at p^{\max} and, therefore, it prefers to cut the price and it sets p^{int} .

Figure 2 represents the optimal pricing policy by the innovative firm for every combination of (x^1, h^1) , for a given value of Δ_c , b and l . The figure is drawn for $\Delta_c = 0$. For a positive (resp. negative) Δ_c , all the figure would move upwards (resp. downwards) in a proportion $\frac{1}{b}\Delta_c$. Figure 2 is not symmetric with respect to the vertical x^0 because, in this example, $x^0 \neq \frac{1}{2}$.

It is worth noticing that although the optimal price decision p^{1*} of firm 1 is continuously increasing in the quality of the drug (i.e., in Δ_y) inside each region, it is neither continuous nor increasing when Δ_y moves from Region b to Region c. At the border of the two regions, where $\Delta_y = lg(\Delta_x)$, $p^{\max} > p^{\text{all}}$ (in Region b.i) and $p^{\text{int}} > p^{\text{all}}$ (in Region b.ii). Therefore, the price p^{1*} decreases from either p^{\max} or p^{int} to p^{all} if a marginal increase in Δ_h moves the drug from Region b to Region c. On the other hand, the price p^{1*} is continuous in Δ_y (and in Δ_x) at the border of the Region b.i and Region b.ii. Finally, the optimal price p^{1*} increases with the horizontal differentiation Δ_x in Region b.i, but decreases with Δ_x in Region b.ii and Region c.

Corollary 1 provides some comparative statics of firm 1's profits with respect to the level of differentiation between the two drugs, Δ_x and Δ_h . We will use these expressions afterwards in section 7 and subsection 7.3 where we compare the private and social incentives to develop a new drug.

Corollary 1 *The comparative statics of firm 1's profits are as follows:*

- Region b.i: Π^1 is increasing and linear in Δ_y . It is a concave function in Δ_x , increasing up to $\Delta_x = \frac{1}{2l}(lM - \Delta_y)$ and decreasing afterwards.
- Region b.ii: Π^1 is increasing and convex in Δ_y and it is decreasing and convex in Δ_x .
- Region c: Π^1 is increasing and linear in Δ_y and it is decreasing and linear in Δ_x .

Corollary 1 shows that the profits of the innovative firm are always larger the more cost-effective the new drug is, that is, the larger the difference $h^1 - h^0$ and/or the smaller the difference $c^1 - c^0$. Regarding the horizontal characteristic of the drug, firm 1's profits are decreasing in Δ_x , except in Region b.i, where they are increasing for low values of Δ_x .

5 Patients' Benefit

The launch of the new drug allows firm 1 to obtain profits and it may also increase patient surplus. We now move to discuss patient surplus in the different price regions.

Recall that in our model doctors are perfect agents for the patients and the health system. Therefore, they only prescribe the new drug when the surplus of a patient is equal or strictly positive. We denote by ΔCS the gain in patient surplus, net of the price, as a consequence of the launch of the new drug. Proposition 2 provides the expressions for ΔCS in the regions identified in Proposition 1.

Proposition 2 *Given the optimal pricing policy p^{1*} of firm 1, the variation of the patient surplus ΔCS due to the launch of the new drug is:*

Region a: $\Delta CS(c^1) = 0$.

Region b.i: $\Delta CS(p^{\max}) = 0$.

Region b.ii: $\Delta CS(p^{int}) = \frac{b}{16l} (\Delta_y + 3l\Delta_x - 2lM) (\Delta_y - 5l\Delta_x + 6lM)$.

Region c: $\Delta CS(p^{all}) = bl\Delta_x (2M - \Delta_x)$.

Some interesting insights can be extracted from Proposition 2. In the scenario where the new drug is sold at p^{\max} (Region b.i), total patient surplus is the same before and after the launch of the new drug. The reason is that the firm charges the largest potential price, p^{\max} , and it therefore extracts all the surplus from the patients that in equilibrium are prescribed the new drug. In contrast, in Region b.ii, the firm decides to charge $p^{int} < p^{\max}$

to attract some new patients and, thus, patient surplus increases when the new drug is marketed. Finally, in the scenario where drug 1 takes over the market (Region c) patients also benefit from the launch of the new drug when $\Delta_x > 0$. In this case, the health advantages of the new drug outweighs the larger price charged by the innovative firm, so that patients end up strictly better off.

We also provide some comparative statics of the patient surplus with respect to the level of differentiation of the two drugs, Δ_x and Δ_y .

Corollary 2 *The comparative statics of the patient surplus are as follows:*

Regions a and b.i: ΔCS does not change with Δ_y or Δ_x .

Region b.ii: ΔCS is increasing in Δ_y . It is increasing up to $\Delta_x = \frac{1}{15l}(14lM - \Delta_y)$, and then decreasing in Δ_x .

Region c: ΔCS is independent of Δ_y and it is increasing in Δ_x .

Moreover, ΔCS is continuous at the border between Region b.i and Region b.ii but it discontinuously increases when Δ_y increases (or Δ_x decreases) and the drug moves from Region b to Region c.

Corollary 2 shows that the gains derived by the patient from the launch of the new drug depend on the cost-effectiveness of this drug as compared to the pioneering drug (i.e., on Δ_y) only in Region b.ii. In this region, the higher the cost-effectiveness (either due to advantages in quality or costs) of the new drug, the higher the gain in the patient surplus. Patient surplus is certainly more sensitive to the horizontal characteristic Δ_x . In particular, it tends to be increasing in Δ_x , except at the region where the new drug replaces the pioneer for some patients (Region b.ii), where patient surplus is decreasing for high values of Δ_x .

6 Private versus Social Incentives

R&D incentives in our model come from the benefits that the parties involved derive from the launching of the potential new drug. To discuss the difference between private and social R&D incentives, it is worth noting that the innovative firm and the patients have different preferences regarding the characteristics of their best drug, given the pioneering drug on the market. Figure 3 represents the comparative statics of firm 1's profits and consumer surplus with respect to vertical and horizontal changes in the new drug as compared to the pioneer (corollaries 1 and 2).¹⁵ In general, the firm cares a great deal

¹⁵Figure 3 only represents the comparative statics for drugs on the right-hand side of x^0 ; for those drugs on the left-hand side of x^0 the comparative statics are similar. Moreover, the figure does not point out the effects that are zero. For example, in Region c, ΔCS is independent of Δ_y and the figure does not include information about the behavior of CS with respect to Δ_y in that region.

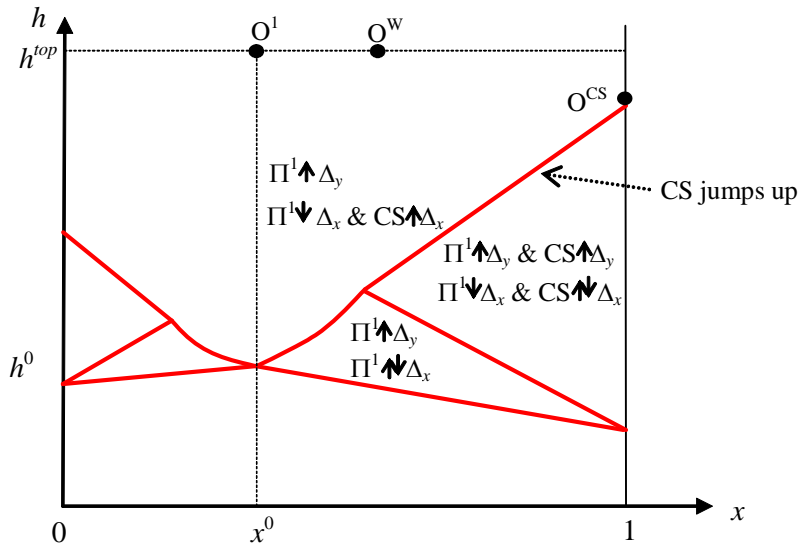


Figure 3: Comparative statics and most preferred drugs.

about increasing the quality of the drug, either through a more effective or a less costly drug, whereas consumers (taking the price into account) only benefit from the launching of a better-quality drug if such a drug is well-suited for those patients far from x^0 (Region b.ii). Additionally, the profits of the innovative firm decrease with the degree of horizontal differentiation between the two drugs, except when the differentiation is small and the quality of the two drugs is similar (the left-hand part of Region b.i). On the contrary, consumers always benefit if new drugs focus on patients for whom the effectiveness of the old drug is low.

Figure 3 helps us to better understand the difference between incentives to innovate for the firm and for a social planner. For example, if the innovative firm could launch a drug through a deterministic process then it would have an incentive to choose a drug with a higher quality but the same horizontal characteristic than the pioneering drug. In fact, if the new drug were available without costs (or at a low cost), the optimal drug for firm 1 would be $O^1 = (x^0, h^{top})$ in Figure 3, where h^{top} is the highest possible quality. In contrast, consumers prefer drugs that are horizontally differentiated from the pioneering drug. In terms of consumer surplus, the optimal drug would be O^{CS} , the farthest from x^0 in the horizontal dimension and at the border of Region c. Finally, if we define the increase in social welfare as the sum of firm 1's profits and the increase in patient surplus,¹⁶ that is, $\Delta W \equiv \Pi^1 + \Delta CS$, then balancing both firm and consumer interests, the optimal

¹⁶We have assumed that the pioneering drug is sold at its marginal cost. Thus, the firm (or any of the firms) selling drug 0 always has zero profits.

drug from a total welfare point of view would be O^W in Figure 3. Notice that the drug O^W improves upon the pioneering drug as much as possible in the vertical dimension and it is also horizontally differentiated from the pioneering drug, although the optimal differentiation is far from the maximal one.

7 A Model of Pharmaceutical Innovation

In this section, we analyze the first stage of our game, the innovation stage, where firm 1 chooses its R&D investment in order to maximize its expected profit. The firm takes into account that its R&D decision will affect the probability of success and the characteristics of the drug that might be marketed at the beginning of the second stage. The innovation strategy of the firm involves both the type of innovative process that the firm wants to adopt and the level of the resources invested in that process. Different types of processes aim at obtaining (with some probability) drugs with different characteristics. We now describe a typology of drugs and innovation processes and then discuss the firm's innovation decision.

7.1 Me-Too versus Breakthrough Drugs

In the pharmaceutical industry, a crucial distinction is made between me-too and breakthrough innovations. The traditional distinction between these two types of innovation relies on the mechanism of action of the drugs. Under this view, a breakthrough innovation would consist of a “first-in-class” medicine, based on a new mechanism of action. In contrast, a me-too (or “follow-on”) innovation would refer to a new drug within an existing class, with a mechanism of action similar to that of the old drug but which differs in features such as therapeutic profile, metabolism, adverse effects, dosing schedules or delivery systems.

Several voices in the pharmaceutical industry have called for the adoption of a broader perspective when evaluating innovation in medicines. The definition of a drug is complex and multidimensional and some advantages of the new drugs (some aspects of value as well as of the costs) are not recognized in the traditional view.¹⁷ We follow this approach. Our interpretation is that the difference between me-toos and breakthrough innovations rests on the distance between the new drug and the pioneer at either the horizontal axis and/or the vertical one. A me-too innovation represents a small change for some (or all) patients either in terms of quality of treatment, cost savings or side effects. For example, a me-too innovation may open the possibility of administering smaller or fewer doses, or

¹⁷See the report by Mestre-Ferrandiz, Mordoh, and Sussex (2012) for a thorough discussion of the nature of innovation in medicines.

it might imply a slightly less invasive delivery (which we interpret as a small increase in y).¹⁸ A me-too innovation may also cause slightly lower side effects for a subpopulation of patients (a small change in x). In contrast, a breakthrough innovation ensures a significant increase in the quality of the new drug or a drug whose characteristics make it well-suited for patients who could not be well treated under the existing treatment.¹⁹

7.2 Incremental versus Radical Innovation Processes

A firm seeking a me-too or a breakthrough innovation would rely on different processes. In this sense, we define an incremental innovation process as a process aiming at a me-too innovation with respect to the existing drug.²⁰ Similarly, when the process pursues a significant improvement (for all or a subpopulation of patients), or a substantial cost reduction as compared to the old drug, then we say that the innovation process is radical. Formally, an incremental innovation process (that we will denote by the sub-index *in*) pursues a drug with a small Δ_x or Δ_y whereas a radical innovation process (denoted by the sub-index *ra*) aims at a drug with a large Δ_x and/or Δ_y .

7.3 A Simple Model of Innovation

To further understand the differences between private and social incentives in terms of innovation, in this subsection we propose a highly stylized model of R&D investment with uncertainty. That is, we analyze the first stage of the game, where firm 1 chooses its innovation strategy.

We assume that the result of the innovation process is uncertain and its outcome can only be poorly predicted, if at all. The uncertainty is greater for ground-breaking and pioneering processes. Thus, we assume that an innovation process may lead (or not) to an innovation and that, even if the process is successful, the specific characteristics of the innovation (i.e., the new drug) are ex-ante random.

There are two types of innovation processes available to the pharmaceutical firm: an incremental innovation process, pursuing a me-too drug, and a radical innovation process, which targets a major innovation as compared to the pioneering drug (x^0, y^0)

¹⁸Insulin pens, for instance, are minimally invasive and have largely superseded the conventional insulin syringe.

¹⁹For instance, omeprazole, the first proton pump inhibitor, proved to be more effective than histamine-2 receptor antagonists to the management of peptic ulcer diseases.

²⁰Sometimes, follow-on drugs are simply the natural outcome of simultaneous research programs into the same therapeutic target (diMasi and Paquette, 2004). In other cases, they are the result of an intentionally imitative research program (Garnier, 2008). The approach adopted in this paper fits better with this second view of the innovation process.

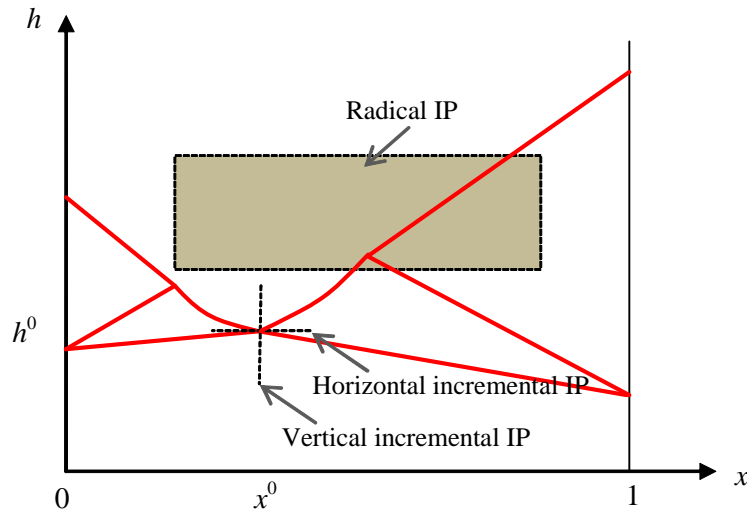


Figure 4: Incremental and radical innovation processes.

(i.e., a breakthrough drug). The incremental innovation process can lead to a drug that is similar to the pioneering drug, in the sense that it would have the same horizontal (or vertical) characteristic, that is, $x^1 = x^0$ (or $y^1 = y^0$) and a minor improvement in quality (or small differences in side effects). On the other hand, the radical innovation process results, in case of success, in a drug (x^1, y^1) where y^1 is random but higher than y^0 and x^1 is also random. This reflects the idea that the innovative process represents a significant departure from the old products or technologies. Moreover, the side effects may also be very different from those of the pioneering drug. We depict the two categories of processes in Figure 4.²¹ The crucial differences between the two categories of processes is that the distribution of the outcome of a radical innovation process takes values on large support and has a greater variance, which leads to higher chances of getting a breakthrough drug (Girotra et al., 2007; Singh and Fleming, 2010).

Let us first consider incremental innovation processes pursuing me-too drugs. If an investment I_{in} is realized, then there is a probability $q_{in}(I_{in})$, increasing in I_{in} , of obtaining an innovation, while no drug is obtained with probability $1 - q_{in}(I_{in})$. If it is a vertical innovation process then, in the case of success, the new drug has characteristics (x^1, y^1) , with $x^1 = x^0$ and y_1 is distributed according to the density function $f_{in}(y^1)$ that takes values in the interval $[y^0 - \gamma_{ver}, y^0 + \delta_{ver}]$, where γ_{ver} and δ_{ver} are small (in the sense that the new drug cannot be much more cost-effective than the pioneer). For horizontal

²¹Note that our innovation process in the vertical dimension differs notably from the vertical innovation analyzed by Bardey et al. (2010), where a vertical innovation is always synonymous with a breakthrough innovation.

incremental processes, the R&D investment leads, in the case of success, to a drug (x^1, y^1) with $y^1 = y^0$ and x^1 is distributed according to a density function $h_{in}(x)$ that takes values in the interval $[x^0 - \gamma_{hor}, x^0 + \delta_{hor}]$, where γ_{hor} and δ_{hor} are positive and small (in the sense that the new drug would be suitable for similar types of patient than the pioneer).

Proposition 3 compares private and social incentives to adopt an incremental innovation process.

Proposition 3 *For an incremental innovation process pursuing a me-too drug, the optimal investment decision for firm 1, I_{in}^1 , and for the social planner, I_{in}^* , coincide: $I_{in}^1 = I_{in}^*$.*

The intuition for this result derives from the fact that, for drugs close to the pioneering drug, the innovative firm is always able to extract all the patient surplus. Consider a vertical innovation process. If it is successful but the realization of \tilde{y}^1 is $y^1 < y^0$ then the new drug is not prescribed and trivially $\Delta CS = 0$. On the other hand, if a vertical innovation has a characteristic $y^1 > y^0$, firm 1 gets some monopoly power because the new drug lies in Region c. The firm will impact all the extra quality of the drug into a higher price and it is able to extract all the patient surplus ($\Delta CS = 0$ in Region c when $\Delta_x = 0$). As patients never benefit from the new drug, social and private incentives to innovate coincide as do their optimal investment levels. The intuition for the horizontal incremental process is similar: when the realization of \tilde{x}^1 is close to x^0 , firm 1 sets the maximum prize that consumers are willing to pay and it extracts all the patient surplus ($\Delta CS = 0$).

We now consider a radical process pursuing a breakthrough drug, which implies revolutionary breakthroughs in disease therapy leading to a drug with a larger cost-effectiveness. If an investment I_{ra} is made then there is a probability $q_{ra}(I_{ra})$, increasing in I_{ra} , of success at getting a new drug (x^1, y^1) .²² If the process is successful, the new drug has characteristics (x^1, y^1) , where y^1 takes values in the interval $[y^0 + \kappa, y^0 + v]$ and x^1 takes values in the interval $[0, 1]$, where κ can be positive or negative, v is positive, $\kappa < v$ and (x^1, y^1) is distributed according to a density function $f_{ra}(x^1, y^1)$.²³

Proposition 4 *For a radical innovation process pursuing a breakthrough drug the optimal investment decision for firm 1, I_{ra}^1 , is smaller than that for the social planner, I_{ra}^* : $I_{ra}^1 < I_{ra}^*$.*

A radical innovation process leads, with a certain probability, to a drug that improves patient surplus. First, patients benefit when the process is successful if the new drug lies

²²Typically, $q_{ra}(I)$ would be (much) lower than $q_{in}(I)$ for any I .

²³The function $f_{ra}(x^1, y^1)$ may not take values for some intervals of characteristics. Figure 4, for instance, illustrates a process that never leads to very extreme drugs on the horizontal axis. In this figure, $\kappa > 0$ but this is not necessary in general.

in Region c and $x^1 \neq x^0$. In this case, the innovative firm cannot appropriate all the patient surplus: at the maximum price at which it can serve all the market (p^{all}), some patients are strictly better-off with the new drug than with the pioneer. Similarly, if the successful process leads to a drug in Region b.ii, then the interior price p^{int} allows patients to benefit. A social planner would also take into account this surplus that the firm cannot extract from patients and it would choose a higher investment level than the firm.

As shown in propositions 3 and 4, the firm fully appropriates all the benefits derived from me-too drugs, whereas patients can appropriate some surplus if a radical innovation process goes successfully. Proposition 5 uses these results to state the main policy implication of our analysis of the innovation incentives.

Proposition 5 *If firm 1 can only adopt one category of innovation process (due, for instance, to financial or capacity constraints) then:*

- (i) *There are cases where firm 1 prefers to adopt an incremental innovation process whereas the social surplus is larger adopting a radical innovation process.*
- (ii) *There is no case where firm 1 prefers to adopt a radical innovation process whereas adopting an incremental innovation process would generate higher welfare.*

Proposition 5 provides some theoretical support to the social concern that pharmaceutical firms devote too many resources to market me-too drugs and too few to launch breakthrough drugs. Our model suggests that this disalignment between private and social incentives is due to the lack of private incentives to pursue radical innovations. While private and social incentives to devote resources to pursue me-too drugs are aligned because the firm is able to appropriate all the benefits through the price, this is not true for first-in-class drugs. In the latter case, the firm sets a price for the new drug which is low enough to serve all (or a good part of) the market and the consumers benefit from it.

8 Conclusion

The proliferation of follow-on drugs is nowadays the subject of some debate. Proponents of follow-on drugs highlight that some of them are therapeutically superior to the pioneer. Moreover, patients and physicians benefit from the access to a larger pool of therapeutic choices. But there are also voices warning that imitative drug development poses a threat, as it could reduce the incentive for firms to develop first-in-class drugs. As evidence of this, an example often cited is that of the protein kinases. These cellular proteins are among the most common targets for drug discovery. However, although there are 518 protein kinases in the human genome, more than half the current drug discovery programs focus

on the handful of kinases for which there is already an existing drug (Fedorov, Müller, and Knapp, 2010).

This paper seeks to contribute to this social debate. We have constructed a simple model where drugs are vertically and horizontally differentiated. After studying the optimal price decision of a firm introducing a new drug, we have analyzed the firm incentives to invest in R&D when a pioneer drug is already on the market. In particular, we have disentangled the quest for breakthrough drugs from the search of follow-on drugs. In our model, both breakthrough and follow-on drugs are always socially valuable.

While private and social incentives to invest in R&D processes coincide for incremental innovation ventures, private incentives are lower than social ones when the process is radical. Moreover, we interestingly find that pharmaceutical firms are too prone to devote resources (if scarce) to pursue incremental innovation processes so as to fully appropriate all the benefits derived from me-too drugs. Thus, these results somehow reproduce the social concern that the main problem regarding the rapid increase of me-too drugs is that they diminish the incentives for innovation in pioneering drugs.

Our conclusions are obtained under some simplifying assumptions; we now discuss some of them. First, we have assumed that the pioneering drug is sold on the market at its marginal price. This hypothesis fits well in markets where the patent for the pioneering drug has already expired and several generic drugs have entered the market. However, there are other markets where the pioneering drug is sold by one firm at a price over its marginal cost. In such cases, the incumbent firm may react to the introduction of a new drug by reducing the price of its drug. Price competition between the two firms will be fiercer the lower the horizontal distance is between the two drugs. This seems to suggest that under price competition the incentives of the innovative firm to differentiate itself in the horizontal axis increase, which may translate into more incentives to adopt radical innovation processes.

Secondly, our innovative firm freely chooses the price of the new drug. However, regulations worldwide to control excessive market power of pharmaceutical firms abound. It would be worth investigating whether price regulations (such as price ceilings or reference prices) would be an effective tool to align the incentives of the firm and the society. Price ceilings in our model, for instance, would undermine private incentives to innovate. Moreover, our analysis seems to suggest that price ceilings would have a larger impact on vertical innovations. Since vertical improvements allow the pharmaceutical firm to charge larger prices, the firm will be more constrained by the regulated price if it launches a new drug that is an improvement over the pioneer in the vertical dimension.

Third, our results have been obtained under the assumption of risk neutrality for all the players involved. However, it is natural to think that pharmaceutical firms exhibit a

certain degree of risk aversion. If we relax the risk neutrality assumption to accommodate more realism, our main result would be reinforced as risk aversion would make firms even more prone to adopt (more safety) incremental innovations.

Finally, in our model there is only one innovative firm so we have not considered competition in research (in Bardey et al., 2010, several laboratories compete in the research sector of the pharmaceutical industry). If firms compete in adopting incremental innovation processes, they will overinvest in comparison with the socially efficient level, leading to more me-too drugs than the socially optimal number. At the same time, the innovative firms will also increase their investment in radical innovation processes under the pressure of competition. And this could lead to investment levels closer to the social optimal one.

The spirit of this work is eminently positive. In a simple setting, we have identified a problem of misalignment between private and social incentives to innovate that results in a bias toward me-too drugs. A more normative analysis, in which different solutions to the problem can be addressed, is left for further research. Such policies could include direct R&D tax incentives, nonprofit tax exemptions for research institutions, public financing of R&D activity, as well as many other instruments that attempt to stimulate various forms of research and innovative activity. Our analysis suggests that the optimal R&D policy should induce firms to pursue more radical innovation ventures.

Finally, although there seems to be a great social awareness of the proliferation of me-too drugs, to date economists have done little theoretical research on this issue. We hope that this study opens the door to further research into that area and that it will also stimulate the ongoing debate over the excessive launching of me-too drugs.

References

- [1] Angell, M., 2004. The truth about the drug companies. New York: Random House.
- [2] Antoñanzas, F., Juárez-Castelló, C., Rodríguez-Ibeas, R., 2011. Innovation, loyalty and generic competition in pharmaceutical markets. *SERIEs* 2, 75-95.
- [3] Avorn, J., 2004. Powerful medicines. New York, Knopf.
- [4] Bardey, D., Bommier, A., Jullien, B., 2010. Retail Price Regulation and Innovation: Reference Pricing in the Pharmaceutical Industry. *Journal of Health Economics* 29 (2), 303-316.
- [5] Cohen, J., Cabanilla, L., Sosnov, J., 2006. Role of follow-on drugs and indications on the WHO Essential Drug List. *Journal of Clinical Pharmacy and Therapeutics* 31, 585-592.

- [6] diMasi, J., Paquette, C., 2004. The Economics of Follow-on Drug Research and Development Trends in Entry Rates and the Timing of Development. *Pharmacoeconomics* 22 (Suppl. 2), 1-14.
- [7] European Commission, 2013. The 2012 EU Industrial R&D Investment Scoreboard. European Commission. Joint Research Centre (JRC). Luxembourg: Publications Office of the European Union.
- [8] Fedorov, O., Müller, S., Knapp, S., 2010. The (un)targeted cancer kinome. *Nature Chemical Biology* 6(3), 166-169.
- [9] García, R., Calantone, R., 2002. A Critical look at technological innovation typology and innovativeness terminology. A literature review. *Journal of Product Innovation Management*, 19(2), 110-132.
- [10] Ganuza J., Llobet G., Dominguez B., 2009. R&D in the Pharmaceutical Industry: A World of Small Innovations. *Management Science* 55 (4), 539-551.
- [11] Garnier, J.P., 2008. Rebuilding the R&D engine in big pharma. *Harvard Business Review* 86(5), 68-70, 72-76, 128.
- [12] Girotra, K., Terwiesch, C., Ulrich, K.T., 2007. Valuing R&D projects in a portfolio: Evidence from the pharmaceutical industry, *Management Science* 5 (9), 1452-1466.
- [13] Goozner, M., 2004. The \$800m pill: The truth behind the cost of new drugs. Berkeley, University of California Press.
- [14] Hernandez A.F., Harrington, R.A., 2008. Comparative effectiveness of angiotensin-converting-enzyme inhibitors: is an ACE always an ace? *Canadian Medical Association Journal* 178 (10), 1316-1319.
- [15] Kapur, N.K., Musunuru, K., 2008. Clinical efficacy and safety of statins in managing cardiovascular risk. *Journal of Vascular Health and Risk Management* 4(2), 341-353.
- [16] Mestre-Ferrandiz, J., Mordoh, A., Sussex, J., 2012. The many faces of innovation. Office for Health Economics. Available at: <http://www.ohe.org/publications/article/the-many-faces-of-innovation-119.cfm>
- [17] Miller, H.I., 2014. Critics of 'Me-Too Drugs' Need to Take a Chill Pill. *Wall Street Journal*, January 1st.
- [18] Pharmaceutical Research and Manufacturers of America, 2013. 2013 Biopharmaceutical Research Industry Profile. Washington, DC, PhRMA, July.

- [19] Singh, J., Fleming L., 2010. Lone Inventors as Sources of Breakthroughs: Myth or Reality? *Management Science* 56 (1), 41-56.
- [20] Wertheimer, A.I., Santella T.M., 2009. Pharmaceutical Evolution: The Advantages of Incremental Innovation in Drug Development. Competitive Enterprise Institute. Issue Analysis, no.2, April. Washington, DC.
- [21] Yin, N., 2012. Pharmaceuticals, Incremental Innovation and Market Exclusivity. Mimeo, Toulouse School of Economics.

9 Appendix

Proof of Proposition 1. We distinguish three zones of parameters: $c^1 \geq p^{\max}$, $c^1 \in (p^{\text{all}}, p^{\max})$, and $c^1 \leq p^{\text{all}}$. We note that $c^1 \geq p^{\max}$ if and only if $\Delta_c \geq b(l\Delta_x + \Delta_h)$, i.e., $\Delta_y \leq -l\Delta_x$. Similarly, $c^1 \leq p^{\text{all}}$ if and only if $\Delta_c \leq b(\Delta_h - l\Delta_x)$, i.e., $\Delta_y \geq l\Delta_x$. We analyze the optimal firm's pricing in the three zones.

Zone $\Delta_y \leq -l\Delta_x$. Given that $c^1 \geq p^{\max}$, there is no price below marginal cost at which the physician prescribes the new drug to some patients. Therefore, the best strategy for firm 1 is not to sell its drug.

Zone $\Delta_y \in (-l\Delta_x, l\Delta_x)$. In this zone, $c^1 > p^{\text{all}}$ and firm 1 cannot select a price at which it makes a profit by selling to all the patients. If $x^1 \geq x^0$, the firm chooses the price p^1 that maximizes its profits, taking into account that its drug will be prescribed to those patients with characteristic $x \in [\tilde{x}, 1)$. Hence, $p^1 \in [c^1, p^{\max}]$ maximizes

$$\Pi^1(p^1) = (p^1 - c^1)(1 - \tilde{x}) = (p^1 - c^1) \left[1 - x^0 - \frac{(l\Delta_x - \Delta_h)}{2l} - \frac{(p^1 - c^0)}{2bl} \right].$$

We write the expression for the profits as

$$\Pi^1(p^1) = (p^1 - c^1) \left[M - \frac{(l\Delta_x - \Delta_h)}{2l} - \frac{(p^1 - c^0)}{2bl} \right],$$

which is also valid for the situation where $x^1 < x^0$.²⁴

The first-order and second-order conditions are

$$\Pi_{p^1}^1(p^1) = M - \frac{(l\Delta_x - \Delta_h)}{2l} - \frac{(2p^1 - (c^0 + c^1))}{2bl},$$

²⁴If $x^1 < x^0$, then

$$\tilde{x} = x^0 - \frac{(l\Delta_x - \Delta_h)}{2l} - \frac{(p^1 - c^0)}{2bl} = M - \frac{(l\Delta_x - \Delta_h)}{2l} - \frac{(p^1 - c^0)}{2bl}.$$

$$\Pi_{p^1 p^1}^1(p^1) = \frac{-1}{bl} < 0.$$

The interior candidate to solution p^1 satisfies $\Pi_{p^1}^1(p^1) = 0$ and is given by

$$p^{int} = \frac{1}{2}(c^0 + c^1) + blM - \frac{1}{2}b(l\Delta_x - \Delta_h).$$

The concavity of the function $\Pi^1(p^1, \Delta_x, \Delta_h)$ implies that the candidate p^{int} is the optimum if and only if it lies in the interval $[c^1, p^{\max}]$. First, $p^{int} > c^1$ if and only if

$$\frac{1}{2}\Delta_c + blM - \frac{1}{2}b(l\Delta_x - \Delta_h) > 0,$$

that is, $\Delta_y > l\Delta_x - 2lM$. Given that $M \geq \Delta_x$, the previous inequality is implied by $\Delta_y > -l\Delta_x$, which is satisfied in this zone. Second, $p^{int} < p^{\max} = c^0 + b(l\Delta_x + \Delta_h)$ if and only if

$$\frac{1}{2}\Delta_c + blM - \frac{1}{2}b(l\Delta_x - \Delta_h) < b(l\Delta_x + \Delta_h),$$

from which it easily follows that $p^{int} < p^{\max}$ if and only if $M < \frac{1}{2l}(3l\Delta_x + \Delta_y)$. Therefore, in this zone, $p^{1*} = p^{int}$ if $M < \frac{1}{2l}(3l\Delta_x + \Delta_y)$ and $p^{1*} = p^{\max}$ otherwise.

Zone $\Delta_y \geq l\Delta_x$. In this zone, firm 1 can set the price p^{all} (or $p^{all} - \varepsilon$, for ε infinitesimal) that would allow it to attract all patients in $[0, 1]$. It can also choose any price in the interval $p^1 \in (p^{all}, p^{\max}]$, in which case the new drug will be prescribed to a subset of the patients. In this interval, the interior candidate to solution is p^{int} , as in the previous zone. We check the conditions under which $p^{int} \in (p^{all}, p^{\max}]$.

First, $p^{int} > p^{all}$ if and only if $\frac{1}{2}(c^0 + c^1) + blM - \frac{1}{2}b(l\Delta_x - \Delta_h) > c^0 + b(\Delta_h - l\Delta_x)$, that is,

$$M > \frac{1}{2l}(\Delta_y - l\Delta_x).$$

Note that at $p^1 = p^{all}$, firm 1 serves a market of size M . However, this cannot be the best pricing strategy for the firm because it can serve the whole market and obtain larger profits by marginally decreasing the price. Therefore, if $M \leq \frac{1}{2l}(\Delta_y - l\Delta_x)$, setting the price p^{all} and serving the whole market is certainly the optimal decision.

Second, as we show in the analysis of Zone $\Delta_y \in (-l\Delta_x, l\Delta_x)$, $p^{int} \leq p^{\max}$ if and only if $M \leq \frac{1}{2l}(3l\Delta_x + \Delta_y)$.

Thus, for $M \geq \frac{1}{2l}(3l\Delta_x + \Delta_y)$, the candidates for solution are p^{\max} and p^{all} ; for $M \in (\frac{1}{2l}(\Delta_y - l\Delta_x), \frac{1}{2l}(3l\Delta_x + \Delta_y))$ the candidates for solution are p^{int} and p^{all} ; and for $M \leq \frac{1}{2l}(\Delta_y - l\Delta_x)$, the optimal price is p^{all} .

We now analyze the conditions for $\Pi^1(p^{\max}) \geq \Pi^1(p^{all})$, that is, $b(M - \Delta_x)(\Delta_y + l\Delta_x) \geq b(\Delta_y - l\Delta_x)$, or

$$\Delta_y \leq \frac{(1 + M - \Delta_x)}{(1 - M + \Delta_x)}l\Delta_x = lg(\Delta_x).$$

We note that the border $\Delta_y = lg(\Delta_x)$ always lies in the zone $\Delta_y \geq l\Delta_x$, because $b\Delta_y - blg(\Delta_x) \leq b(\Delta_y - l\Delta_x)$ if and only if $g(\Delta_x) \geq \Delta_x$, which is equivalent to $\frac{(1+M-\Delta_x)}{(1-M+\Delta_x)} \geq 1$, or $M \geq \Delta_x$, which holds.

Similarly, $\Pi^1(p^{int}) \geq \Pi^1(p^{all})$ if and only if $\frac{b}{2l} \left(lM + \frac{1}{2}(\Delta_y - l\Delta_x) \right)^2 \geq b(\Delta_y - l\Delta_x)$ that, denoting $t \equiv (\Delta_y - l\Delta_x)$, can be written as $\left(lM + \frac{1}{2}t \right)^2 - 2lt \geq 0$, that is,

$$f(t) \equiv \left(t - 2l \left(2 - M + 2\sqrt{1 - M} \right) \right) \left(t - 2l \left(2 - M - 2\sqrt{1 - M} \right) \right) \geq 0.$$

We are interested in the cases where $M > \frac{1}{2l}(\Delta_y - l\Delta_x)$ (because p^{all} is certainly optimal otherwise), that is, $t < 2lM$. The function $f(t)$ satisfies $f(0) > 0$. Moreover, $f(2lM) < 0$. Therefore, $f(t) \geq 0$ if and only if t is lower than the first root, that is, $\Delta_y - l\Delta_x \leq 2l \left(2 - M - 2\sqrt{1 - M} \right)$, or

$$\Delta_y \leq l\Delta_x + 2l \left(2 - M - 2\sqrt{1 - M} \right) = lg(\Delta_x).$$

We note that the change in the definition of the function $g(\Delta_x)$ happens at the point where $M = \frac{1}{2l}(\Delta_y + 3l\Delta_x)$ (which separates the regions where either p^{int} or p^{max} are candidates). Then, the value Δ_x where the change happens is the solution of the following system of equations (in Δ_x and Δ_h):

$$\begin{aligned} M &= \frac{1}{2l}(\Delta_y + 3l\Delta_x) \\ \Delta_y &= l\Delta_x + 2l \left(2 - M - 2\sqrt{1 - M} \right) \end{aligned}$$

that is,

$$\Delta_x = \sqrt{1 - M} - (1 - M),$$

which is a positive value. The function $g(\Delta_x)$, as it is defined in the main text just before Proposition 1, is continuous because it is continuous at the point $\sqrt{1 - M} - (1 - M)$, and it is also continuously differentiable.

Once we analyze the optimal price in each of the three zones, it easily follows that the solution is continuous, in the sense that if the optimal price is p^{all} (resp. p^{max}) in the second zone, then it is also optimal if we decrease Δ_c and enter the third zone. Therefore, the optimal firm 1's pricing policy is the one described in the proposition.

Finally, it is easy to check that the profits are continuous: (a) over the line $\Delta_y = 2lM - 3l\Delta_x$, we have $\Pi^1(p^{max}) = \Pi^1(p^{int})$. (b) If the condition $\Delta_y = lg(\Delta_x)$ holds we have two cases: (b.1) For $g(\Delta_x) \equiv \frac{(1+M-\Delta_x)}{(1-M+\Delta_x)}\Delta_x$, $\Pi^1(p^{all}) = \Pi^1(p^{max})$ and (b.2) for $g(\Delta_x) \equiv \Delta_x + 2 \left(2 - M - 2\sqrt{1 - M} \right)$, then $\Pi^1(p^{all}) = \Pi^1(p^{int})$. ■

Proof of Corollary 1. We denote $\Pi^1(\Delta_x, \Delta_y)$ the firm 1's profits as a function of Δ_x and Δ_y .

In Region b.i, where $p^{1*} = p^{max}$:

$$\Pi_{\Delta_y}^1(\Delta_x, \Delta_y) = b(M - \Delta_x) > 0; \quad \Pi_{\Delta_y \Delta_y}^1(\Delta_x, \Delta_y) = 0.$$

$\Pi_{\Delta_x}^1(\Delta_x, \Delta_y) = b(lM - (2l\Delta_x + \Delta_y))$, which is positive for $\Delta_y = -l\Delta_x$ and negative for $\Delta_y = 2lM - 3l\Delta_x$; $\Pi_{\Delta_x \Delta_x}^1(\Delta_x, \Delta_y) = -2bl < 0$.

In Region b.ii, where $p^{1*} = p^{int}$:

$$\Pi_{\Delta_y}^1(\Delta_x, \Delta_y) = \frac{b}{4l}(2lM + \Delta_y - l\Delta_x) > 0; \quad \Pi_{\Delta_y \Delta_y}^1(\Delta_x, \Delta_y) = \frac{bl}{4} > 0.$$

$$\Pi_{\Delta_x}^1(\Delta_x, \Delta_y) = -\frac{b}{4}(2lM + \Delta_y - l\Delta_x) < 0; \quad \Pi_{\Delta_x \Delta_x}^1(\Delta_x, \Delta_y) = \frac{bl}{4} > 0.$$

In Region c, where $p^{1*} = p^{all}$:

$$\Pi_{\Delta_y}^1(\Delta_x, \Delta_y) = b > 0; \quad \Pi_{\Delta_y \Delta_y}^1(\Delta_x, \Delta_y) = 0.$$

$$\Pi_{\Delta_x}^1(\Delta_x, \Delta_y) = -lb < 0; \quad \Pi_{\Delta_x \Delta_x}^1(\Delta_x, \Delta_y) = 0. \blacksquare$$

Proof of Proposition 2. Proofs for Region a and Region b.i are immediate.

In Region b.ii the variation in patient surplus, that is, (in case $x^1 \geq x^0$)²⁵ the difference for patients in $[\tilde{x}, 1]$ from being treated with drug (h^0, x^0) at price c^0 and being treated with drug (h^1, x^1) at price p^{int} , is the sum of a triangle for $[\tilde{x}, x^1]$ and a rhomboid for $[x^1, 1]$.

a) For $[\tilde{x}, x^1]$, $\Delta CS = \frac{1}{2}(x^1 - \tilde{x}(p^{int}))(bh^1 - p^{int} - b(h^0 - l(x^1 - x^0)) + c^0)$. Substituting $\tilde{x}(p^{int})$ and p^{int} , we have

$$\Delta CS = \frac{b}{16l}(\Delta_y + 3l\Delta_x - 2lM).$$

b) For $[x^1, 1]$, $\Delta CS = (1 - x^1)(bh^1 - p^{int} - b(h^0 - l(x^1 - x^0)) + c^0)$. Substituting p^{int} and using $(1 - x^1) = (M - \Delta_x)$, we have

$$\Delta CS = \frac{b}{2}(M - \Delta_x)(\Delta_y + 3l\Delta_x - 2lM).$$

Hence, adding both intervals, we obtain

$$\Delta CS(p^{int}) = \frac{b}{16l}(\Delta_y + 3l\Delta_x - 2lM)(\Delta_y - 5l\Delta_x + 6lM).$$

In Region c the variation in patient surplus is similar to the one previously analyzed, but taking into account that for p^{all} (considering again that $x^1 > x^0$) patients in $[0, x^0]$ have the same surplus than without the new drug. For the patients in $[x^0, x^1]$,

$$\Delta CS = \frac{1}{2}(x^1 - x^0)(bh^1 - p^{all} - b(h^0 - l(x^1 - x^0)) + c^0).$$

Substituting p^{all} we have

$$\Delta CS = bl\Delta_x^2.$$

²⁵The case $x^1 < x^0$ is similar.

For $[x^1, 1]$, $\Delta CS = (1 - x^1) (bh^1 - p^{all} - b(h^0 - l(x^1 - x^0)) + c^0)$. Substituting p^{all} and using $(1 - x^1) = (M - \Delta_x)$, we obtain

$$\Delta CS = 2bl\Delta_x (M - \Delta_x).$$

Adding the two parts, we have

$$\Delta CS(p^{all}) = bl\Delta_x (2M - \Delta_x).$$

■

Proof of Corollary 2. We denote $\Delta CS(\Delta_x, \Delta_y)$ the variation in consumer surplus as a function of Δ_x and Δ_y .

In Region a and Region b.i all the derivatives are zero.

In Region b.ii:

$$\Delta CS_{\Delta_y}(\Delta_x, \Delta_y) = \frac{b}{8l} (\Delta_y - l\Delta_x + 2lM) > \frac{b}{8l} (-2l\Delta_x + 2lM) > 0.$$

$$\Delta CS_{\Delta_y\Delta_y}(\Delta_x, \Delta_y) = \frac{b}{8l} > 0.$$

$\Delta CS_{\Delta_x}(\Delta_x, \Delta_y) = \frac{1}{8} (-\Delta_y - 15l\Delta_x + 14lM)$. We note that, at $M = \frac{1}{2l} (\Delta_y + 3l\Delta_x)$, $\Delta CS_{\Delta_x}(\Delta_x, \Delta_y) = \frac{3}{4} (\Delta_y + l\Delta_x) > 0$ because $\Delta_y > -l\Delta_x$. Moreover, $\Delta CS_{\Delta_x}(\Delta_x, \Delta_y) < \frac{1}{8} (-14l\Delta_x + 14lM)$. Hence, $\Delta CS_{\Delta_x}(\Delta_x = M, \Delta_y) < 0$.

$$\Delta CS_{\Delta_x\Delta_x}(\Delta_x, \Delta_y) = -\frac{15}{8}bl < 0$$

In Region c:

$$\Delta CS_{\Delta_y}(\Delta_x, \Delta_y) = 0; \Delta CS_{\Delta_y\Delta_y}(\Delta_x, \Delta_y) = 0.$$

$$\Delta CS_{\Delta_x}(\Delta_x, \Delta_y) = 2bl(M - \Delta_x) > 0.$$

$$\Delta CS_{\Delta_x\Delta_x}(\Delta_x, \Delta_y) = -2bl < 0.$$

Finally, the behavior of ΔCS at the borders between regions follows from the behavior of the optimal price p^{1*} that we have discussed following Proposition 1. ■

Proof of Proposition 3. We denote $\Pi^1(x, y)$ and $\Delta CS(x, y)$ the firm 1's profits and the increase in consumer surplus when the new drug has characteristics (x, y) . For all I invested in a vertical incremental innovation process, expected private profits $\Pi_{in}(I)$ and social welfare $W_{in}(I)$ are

$$\Pi_{in}(I) = q_{in}(I) \int_{[y^0 - \gamma_{ver}, y^0 + \delta_{ver}]} \Pi^1(x = x^0, y) f_{in}(y) dy,$$

$$W_{in}(I) = q_{in}(I) \int_{[y^0 - \gamma_{ver}, y^0 + \delta_{ver}]} (\Pi^1(x = x^0, y) + \Delta CS(x = x^0, y)) f_{in}(y) dy.$$

In case of a successful project, the new drug $(x = x^0, y)$ lies either in Region a (for $y \leq y^0$) or Region c with $x = x^0$ (for $y > y^0$). In the first case (see Proposition 2), $\Delta CS(x = x^0, y) = \Delta CS(c^1) = 0$. In the second case, $\Delta CS(x = x^0, y) = \Delta CS(p^{all}) =$

$b\Delta_x(2M - \Delta_x) = 0$ because $\Delta_x = 0$. Therefore, $\Pi_{in}(I) = W_{in}(I)$ for all I and the optimal investment levels for the two functions coincide.

The argument for a horizontal incremental innovation process is similar because the new drug would lie in Region b.i where $\Delta CS(x, y = y^0) = \Delta CS(p^{\max}) = 0$. ■

Proof of Proposition 4. For each investment I in a radical innovation process,

$$\Pi_{ra}(I) = q_{ra}(I) \int_{[0,1]} \int_{[y^0+\kappa, y^0+v]} \Pi^1(x^1, y^1) f_{ra}(x^1, y^1) dy^1 dx^1$$

and

$$W_{ra}(I) = q_{ra}(I) \int_{[0,1]} \int_{[y^0+\kappa, y^0+v]} (\Pi^1(x^1, y^1) + \Delta CS(x^1, y^1)) f_{ra}(x^1, y^1) dy^1 dx^1.$$

There is some positive probability that the new drug (x^1, y^1) lies in Region c with $x^1 \neq x^0$. Depending on the level of y^1 , there can also be a positive probability that the new drug lies in Region b.ii). In both regions, $\Delta CS(x, y^1) > 0$. Given that $\Delta CS(x, y) \geq 0$ for every new drug, we have $\frac{d\Pi_{ra}(I)}{dI} < \frac{dW_{ra}(I)}{dI}$ for every I , which implies that $I_{ra}^1 < I_{ra}^*$. ■

Proof of Proposition 5. Denoting by I_Z^1 and I_Z^* for $Z = in, ra$ the optimal level of investment in a process of type Z for firm 1 and the social planner, respectively, then

$$\begin{aligned} \Pi_{in}(I_{in}^1) &= W_{in}(I_{in}^*) \\ \Pi_{ra}(I_{ra}^1) &< W_{ra}(I_{ra}^*). \end{aligned}$$

It may happen that

$$W_{ra}(I_{ra}^*) > W_{in}(I_{in}^*) = \Pi_{in}(I_{in}^1) > \Pi_{ra}(I_{ra}^1)$$

but it can never be the case that

$$W_{ra}(I_{ra}^*) < W_{in}(I_{in}^*) = \Pi_{in}(I_{in}^1) < \Pi_{ra}(I_{ra}^1),$$

which implies the result. ■