Controlling Excessive Off-Label Medicare Drug Costs Through the False Claims Act

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Controlling Excessive Off-Label Medicare Drug Costs Through the False Claims Act

David Kwok†

Abstract

High prescription drug prices are driving ever-increasing United States healthcare costs, and the federal government is following this alarming trend with ninety-five billion dollars in expenditures for prescription drugs under Medicare Part D. Even accepting arguments that high drug prices are necessary to encourage the development of safe and effective drugs, Medicare Part D is flawed in that it will pay top dollar for ineffective drugs. Because Part D lacks adequate oversight for off-label drug usage, pharmaceutical companies obtain windfall profits for drugs that have not been proven effective for off-label conditions. Permitting companies to reap such profits without incurring the costs of demonstrating efficacy creates a distorted marketplace that leads to excessive Medicare drug expenditures. In addition to the financial burden to taxpayers and the risks to Medicare patients’ health, the flaws in Medicare Part D also increase the risk that non-Medicare patients will be prescribed ineffective and expensive drugs. This article proposes a theoretical reimbursement scheme that encourages fairness and restrains excessive off-label drug reimbursement by tying reimbursement rates to competitive products. Fully correcting this systemic problem will require substantial statutory, regulatory, and institutional reforms that are not immediately likely. In the interim, courts and regulators should embrace the civil False Claims Act to begin to correct the incentive problems created under the present off-label reimbursement structure, thereby immediately curbing excessive Medicare spending on prescription drugs.

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I. Introduction

High prescription-drug prices have been generating alarm in the media and in Congress, and the federal government is no stranger to prescription-drug purchases. Medicare spending on outpatient prescription drugs through Medicare Part D was ninety-five billion dollars in 2016, amounting to thirteen percent of overall Medicare benefit payments. The Centers for Medicare & Medicaid Services (“CMS”) have acknowledged that total Part D costs per capita have been rising at a troubling eleven-percent rate. Pharmaceutical companies (“manufacturers”) typically justify high drug prices by citing the costs of research and development. The high prices offset the substantial costs incurred in conducting scientific studies to demonstrate that drugs are safe and effective in treating particular conditions. High expenditures on drugs are arguably good for society and not excessive if patients are obtaining sufficient benefit from those drugs.


The problem, however, is that Medicare Part D can pay manufacturers these high drug prices without a full demonstration that the drugs are effective. By law, Medicare reimbursements are limited to drugs prescribed for medically accepted indications—conditions for which there is scientific evidence that a drug will be safe and effective. In practice, however, there is no systemic mechanism to ensure that drugs are actually prescribed for such an indication.7 Drug prescriptions are not required to include the indication for which the drugs are prescribed under Part D.8 Furthermore, a physician may legally prescribe a drug for various “off-label” indications—conditions not formally approved by the Food & Drug Administration (“FDA”).9

This system is unfair, and it creates perverse incentives for manufacturers. A manufacturer who has not incurred the costs of completing scientific studies demonstrating its drug’s effectiveness can earn greater revenue than a competing manufacturer who has completed such studies for its alternative drug. This distorted system drives excessive Part D spending, because the government pays more for the unproven drug in comparison to the fully tested drug.10 In a rational system, the government would not pay higher prices for a drug with less evidence to indicate that it is safe and effective. These high prices are likely to drive aggressive manufacturer’s marketing efforts to physicians, and patients may be exposed to higher probabilities of expensive and unproven off-label drug usage.

This article proposes a theoretical reimbursement framework that eliminates this distortion and unfairness by capping off-label reimbursements at a competitive level. A drug that is prescribed for its on-label, FDA-approved condition will continue to receive existing full reimbursement. If a drug is prescribed for an off-label condition, however, its reimbursement level will be tied to the competitive market for the off-label condition. A manufacturer that has not completed scientific studies regarding off-label drug usage will not receive a higher reimbursement than a competing manufacturer that has completed those studies. This


8. See OIG PART D REIMBURSEMENT REPORT, supra note 6.


framework is superior to existing proposals in that it allows room for the development of optimal levels of off-label drug usage. Medicare beneficiaries will still have access to drugs for off-label purposes and manufacturers will have the proper incentives for research.

While attractive in theory, fully implementing this theoretical reimbursement framework is challenging in the short term. Like other proposals, it requires the integration of prescription and diagnosis information. There are difficult structural and statutory barriers to such integration; state law generally governs prescription information11 and the federal government has been hesitant to interfere.12 Additionally, the present standard for tracking patient diagnoses does not correspond with Part D reimbursement rules.13

In the interim, this article proposes civil False Claims Act (“FCA”) liability as a claw-back mechanism to control Part D expenditures by limiting the present system’s distortions and inequity. While manufacturers may temporarily enjoy excessive profits through ever-growing levels of off-label drug reimbursement, civil liability under the FCA will allow the government to reclaim, or “claw back,” those unfair profits and help fund the need for better links between diagnosis and prescription. Unlike other forms of immediately available civil litigation, such as tort liability, the FCA incorporates a whistleblower cause of action.14 Whistleblowers are critical in supplying the core missing information linking prescriptions to diagnoses. Without such whistleblowers, litigation lacks the funding and support to compile the missing information.

FCA liability has already been applied to subset of off-label reimbursement scenarios: off-label promotion cases against manufacturers.15 Under the existing theory, manufacturers are liable for excessive Medicare expenditures because their promotional efforts induce physicians to prescribe off-label drugs that result in improper Medicare reimbursement.16 This article’s proposed FCA solution expands on this theory by including any manufacturer behavior that is a cause-in-fact of excessive Medicare expenditures. Furthermore, FCA liability should be

11. See OIG PART D REIMBURSEMENT REPORT, supra note 6.
12. See James M. Beck & Elizabeth D. Azari, FDA, Off-Label Use, And Informed Consent: Debunking Myths And Misconceptions, 53 FOOD DRUG LAW J. 71, 76 (stating that the FDCA was “not intended as a medical practices act and [did] not interfere with the practice of the healing art.”); See also Weber et al., supra note 7 (quoting Jonathan Blum, director of Medicare, that agency philosophy “really has been to defer to physicians.”).
13. See OIG PART D REIMBURSEMENT REPORT, supra note 6
16. Id.
calibrated to the competitive market for the patient’s diagnosis; manufacturers should be liable for any windfall profits that result when their off-label reimbursement exceeds the competitive market rate for drugs with scientifically proven efficacy.

The FCA can be a surprisingly effective claw-back mechanism as a short-term solution. Applying a claw-back mechanism to all manufacturers who benefit from off-label reimbursements would provide properly aligned incentives and eliminate inequity from the present system.

Part II describes the regulatory environment and structure that leads Medicare Part D to be susceptible to excessive drug costs through off-label drug reimbursement. Part III highlights that even if we give manufacturers the benefit of the doubt, their legal actions will still lead to excessive drug costs and unfairness because of the existing reimbursement system. Part IV proposes a theoretically superior reimbursement system that acknowledges the potential societal value of off-label drug reimbursement. Part V discusses how the FCA can be used as an interim claw-back solution to reduce excessive drug costs. Part VI addresses some concerns about this expanded use of the FCA, and Part VII is the conclusion.

II. Background on Off-label Drugs and Medicare Part D

The United States healthcare system attracts extensive criticism for its high costs and comparatively inferior results. Medicare provided nearly six hundred billion dollars in benefits in 2014. Critics often point to expensive pharmaceuticals as contributing to high U.S. healthcare costs. Nonetheless, manufacturers generally defend high pharmaceutical prices by citing the need for expensive research to develop safe and effective

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18. See also KAISER FAMILY FOUND., supra note 2.

Medicare reimburses for prescription drugs through numerous mechanisms; this article focuses upon a particular flaw in Medicare Part D reimbursement.

Medicare Part D covers outpatient prescription drugs, also known as self-administered prescription drugs. Part D expenditures were ninety-five billion dollars in 2016, amounting to roughly thirteen percent of overall Medicare benefit payments. CMS has acknowledged that total Part D costs per capita have been rising at a troubling eleven-percent rate. Part of the core problem with Part D spending is that, while reimbursements are legally limited to drugs provided for medically accepted indications, there is no systemic mechanism to ensure that drugs are actually prescribed for such an indication. The written prescription contains no direct link between the drug and the indication for which it was prescribed under Part D. A physician will diagnose a patient with a certain condition and then prescribe a drug to treat that condition, but the prescription itself simply specifies the drug and dosage information. The patient then brings the prescription to a pharmacist who fills the prescription and files paperwork for reimbursement to Medicare.

CMS manages Part D, but much of the execution is delegated to Medicare Part D sponsors: private insurance companies. Each plan sponsor has substantial autonomy in the coverage of Medicare Part D patients. By delegating such authority to sponsors, Medicare attempts to benefit from the private competitive market. These sponsors can compete to provide superior drug coverage to patients while also competing to hold drug costs down. The sponsors decide upon formularies, which are the lists of drugs covered by the sponsor. The sponsors may set different levels of cost-sharing with patients. For example, sponsors decide on

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21. See CONGRESSIONAL BUDGET OFFICE, supra note 2; see also KAISER FAMILY FOUND., supra note 2.

22. See Centers for Medicaid and Medicare Services, supra note 3.

23. OIG PART D REIMBURSEMENT REPORT, supra note 6, at 1.

24. See id. at 5.


27. See MEDPAC 2014, supra note 25, at 362-363. (noting that the sponsors have had some success in reducing drug prices when generic competition was available, but face challenges when drugs are unique treatments).
Sponsors receive a variety of payments from Medicare.\textsuperscript{28} The core payment is the direct subsidy, a monthly payment to sponsors, adjusted for individual enrollee risk.\textsuperscript{29} Medicare also pays for eighty percent of drug spending that exceeds the out-of-pocket threshold for any patient;\textsuperscript{30} this payment is known as reinsurance. The third major payment is the low-income subsidy ("LIS") through which Medicare covers enrollee costs for those who would have trouble paying for coverage.\textsuperscript{31}

To be clear, physicians are free to exercise their own judgment in prescribing drugs for various conditions; the FDA does not want to be seen as interfering with a physician’s practice of medicine.\textsuperscript{32} Similarly, CMS’s focus is not upon physician decision-making, but on managing drug reimbursement for Medicare, and the agency’s decision is heavily entwined with the FDA’s processes. I begin with a discussion of the FDA’s drug approval process.

A. FDA Approval of Drugs

Before pharmaceuticals enter into interstate commerce, the Federal Food, Drug, and Cosmetic Act ("FDCA") requires FDA approval for specific uses, including indication, population, dosage, and duration.\textsuperscript{33} To obtain FDA approval, manufacturers must demonstrate the safety and efficacy of a new drug for each intended use or indication.\textsuperscript{34} Pharmaceutical manufacturers pay for scientific studies to establish the safety and efficacy of the drugs for particular indications. In deciding whether to approve a new drug, the FDA compares the drug’s benefits against its risks by asking whether the drug offers sufficient benefits to justify the risk of side effects.\textsuperscript{35}

Once approved, the FDCA requires manufacturers to label the drug in a fashion consistent with the FDA-approved usage. The FDA similarly places limits on manufacturers’ promotional activities regarding the drugs. Physicians, however, do not fall within the FDA’s jurisdiction. Physicians

\textsuperscript{28} See id. at 375.

\textsuperscript{29} See id; see also id. at 362 (explaining how there are also risk corridors that address market-based risks as opposed to individual patient risk).

\textsuperscript{30} See MEDPAC 2016, supra note 26, at 174.

\textsuperscript{31} See id. at 157.


\textsuperscript{35} Bruce Patsner, Marketing Approval Versus Cost of New Medical Technologies in the Era of Comparative Effectiveness: CMS, not FDA, Will Be the Primary Player, 3 J. HEALTH & LIFE SCI. L. 38, 55 (2010).
are free to prescribe an approved drug in a manner that differs from the approved usage.\textsuperscript{36} Prescribing a drug for an alternative usage is known as off-label prescription.

Off-label prescription is common, potentially comprising over twenty percent of prescriptions.\textsuperscript{37} Off-label usage is particularly frequent in psychiatry, oncology, and pediatrics.\textsuperscript{38} The U.S. General Accounting Office\textsuperscript{39} (“GAO”) found that one third of cancer drugs were off-label and that more than half of all cancer patients received at least one drug for an off-label indication.\textsuperscript{40} Patients with rare diseases—also called orphan diseases—are also often dependent on off-label uses for their treatment because the number of patients with each orphan disease is often too low to justify the tremendous expense associated with seeking FDA approval for those indications.\textsuperscript{41} Approximately twenty-one percent of all drugs prescribed to treat orphan diseases are off-label.\textsuperscript{42}

B. Government reimbursement for off-label drug usage

The fact that the FDA has not approved a drug for an off-label indication is not determinative as to reimbursement eligibility. CMS decides whether or not to reimburse for a drug and at what price to reimburse for a drug.\textsuperscript{43} Historically, CMS would generally reimburse for a drug that the FDA had approved, but the two agencies’ decisions have shown some divergence more recently.\textsuperscript{44} Aside from differences in standards and procedures, CMS’s mission explicitly incorporates financial security; cost-effectiveness is a consideration beyond the benefits and risk of the drug.\textsuperscript{45} CMS will consider off-label uses and clinical data that are not part of the FDA approval process, which leads to CMS approval of a drug for an indication that has not received FDA approval.\textsuperscript{46} Conversely, if the

\begin{itemize}
  \item \textsuperscript{36} See 21 U.S.C. § 396 (2012).
  \item \textsuperscript{37} See David C. Radley et al., \textit{Off-Label Prescribing Among Office-Based Physicians}, 166 ARCHIVES INTERNAL MED. 1021, 1023 (2006).
  \item \textsuperscript{39} Now known as the Government Accountability Office.
  \item \textsuperscript{41} Bryan A. Liang & Tim Mackey, \textit{Reforming Off-Label Promotion to Enhance Orphan Disease Treatment}, 327 SCI. 273, 273 (2010).
  \item \textsuperscript{42} \textit{id.}
  \item \textsuperscript{43} See Patsner, supra note 35, at 41.
  \item \textsuperscript{44} See \textit{id.} at 43.
  \item \textsuperscript{45} See \textit{id.} at 55.
  \item \textsuperscript{46} See \textit{id.} at 56 (citing Jeffrey A Kelman, M.D., Chief Medical Officer, Center for Beneficiary Choices, CMS).
\end{itemize}
costs of an FDA-approved drug are too high and the drug does not provide a superior benefit-risk calculus compared to existing competitors, CMS might decline to cover the FDA-approved drug.47

Patient-administered drugs fall under Medicare Part D, in contrast to professionally administered drugs under Part A and Part B.48 Also known as outpatient drugs, these patient-administered drugs are first prescribed by a physician. The patient then typically brings the prescription to a pharmacy that fills the prescription and bills the insurer, here Medicare Part D. While the patient’s medical records with the physician contain the patient’s diagnosis, the prescription that the pharmacy sees does not. Thus, under Part D, reimbursement is linked to the price of the drug and not to the patient’s diagnosis.49 The government knows the price of the drug to be reimbursed under Part D, but it does not explicitly know why the patient should be taking that drug.50

As a formal matter, for outpatient drug claims to qualify for Medicare Part D reimbursement, the drugs must be provided for medically accepted indications. Medically accepted indications include both uses approved by FDA and uses supported by one or more of three publications, known as compendia, specified in section 1927(g)(1)(B)(i) of the Social Security Act.51 Medically accepted indications may also be established through Local Coverage Decisions, by which CMS contracts with private organizations to make regionally limited decisions. Finally, CMS also establishes medically accepted indications through annually published National Coverage Decisions.52

To summarize, some off-label use of drugs may be reimbursable under Medicare Part D, but since the drugs are prescribed and reimbursed without a direct link to the diagnosis, CMS does not immediately know whether the drug prescription is legally reimbursable.

47. See id. at 57.

48. See Which Part of Medicare Will Cover My Prescription Drugs (A, B, or D)?, MEDICARE INTERACTIVE, https://www.medicareinteractive.org/get-answers/medicare-covered-services/prescription-drugs/which-part-of-medicare-will-cover-my-prescription-drugs-a-b-or-d (last visited Nov. 20, 2016); While this article focuses on Medicare Part D, it should be noted that Medicaid rules for off-label drug reimbursement are similar, and much of the article’s reasoning can thus be applied to Medicaid reimbursement. See, e.g., United States ex rel. Franklin v. Parke-Davis, No. Civ. A. 96-11651PBS, 2003 WL 22048255 at *3 (D. Mass. Aug. 22, 2003).

49. See OIG PART D REIMBURSEMENT REPORT, supra note 6, at 5.

50. See id. at 5-6.

51. Id. at 1.

C. Criticism of manufacturers

Criticism relating to off-label drug usage has generally focused upon the role of manufacturers. Manufacturers do not prescribe drugs, but they certainly develop and promote drugs. In an oft-cited example, the manufacturer of Neurontin, an FDA-approved epilepsy drug, pursued an off-label marketing strategy that brought in over two billion dollars a year with roughly ninety percent of Neurontin prescriptions for off-label use. Commentators and courts criticize manufacturers for egregious promotional efforts, including practices such as giving misleading information about drugs to physicians and offering them bribes and kickbacks. Manufacturers may be supporting the publication and dissemination of articles that suggest off-label drug usage with insufficient scientific support. Excessive off-label drug promotion and usage threatens to circumvent public oversight of drug safety and efficacy. Off-label drug use itself may be dangerous and ineffective, and manufacturers may be exacerbating the problem through their off-label promotional efforts.

As noted earlier, physicians may freely prescribe off-label, but there are restrictions on manufacturers’ ability to promote off-label usage of their products. The FDA allows manufacturers to distribute copies of peer-reviewed journal articles discussing off-label usage, but summarizing such articles might subject manufacturers to prosecution. Manufacturers may also be allowed to discuss off-label usage in response to unsolicited demands.


55. See, e.g., Meier, supra note 53; Rodwin, supra note 38, at 657.

56. See Rodwin, supra note 38, at 656; see also Sergio Sismondo, Key Opinion Leaders and the Corruption of Medical Knowledge: What the Sunshine Act Will and Won’t Cast Light On, 41 J. L. MED. & ETHICS 1, 640 (2013).

57. See Rodwin, supra note 53 at 659.


requests from physicians. The FDA has been criticized for offering insufficiently clear guidance as to appropriate promotional behavior regarding off-label drug usage.

If manufacturers improperly promote off-label usage, they can be held criminally culpable for misbranding under the FDCA. The government has repeatedly convicted pharmaceutical companies and their representatives based on their off-label promotional activities. As discussed further in Part V, manufacturers also face sanctions for off-label promotion under the False Claims Act.

If Medicare did not reimburse for off-label drug usage, the government would not be providing a direct incentive for manufacturers to promote off-label usage. Both CMS and states have acknowledged, though, that there are some off-label uses that are beneficial, and there are, therefore, benefits to legal reimbursement for some off-label prescriptions. Complicating matters is that the government often has weak and incomplete information regarding off-label drug usage and may not even know when reimbursements are for off-label usage. The Inspector General of the Department of Health and Human Services ("HHS") has already proposed a clear reform to improve tracking of off-label usage, allowing CMS to determine promptly whether reimbursements are appropriate. As proposed in Part IV, this reform should be supplemented by tying reimbursement rates to the indication for which a drug is used, rather than to only the drug itself.

The tension between a regulatory scheme that attempts to restrict manufacturers’ encouragement of off-label drug usage while acknowledging the value in physicians prescribing off-label drug usage has manifested in various judicial decisions. The Supreme Court recognized that off-label prescribing “is an accepted and necessary corollary of the FDA’s mission to regulate.” While Neurontin’s manufacturer was fined for its off-label marketing efforts, the FDA also approved some off-label

61. Id.
64. See, e.g., United States v. Caronia, 703 F.3d 149, 152 (2d Cir. 2012).
65. See Ausness, supra note 58, at 1325-1326.
66. See OIG PART D REIMBURSEMENT REPORT, supra note 6, at 2.
67. See id. at 1.
68. See id. at 6.
70. Buckman Co., 531 U.S. at 350.
uses of Neurontin. Courts have debated the importance of scientific truth in manufacturer’s promotional efforts, but this is typically difficult in off-label cases, because the FDA itself does not know the truth about whether a drug’s off-label use is effective, and it is unlikely that a court could do better than an expert agency in evaluating drug effectiveness.

III. An Inherently Flawed Reimbursement System Drives Excessive Costs

The present focus on scientific truth and punishing deceptive manufacturer behavior overlooks the broader problem: because the present reimbursement system is flawed, all profit-seeking behavior contributes to excessive reimbursements under Medicare Part D. Given the system’s present design, Medicare will end up spending excessively on off-label conditions, and even honest manufacturers will naturally over-invest in driving off-label drug usage.

Focusing on manufacturer violations of pharmaceutical promotional rules may help limit disinformation, but it is unlikely to stem the tide of excessive reimbursements under Medicare Part D. Manufacturers can follow every rule and regulation regarding off-label promotion and they will still have every incentive to over-invest in encouraging and developing off-label drug usage. Investment is a broad concept covering a manufacturer’s behavior in pursuing revenue from off-label drugs; it includes research into off-label efficacy, for example, and it is not limited to promotional behavior that directly engages physicians. This over-investment will continue to drive excessive Part D reimbursements.

This part demonstrates the danger of this over-investment contributing to excessive Medicare reimbursements. The article here makes every assumption in favor of manufacturers and demonstrates that, nonetheless, the present system will continue to drive excessive drug reimbursement levels. The situation may actually be worse in reality, given that manufacturers may not always act in the public interest as assumed here.

A. A standard for excessive costs

Analyzing excessive costs begins with a general principle that drug costs are excessive if they exceed the social benefit obtained from a patient utilizing the drug. If there is no scientific evidence that a drug is


72. See, e.g., Caronia, 703 F.3d at 166-67; See Mark Ratner & Trisha Gura, Off-Label or Off-Limits?, 26 NATURE BIOTECHNOLOGY 867, 873-74 (2008).

73. See Christopher Robertson, When Truth Cannot Be Presumed: The Regulation of Drug Promotion Under an Expanding First Amendment, 94 B.U. L. Rev. 545, 560 (2014) (“FDA defers to physician discretion to prescribe off label, because it remains ignorant about safety and efficacy claims until they are proven.”).
safe and effective for an off-label condition, then any Medicare spending on that drug for that off-label condition is excessive. This would be the archetype of purely wasteful spending, as patients do not benefit at all from taking unsafe, ineffective drugs. At the other end of the spectrum are FDA-approved drugs for the treatment of an on-label condition for which CMS is willing to reimburse. Given that cost-benefit analysis is part of CMS’s approval process, it is safe to presume that spending on such drugs for the on-label condition is not excessive.

Drugs prescribed for off-label conditions will fall somewhere along this spectrum. Some drugs may have an extremely limited number of scientific studies supporting their efficacy for off-label conditions—those drugs will fall closer to the wasteful end of the spectrum. Other drugs may have excellent studies supporting the off-label usage—these drugs will lie close to the FDA-approved on-label end. There is therefore some optimal level of Medicare spending on the drug for the off-label condition that balances the costs of the drug against the benefits patients may obtain from the drug. Note that this optimal level of Medicare spending is specific to a drug-condition combination. If a drug can treat two distinct conditions, there will be an optimal level of Medicare spending for condition A and a separate optimal level of spending for condition B. Similarly, if a condition can be treated by two distinct drugs, there will be an optimal level of spending for drug X for that condition and a separate optimal level of spending for drug Y for the same condition.

Expanding this analysis, consider that Medicare spending on drugs has at least two related purposes. First, CMS has an immediate interest in ensuring that patients receive safe, effective treatment that is presently available. Second, CMS has a long-term interest in manufacturers producing new safe and effective drugs.

Regarding CMS’s immediate interest in ensuring that patients receive treatment, an optimal level of spending corresponds to the safety and effectiveness of the drugs. If the drug is highly beneficial to patients suffering from a costly condition, the optimal level of spending would likely be higher. If a drug produces only limited benefits for a small portion of patients suffering a mild condition, the optimal level of spending may be lower. If there are two potential drugs for the treatment of one condition and they are identical in safety and effectiveness, any spending on the more expensive drug is excessive unless there is some other justification for such spending. For example, CMS might value having competition in the market supply of the drugs. To maintain the viability

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74. If the drug is actually harmful to patients, society should actually invest in preventing access to such drugs, even if they were free to the Medicare system.

75. See Pattners, supra note 35, at 55.

76. This competitive interest may be linked to CMS’s below long-term interest in new drug development. Competition may also be important in negotiating prices for existing drugs.
of the manufacturer producing the more expensive but otherwise identical drug, CMS might agree to reimburse for some limited level of the more expensive drug. Spending may be excessive if it is heavily concentrated on relatively ineffective drugs when safe, more effective alternatives are available.

CMS’s long-term interest in manufacturers producing new safe and effective drugs interacts with its immediate interest in patient treatment. A naïve method of compensating manufacturers for drugs would be to reimburse at the marginal cost of production for those drugs. Manufacturers, in theory, would continue to produce existing drugs, but such a strategy would strip out the profit incentive for future drug research and development. The reimbursement rates for drugs thus must be sufficiently high to induce manufacturers to conduct ongoing research and development of drugs. Excessive spending, considering CMS’s long-term interest in ongoing research, could come in the form of market distortion. For example, CMS’s willingness to reimburse for a high-priced drug is a signal to the marketplace that there is a strong need for safe, effective treatment of the particular condition the drug treats. CMS would expect manufacturers to react to the high price signal by investing in new drug development for that particular condition. If CMS is reimbursing for a high-priced drug, but there is already a safe, equally effective, and lower-cost treatment for the same condition, CMS expenditures on the high-priced drug might induce other manufacturers to continue emphasizing drug research for that same condition. Such expenditures and investments might be excessive, as the existence of the lower-cost option would suggest that other conditions should be research priorities. Under these principles, CMS should place the strongest reimbursement incentives upon FDA-approved drug treatments for on-label conditions. These are drugs that have crossed a threshold of scientific evidence in establishing safety and efficacy for certain conditions. At the other end, if there is no scientific evidence of safety and efficacy for the treatment of another condition, there should be no Medicare reimbursement for that drug-condition combination.

In between, a drug for which there is limited scientific evidence of off-label efficacy should receive an intermediate level of reimbursement incentive. If CMS were to allow reimbursement at the same level as an FDA-approved on-label treatment, there would be no incentive for the manufacturer to continue research and testing to satisfy FDA standards for an off-label use. If CMS were to prohibit any reimbursement for this intermediate case, patients might not have access to the drug. Allowing intermediate reimbursement strikes a balance for patients who might benefit from a drug that has not completed scientific-efficacy studies; those patients will receive the drug, but the manufacturer will not receive more reimbursement than a competing manufacturer that has completed those scientific studies demonstrating efficacy.

There are thus multiple criteria by which Medicare drug expenditures may be excessive. This section is not a comprehensive list of parameters in
determining optimal Medicare spending on off-label drugs, nor is this a claim that society can necessarily establish the precise, optimal level of spending on any particular drug. Rather, it is rather a claim that there are different levels of scientific evidence supporting the efficacy of drugs and that it is excessive to spend more money for drugs that have lower levels of scientific evidence supporting their efficacy. The Medicare spending framework will have a tendency to drive patients, physicians, and manufacturers to either better levels of spending that are closer to optimal or worse levels of spending that stray further from optimal.

B. Even an idealized manufacturer contributes to excessive drug reimbursement under the present system.

The next step in analyzing off-label drug costs under Medicare Part D is to consider manufacturer behavior. Instead of focusing on manufacturer misbehavior, a problem both courts and commentators have discussed at length, this section discusses the impact of ethical profit-seeking manufacturers on Medicare Part D off-label expenditures. While unethical, avaricious manufacturers can cause excessive drug expenditures, this section demonstrates that even ethical profit-seeking manufacturers within the existing reimbursement framework lead to excessive Part D off-label expenditures.

One beginning premise is that off-label drug usage is, in the short term, good for society. Doctors can be trusted to treat their patients properly, and if a doctor believes that prescribing a drug for an off-label use is a good choice, this section assumes that the patient will benefit from taking that drug. Nonetheless, it is important to recognize that physicians are limited in their availability and capacity to learn about new drug uses. Physicians do not instantaneously learn about new drug uses; they have limited time to both treat patients and study new treatment developments. Moreover, physicians have very limited information regarding drug prices.

77. See supra Part II.C.
78. See 21 U.S.C. § 396 (2012). While the FDA’s lack of involvement in physician off-label prescription can be described as a reluctance to interfere with the practice of medicine, this similarly suggests that there are positive aspects to off-label prescription. If a particular off-label prescription were generating consistently bad outcomes for patients, it is difficult to believe that a regulatory agency would not take action.
The second assumption is that pharmaceutical firms are broadly acting in the public interest. This assumption automatically rules out deliberate lies and deception regarding off-label drug usage. Moreover, this implies that any increased investment that the firms make in off-label research corresponds to increased public good. For example, this means that if a firm increases its spending on off-label research, it conducts legitimate clinical research on safety and efficacy and it disseminates the results of that research.

The third premise of this section is a focus on off-label usage of a patent-protected drug in a market of patent-protected drugs. Once patent protection expires, competition from generics and other manufacturers may reduce prices and may even provide insufficient incentive for off-label research. Thus, in this section, the market price of drugs refers specifically to the competitive market of patent-protected drugs.

1. Manufacturer pricing

First consider the manufacturer’s pricing mechanism. After completing scientific studies and obtaining FDA approval for a drug, how does a manufacturer set the price of its patent-protected drug? As many have acknowledged, manufacturers set prices at whatever the market will bear. This is standard profit-seeking behavior, and this piece is not criticism of such behavior. Following general market theory, the drug will be sold for a price that corresponds to the benefits that a patient expects to receive from the drug. In other words, a drug that offers little benefit will not command a high price.

Rather, the challenge leading to excessive Medicare reimbursements is that the manufacturer will set the drug price at a level tied solely to the market for the on-label condition. There is little reason to believe that the market for a drug’s off-label condition will be tightly linked to the


82. For a brief discussion of this unique environment, see Rebecca S. Eisenberg, The Problem of New Uses, 5 YALE J. HEALTH POL’Y, L. & ETHICS 717, 720 (2005).

83. There is empirical data to suggest, however, that drug prices even with generic competition may not necessarily be lower. See Panos Kanavos, Joan Costa-Font & Elizabeth Seeley, Competition in Off-Patent Drug Markets: Issues, Regulation and Evidence, 23 ECON. POLICY 500, 500-01 (July 2008).


85. The present Medicare Part D reimbursement system does not have visibility into the treated condition, thus limiting downward pressure on pricing. As I discuss later in Part VI, though, it is possible that market pressures outside of the Medicare system may create price pressure on the manufacturer.
market for the on-label condition. To the extent that the market price for drugs treating the off-label condition is actually lower, Medicare reimbursement based on drug prescription independent of condition will be excessive.

For example, begin with a firm that has obtained FDA approval for a new drug X for the treatment of condition A. It sets the price for drug X by evaluating the market of treatments for condition A. If existing treatments for condition A are limited, expensive, and not particularly effective, the firm may be able to charge a high price for drug X. Assume that the firm charges $5000 per dose of drug X given the market for condition A. Parallel research determines an off-label use for drug X in treating condition B, but the market for condition B is much more competitive. Even if the firm obtained FDA approval of drug X for treating condition B, the firm believes it could only charge fifty dollars per dose of drug X if it were selling drug X solely for the treatment of condition B.

In this scenario, Medicare reimbursement for drug X at the $5000 price for an off-label treatment of condition B is excessive. This claim is based on the assumption that market prices are a proxy for the harm associated with the treated condition. If the market price for the off-label condition is significantly lower, it implies that the harm from the off-label condition is significantly lower. Thus, Medicare expenditure at the higher drug price for condition B is likely excessive.

It is nonetheless possible that the drug could be worth the full price of reimbursement, regardless of indication. In the above example, even though the $5000 per dose price is linked to the on-label condition, it is theoretically possible that an educated consumer would be willing to pay that price for the off-label condition. We might think a physician would

86. This depends, of course, on the type of off-label scenario at play. Some types of off-label usage involve patient groups that have not been adequately addressed in studies (i.e., drugs not tested on children under the age of two). See Alexandra Ossola, FDA Allows Company To Market Drug For Off-Label Use: Some Experts Predict That More Companies Will Try To Do The Same, POPULAR SCIENCE (Mar. 10, 2016), http://www.popsci.com/fda-allows-company-to-market-drug-for-off-label-use; see also COMMITTEE ON ACCELERATING RARE DISEASES RESEARCH AND ORPHAN PRODUCT DEVELOPMENT, BOARD ON HEALTH SCIENCES POLICY, RARE DISEASES AND ORPHAN PRODUCTS: ACCELERATING RESEARCH AND DEVELOPMENT 192 (2010).

87. The assumption that market prices are a relative proxy for harm assumes that there is some rationality in the marketplace. This assumption is more credible as a proxy for the minimum level of harm caused by a condition: a patient, insurance carrier, or physician would not purchase the drug if the harm of the condition did not exceed the price of the drug. In contrast, it is possible that the harm of the condition greatly outweighs the price of the drug, and the consumer is getting a great deal by paying a low price to remove a great harm. See Richard E. Caves et al., Patent Expiration, Entry, and Competition in the U.S. Pharmaceutical Industry, 1991 BROOKINGS PAPERS ON ECONOMIC ACTIVITY: MICROECONOMICS 1, 7 (1991).

88. See id. at 5 (describing patients as unlikely to select physicians based on the physicians’ drug prescription behavior).
be best suited to make such a determination. In reality, of course, most physicians are not aware of the particular price or reimbursement rates of drugs, but they at least may be aware of insurance coverage and the ability of patients to actually obtain prescribed drugs. Even if, in the short term, this off-label drug is worth the full price of reimbursement, the next section describes the method by which improper incentives drive excessive drug costs.

As a final note, while this article focuses on utilitarian concerns, there is also an underlying fairness concern that supports this argument. To the extent that there are already drugs that are effective in treating condition B that have obtained FDA approval for the condition, it seems inequitable to allow drug X, which has not obtained FDA approval for condition B, to receive a higher price for treating condition B.

2. Manufacturer investment

One possible criticism of the aforementioned concern with manufacturer pricing is that costs may balance out; sometimes Medicare pays an excessive amount for an off-label treatment, but sometimes Medicare gets a good deal because the off-label drug is actually cheaper than the competitive products for the off-label condition.

This leads to the next problem: the manufacturer’s response to this incentive structure. The above costs are unlikely to balance out because of the manufacturer’s investment decisions given a pharmaceutical market with manufacturers setting prices as described in the prior section. Broadly speaking, manufacturers that see the opportunity to profit from off-label usage are likely to invest in off-label usage, but if Medicare is instead getting a good deal, it is unlikely that manufacturers will similarly invest in such off-label usage.

Following the earlier example, begin with a manufacturer that has obtained FDA approval for drug X in treating condition A. At this point, there may be zero scientific evidence regarding the efficacy of the drug for the off-label condition B. The optimal spending on research into drug X’s efficacy in treating condition B depends on two factors: first, the manufacturer’s beliefs about the future efficacy of such research, and second, the size of the market for condition B.

It is possible that those factors will lead the manufacturer to actually obtain FDA approval of drug X for condition B. Since this paper discusses off-label usage, though, I assume that either the market for condition B or the cost of the research somehow makes obtaining FDA approval for drug X for condition B infeasible for the firm. Nonetheless, because of the

89. See TEMIN, supra note 80, at102-06.
90. It is also possible that the manufacturer may be unwilling to invest in the necessary research because of the risk of discovering some side-effects that would jeopardize its original FDA-approved indication. See COMMITTEE ON ACCELERATING RARE DISEASES RESEARCH AND ORPHAN PRODUCT DEVELOPMENT, supra note 88, at 192.
possibility of off-label usage and reimbursement, the firm will invest some non-zero amount into condition B research. Roughly speaking, the amount the firm invests into condition B research corresponds with the expected revenue from condition B reimbursement. If reimbursement for drug X is fixed regardless of condition, then the firm will invest in condition B research at a level corresponding to the price determined by condition A—in this example, $5000 per dose.

Note, however, that the price of $5000 may have no correlation with the market for condition B. There may be already effective, patent-protected drugs that treat condition B and have a significantly lower price, and drug X might not be any more effective. If the manufacturer estimates the market size for condition B using the $5000 per dose value, the market opportunity for condition B will be much greater than the market value based upon the present, lower-cost drugs available for condition B.

Faced with this incentive structure, the manufacturer will over-invest, leading to societal losses. The price discrepancy is a distorted allocation of research funding. To the extent manufacturer investment is on research, this spending is misallocated; Medicare does not actually prioritize investments in condition B research at a level corresponding with a drug priced at $5000 per dose, and society would be better off if the manufacturer invested in other research. Stated another way, if the market price of a competitor drug in treating condition B is fifty dollars per dose, Medicare would not encourage manufacturers to invest at a market level corresponding to $5000 per dose for condition B; there are other conditions worthier of investment. From the drug X manufacturer’s perspective, though, there is a large revenue opportunity in pushing drug X for condition B. It is possible that drug X might actually be one hundred times more effective and safe in comparison to the existing competitors in treating condition B, but charging a hundred-times higher rate should require FDA approval.

Note that an investment incentive problem still exists if the off-label market price is higher than the on-label market price; i.e., a fifty dollar per dose drug has an off-label use for which competitors are charging $5000 per dose. In this situation, manufacturers face insufficient incentive to invest in off-label usage. The result will be insufficient research and promotion of cost-saving off-label drug usage. In other words, Medicare expenditures will be higher than optimal, as there will be relatively increased usage of the FDA-approved drugs for condition B and less-than-optimal research and information supporting cheaper off-label drugs for condition B.

A further complication is the fact that the manufacturer will split its investment in the off-label condition between research and promotion.

91. Some have described drug regulations as an incentive for producing knowledge about the drug. See Robertson, supra note 73, at 561.

92. See, e.g., Caves, supra note 87, at 2.
Given that research has a downside, there is a serious risk that the investment may favor promotion over research. A manufacturer considering further research must consider the possibility that subsequent clinical research will reveal weaker results or worse side effects. Such negative clinical findings could jeopardize not only use of the drug for the off-label condition, but also its use for the on-label condition. The general problem here is that manufacturers may have a difficult time capturing benefits from further off-label research. If subsequent research reveals weaker results or worse side effects, society is better off learning about the weaker results or worse side effects. Unfortunately for the manufacturer, it is in the business of selling drugs and not information. When its research reveals these negative results, society benefits from the knowledge, but it is difficult for the manufacturer to profit from such negative knowledge.

In contrast to research, manufacturers will likely capture much of the benefit from promotional activity. Promotional activity likely increases physician awareness and thus propensity to prescribe the manufacturer’s drug. Following the basic assumption that manufacturers do not act deceptively, society benefits from the increased physician knowledge. The manufacturer will benefit from revenue due to the reimbursement for the prescribed drug.

The fact that a physician learns about the drug’s off-label uses, however, does not automatically mean that increased promotional activity in distributing knowledge is universally desirable. Physicians have limited time and mental resources; learning new information is constrained by those resources. If physicians are limited in their time to listen to manufacturers’ reps, those physicians may disproportionately favor drugs.

93. See Robertson, supra note 73, at 559-60.
94. Id.
95. In 2004, Merck was forced to remove from the market its $2.5 billion Vioxx product, a drug approved for the treatment of arthritic pain, when clinical studies on an alternative treatment, preventing recurrence of colon polyps, revealed increased cardiovascular side effects from the drug. See Barbara Martinez et al., Merck Pulls Vioxx From Market After Link to Heart Problems, WALL ST. J. (Oct. 1, 2004), http://www.wsj.com/articles/SB109654671320932405.
96. See Robertson, supra note 73, at 561 (“Information is needed to make product markets perform optimally, but if sellers are to provide that information then they must be given an incentive to do so.”) (citing Howard Beales et al., The Efficient Regulation of Consumer Information, 24 J.L. & Econ. 491, 504 (1981)).
97. See, e.g., Caves supra note 87, at 5 (citing TEMIN, supra note 91) (describing physicians’ lack of “ready and well-organized information” regarding drug choices).
98. See id. at 4-8.
99. See id. at 5 (citing TEMIN, supra note 91) (describing physicians’ lack of “ready and well-organized information” regarding drug choices).
whose manufacturers dedicate greater promotional resources.\textsuperscript{100} Regardless of the actual impact on physician prescriptions, there is the social loss of spending physician time on relatively weak scientific studies that do not rise to the same level of the studies supporting FDA-approved drugs and conditions. Stated another way, the result of increased manufacturer investment may be a lot of distracting noise that makes it more difficult for physicians to focus on relevant new information.\textsuperscript{101}

\textbf{C. Existing limits to excessive off-label expenditures}

While off-label reimbursement is difficult to detect under the present Medicare Part D system, there are limits to manufacturers’ ability to profit in such a manner.

One possible limitation is that off-label reimbursements might come to dominate on-label reimbursement. If only a small number of patients suffer from the on-label condition, but Medicare is paying for a much larger volume of the manufacturer’s drug, this would attract significant attention.\textsuperscript{102} Rather than identifying specific prescriptions that are for off-label usage, the aggregate data would provide a conservative estimate of the off-label usage volume. If there are only 100,000 patients with condition A and Medicare is reimbursing for 500,000 patient-doses of drug X, CMS might reasonably be suspicious of drug X. The benign assumption is that such high levels of drug X reimbursement are due to off-label usage, but such high reimbursement might also be a signal of fraudulent billing. Either way, the suspicion could drive CMS to begin requiring preauthorization or other administrative controls on the prescription of the manufacturer’s drug. Such rules would not only limit off-label prescriptions of the drug but also hamper on-label prescriptions.

Manufacturers would not want to attract such attention. They therefore might limit promotional efforts to avoid exceeding some threshold that could trigger CMS investigation.\textsuperscript{103} Given CMS’s general reluctance to take investigative steps, though, I assume that this upper

\textsuperscript{100} Id. at 12 (describing large volume of advertisements as a “signal-jamming” strategy to fight competitor information).

\textsuperscript{101} Id.

\textsuperscript{102} For example, 83% of physician prescriptions for Gabapentin (Neurontin) were for off-label uses. David C. Radley et al., \textit{Off-Label Prescribing Among Office-Based Physicians}, 166 ARCHIVES INTERNAL MED. 1021, 1021 (2006). This led to subsequent litigation. See \textit{id.} at 1026.

\textsuperscript{103} More formally, a manufacturer would conduct research and promote the off-label use of its drug until the marginal benefits from doing equaled the marginal costs of such research & promotion. Those marginal benefits would be severely reduced if CMS instituted investigations into its drug. See \textit{CTR. FOR MEDICARE & MEDICAID SERVS., OFF-LABEL PHARMACEUTICAL MARKING: HOW TO RECOGNIZE AND REPORT IT} (Oct. 2015), available at https://www.cms.gov/Medicare-Medicaid-Coordination/Fraud-Prevention/Medicaid-Integrity-Education/Downloads/off-label-marketing-factsheet.pdf.
limit to the manufacturer’s over-investment is not sufficiently low to justify non-intervention by other means.

D. The role of sponsor competition

The existence of multiple Part D sponsors increases the complexity of the manufacturer’s decision. First, simply to maximize profit and revenue from its drug, the manufacturer should ensure that its drug is included in all sponsors’ formularies. This could be accomplished by ensuring that the drug is part of Part D’s list of required drugs that every sponsor must include, or it could be accomplished through individual negotiation with each sponsor. Working from the assumption that the manufacturer’s drug can command a high price, it is reasonable to believe that there is little competition for the drug and its on-label treatment condition. With little or no competition, inclusion in formularies should be relatively straightforward.

Uniform inclusion of the drug in all sponsors’ formularies is also important in protecting its off-label profitability. If the manufacturer fails to include the drug in a limited number of sponsors’ plans, the manufacturer may create a negative feedback cycle that will damage its off-label earning potential. This negative feedback cycle is triggered by the fact that a sponsor that does include the manufacturer’s drug will be at a relative disadvantage to a sponsor that does not include the manufacturer’s drug. A sponsor that includes the manufacturer’s drug for its on-label condition faces increased costs for the off-label condition. These increased costs may reduce that sponsor’s competitiveness in contrast with a sponsor that does not cover the manufacturer’s drug. Such reduced competitiveness may trigger the sponsor to conduct research that would identify the manufacturer’s drug as the cause of its comparatively higher costs. If all sponsors include the manufacturer’s drug in their formularies, though, there is less risk of this reduced competitiveness triggering investigation of the manufacturer.

E. An instrumental need for off-label revenue

Critics of the argument in III.B. might claim that off-label revenue is important in getting the drug out at all. It is possible that manufacturers rely upon the off-label revenue to support their investment in the FDA-approval process for the on-label condition. The argument, then, is that the manufacturer would not even invest in the drug for any FDA approval, because it believes that there is insufficient potential revenue for the on-label condition to justify its investment. This is an open empirical question, although there are commentators who argue that manufacturers overstate the actual investments necessary to develop FDA-approved drugs.104

104. See, e.g., Donald W Light, Rebecca Warburton, Demythologizing the High Costs of Pharmaceutical Research, 6 Biosocieties 34, 34 (2011); See also Joseph A.
Another potential criticism is that manufacturers naturally under-invest in off-label conditions. The threat of patent expiration and generic competition may generally induce under-investment in off-label research in patent-protected drugs.\textsuperscript{105} Thus, allowing for high reimbursement rates for off-label prescriptions provides an incentive for manufacturers to conduct a limited level of research prior to patent expiration, even if those manufacturers are not conducting sufficient research to satisfy FDA approval requirements.

These arguments are of secondary importance. The present system obfuscates the connection between drugs and the conditions being treated. Moreover, the potential positive instrumental benefits of the existing system are due to chance: a drug happens to be effective for certain on-label and off-label conditions. The above goals can be better pursued via direct, more visible means of subsidy and promotion.

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In summary, the existing government reimbursement system for off-label drug usage is inherently flawed. The improper incentives and unfair windfall profits will lead even scrupulous manufacturers to over-invest in off-label activity, resulting in excessive expenditures under Medicare Part D.

IV. A theoretical solution: reimbursement linked to competitor pricing

The above theoretical model focuses on excess drug expenditures resulting from the present system; in reality, there are more harms that may result. As discussed earlier, physicians might be insufficiently or improperly informed about the costs and benefits of off-label drug use, and manufacturers might not be completely honest and transparent in their promotional efforts. Patient health and safety may also be at risk.

Because of those additional harms, some have suggested eliminating manufacturer profits from off-label prescriptions,\textsuperscript{106} proposing reimbursing off-label prescriptions at the marginal cost of production for the drug, which effectively eliminates any profit from the off-label prescription and sale.\textsuperscript{107} This proposal may be the best solution; a thorough analysis depends on the extent to which firms behave badly in response to the off-label incentives in comparison to the potentially good responses described in Part III. If the potential revenue from off-label drug usage drive manufacturers towards socially harmful activities that

\textsuperscript{DiMasi et. al., The Price of Innovation: New Estimates of Drug Development Costs, 22 J. OF HEALTH ECON. 151, 151 (2003).}

\textsuperscript{105. See, Caves, supra note 87, at 1-2.}

\textsuperscript{106. Rodwin, supra note 38, at 659-660.}

\textsuperscript{107. Id. at 659.}
outweigh beneficial activities, then this proposal to eliminate any off-label reimbursement profit is the right solution.

A. Capping reimbursement by reference to an FDA approved competitor

If it is the case, however, that firms have the public interest in mind, eliminating all incentive for off-label efforts may be detrimental to society. Instead of reimbursing at the marginal cost of production, this article proposes that a fair reimbursement rate would be capped at a rate tied to the competitive, patent-protected market for treatment of the condition. Such a cap would ensure that patients would still have access to drugs for off-label indications, while improving manufacturers’ incentives for research. This cap could take multiple forms.

One option that results in the greatest amount of fairness is to cap at the lowest-priced FDA approved competitor. Under such a system, a manufacturer who has not received FDA approval for the off-label condition could not receive reimbursement higher than any competitor who has received FDA approval for treatment of the condition. This would not eliminate the incentive for incremental off-label research and promotion. Instead, setting such a reimbursement rate would give manufacturers incentive to invest in some level of off-label research and promotion, but such incentive would be no greater than the incentive enjoyed by an FDA-approved competitor.

An alternative cap would be to set the maximum reimbursement rate at the second-highest priced FDA-approved competitor. This might reduce some of the fairness of the first option, in that some FDA-approved competitors might receive a lower reimbursement rate than manufacturers who have not received FDA approval. Nonetheless, it is possible that this cap would provide superior incentives for manufacturers who had not obtained FDA approval. The lowest priced FDA-competitor may be a remarkably low-efficacy product that was approved at a time when no other treatments were available for the relevant indication. The newer off-label drug may be more comparable in efficacy to the best FDA-approved drugs on the market. Allowing the off-label drug to be reimbursed at the second-highest FDA-approved competitor price may provide a better incentive despite the potential unfairness. Some auction theories suggest that this second-highest price may be a good choice.\(^{108}\)

There are other cap proposals that could be justified, such as a cap linked to the mean or median reimbursement rate of the FDA-approved competitors and there will be similar trade-offs between fairness and potential incentives. This article does not take a position as to the best particular cap. Rather, the important core is that the cap must somehow be linked to the market of FDA-approved competitors.

\(^{108}\) See, e.g., William Vickrey, *Counterspeculation, Auctions, and Competitive Sealed Tenders*, 16 J. Fin. 8, 8 (1961).
B. An extended proposal: reimbursement tied to competitive indication

The aforementioned cap proposal seeks to limit excessive Medicare Part D expenditures on off-label drugs that result from the presently flawed reimbursement system. There is a flip side to this discussion, though: the possibility that the flawed reimbursement system also benefits Medicare by obtaining cheaper off-label drugs. The prior section focused on scenarios in which the off-label drug commands a higher price than the prevailing FDA-approved competitors. There is also the possibility that the drug price is substantially lower than the competitive price for the off-label condition. Under the present system, to the extent the lower-price drug is used for an off-label purpose and is safe and effective, Medicare is actually getting a good deal in the short term; the patient is receiving treatment at a substantially lower drug cost than she would have if she were receiving a drug approved for the condition.

The arguments raised in Part III regarding incentives nonetheless apply in this situation, too. In this case, however, manufacturers may under-invest in off-label research under the present regime. Compared to companies that are looking into new, patentable drugs specifically for the off-label condition, manufacturers who have an existing drug at a relatively lower price will not invest as much because of their weakened ability to command a higher price. If the manufacturer unilaterally raises the price for all customers, they may receive tremendous pushback in the marketplace and negative media attention.\footnote{For example, consider the recent outcry regarding Mylan NV and its EpiPen price increase. See Louise Radnofsky, \textit{EpiPen Maker Executive to Testify at House Hearing}, \textit{Wall St. J.} (Sept. 15, 2016), http://www.wsj.com/articles/epipen-maker-executive-to-testify-at-house-hearing-1473894399.} Given the existing disconnect between diagnosis and prescription, though, manufacturers have no way of charging different prices to Medicare for the same drug. Manufacturers dealing with an existing drug will likely be stuck at the lower reimbursement rate of the on-label condition.

Thus, a broader proposal would be to tie all reimbursement rates to the competitive indication rates. Medicare would thus reimburse for a specific indication rather than a specific drug. Rather than a physician prescribing a specific drug and the manufacturer receiving reimbursement at a negotiated price, all manufacturers would receive the same reimbursement price when their drug is used. A manufacturer in the above situation could then benefit from higher rates as long as the patient had the off-label condition, and such a manufacturer would then face comparable incentives for investment.

In the short term, this extended proposal is likely to lead to higher Medicare drug expenditures in comparison with the above cap proposals, because it would allow a manufacturer to benefit from higher reimbursement rates if the price for the off-label condition is higher than the on-label reimbursement rate. In the long term, however, correction of this incentive problem should induce greater research for the off-label
condition, resulting in greater manufacturer competition for the off-label condition and eventually lower prices.

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Note that these proposals to cap reimbursement rates fall under Part III’s theoretical model of an idealized manufacturer. To the extent there are additional harms, the cap should actually be lower. In fact, if the harms from off-label usage are sufficiently severe, even proposals to cap rates at the marginal cost of production may be insufficient. Rather than allowing reimbursement at the marginal cost of production, perhaps no reimbursement should be allowed at all.

Also consider that this proposed solution focuses solely on the market for patent-protected drugs. Once generic equivalents enter the marketplace, there is reason to believe that prices will drop to a point where there may be insufficient manufacturer incentive for further research. Thus, in analyzing the competitive market for any specific condition, the reimbursement cap must focus solely on the patent-protected competitors. For example, if there are five FDA-approved treatments for a condition and two of the treatments have lost patent protection and have generic equivalents available, the reimbursement cap would only consider the three other FDA-approved treatments. Even under this formulation, the existence of the generic products may still have some downward influence on the market price, but excluding such influence is likely to be difficult and of limited benefit.

Regardless of the precise optimal reimbursement rate, all reform proposals hinge upon one critical piece of information: tying patient indication to the prescription. Without this key piece of information, this article’s proposed reforms are infeasible. Reimbursement based upon indication requires that we actually have the indication for which a drug was prescribed.

Implementing such a change in reimbursement would require significant statutory, regulatory, and professional changes. These changes will be difficult, though, and unlikely in the short term. First, the FDA and CMS have demonstrated a long-standing commitment to avoiding interference with the practice of medicine. Forcing physicians to fundamentally change their drug prescription process seems contrary to its non-interference commitment. As CMS has noted, including a diagnosis in a prescription is not presently standard practice. Moreover, information included with prescriptions is generally governed by state law and outside of CMS’s present statutory authority. Finally, present

110. See Beck & Azari, supra note 12, at 76. (stating that the FDCA was “not intended as a medical practices act and [did] not interfere with the practice of the healing art.”); see also Weber et al., supra note 7 (quoting Jonathan Blum, director of Medicare, that agency philosophy “really has been to defer to physicians.”).

111. OIG PART D REIMBURSEMENT REPORT, supra note 6, at 6.

112. Id. at 6.
coding standards for diagnosis are not sufficiently detailed to correspond with CMS rules regarding medically accepted indications.113

V. FCA liability as a claw-back for windfall off-label profits.

Despite the above barriers to a first best solution, there are immediate steps that could be taken to reduce the excessive costs stemming from the flawed Part D reimbursement scheme for off-label uses of drugs. I begin this part with an overview of existing short-term proposals, followed with a proposal for expanding use of the civil False Claims Act.

A. Interim solutions are not priorities

The Office of the Inspector General (“OIG”) of HHS has recognized the core informational disconnect within Medicare Part D and has mentioned a variety of proposals to address the problem.114 These proposals, however, all share one common flaw: they require CMS to shift limited resources towards addressing this challenge. As suggested in the CMS response to the OIG report, such increased resource allocation is unlikely and the detection of improper off-label drug reimbursement faces numerous obstacles.115 A review of the alternative OIG proposed interim solutions follows.

1. Prior authorization

Prior authorization is a prepayment strategy that could provide the missing informational link between prescription and diagnosis. Prior authorization requires explicit authorization from Medicare prior to a patient obtaining drugs.116 Presently, CMS permits Medicare Part D sponsors to use prior authorization for certain drugs that are at high risk for prescription without a medically accepted indication.117 Prior authorization could be expanded, but it is viewed as a cumbersome, time-consuming process that limits patient access to drugs.118 Because of its cumbersome nature, regulations presently prohibit sponsors from using prior authorization for six classes of drugs.119 A more limited proposal is to require prior authorization for drugs exceeding a specific reimbursement cost.120

113. Id. at 8.
114. Id. at 9.
115. Id. at 8-9.
116. Id. at 2.
117. See id. at 8.
120. Gillick, supra note 118, at 346-47.
The present situation suggests that the Part D sponsors are not interested in expanding prior authorization, nor is CMS encouraging them to do so. This may be evidence that CMS and the sponsors are responsive to the patient and physician interest in ease of access to medications.

2. Post-payment Audits

Another option is requiring Part D sponsors to conduct audits of prior payments. Sponsors could obtain diagnosis information from physicians and retroactively compare those notes with drug reimbursements. Again, the main challenge here is that sponsors and CMS apparently do not seem motivated to conduct such audits.

A secondary, more technical problem is that CMS may approve payments that are part of certain drug compendia, but subscribers may only have access to the most recent version of those compendia. Because at least one compendium is updated on a quarterly basis, sponsors would have to complete audits on a timely basis.

Given the reluctance of CMS and its delegates to prioritize either an interim solution or the larger systematic challenge, filling in the information gap requires some third-party action. The civil FCA may fill this role, given its prominent involvement of whistleblowers. The FCA can serve as a temporary transition to a diagnosis-based reimbursement regime.

B. General FCA Background

The False Claims Act has become one of the most prominent tools in combatting fraud against the federal government. The FCA generally prescribes fraud or false claims against the federal government. The relevant mens rea for defendant liability is knowledge; the statute defines “knowledge” to include a person who “acts in reckless disregard of the truth or falsity of the information; and [the statute] require[s] no proof of specific intent to defraud.” The FCA has both civil and criminal provisions; this article focuses solely on the civil FCA.

121. See OIG PART D REIMBURSEMENT REPORT, supra note 6, at 2.
122. See id.
123. See id. at 1-2.
125. 31 U.S.C. §§ 3729-30 (2012). The civil FCA also has a criminal counterpart found in 18 U.S.C. § 287, but for reasons similar to the FDCA, I do not focus on criminal sanctions in this article.
Besides traditional public enforcement, the FCA also contains *qui tam* provisions, which allow private litigants—known as “relators”—to pursue civil actions and prosecute cases of fraud in lieu of the Department of Justice (“DOJ”).\(^{127}\) Today, relators can receive as much as thirty percent of the civil recovery, which can be substantial given the statute’s treble damages provisions. Civil penalties also include $5500 to $11,000 in fines per false claim. A successful relator is also entitled to legal fees from the defendant.\(^{128}\)

As a practical matter, though, the FCA is an information-providing system rather than a private-enforcement system.\(^{129}\) The vast majority of FCA cases in which the relator recovers from the defendants are DOJ-prosecuted cases.\(^{130}\) The law firms that represent relators in FCA actions generally specialize in obtaining DOJ intervention.\(^{131}\) Purely private enforcement of the FCA is generally either not pursued or unsuccessful.\(^{132}\) Thus, the FCA broadly functions as a whistleblower system in which relators provide information to the DOJ and the DOJ decides whether or not to pursue the defendant based on such information.\(^{133}\)

C. The existing theory of off-label promotion as an FCA violation

Roughly sixty percent of FCA cases today involve allegations of healthcare fraud.\(^{134}\) Some of the largest settlements generally involve off-label promotion claims.\(^{135}\) The FCA’s present role is contentious for a variety of reasons and this article’s proposed solution of leveraging FCA liability as a claw-back mechanism is likely to be similarly contentious.


\(^{128}\) Id.


\(^{131}\) See Kwok, *supra* note 129, at 237-38.

\(^{132}\) FALSE CLAIMS ACT STATISTICS, *supra* note 130.

\(^{133}\) See Kwok, *supra* note 129, at 226-30; FALSE CLAIMS ACT STATISTICS, *supra* note 130.

While FCA cases typically address direct fraud against the federal government, such as a healthcare provider billing Medicare for a procedure that was never performed, the courts have recognized FCA cases under an inducement-of-fraud theory for off-label promotion. Under this theory, “[a]ny person who . . . knowingly . . . causes to be presented a false or fraudulent claim for payment or approval . . . is liable.” As applied to the case of off-label uses of pharmaceuticals, the false claim is the healthcare provider’s paperwork billing Medicare for a drug used in a non-reimbursable manner. While Medicare would be willing to reimburse for an on-label, medically acceptable use, it would not reimburse for an off-label, non-medically-acceptable use. Billing Medicare for the on-label use while using it for the non-medically-acceptable use would be considered a false claim, as Medicare would not have reimbursed had it known the truth about the drug’s usage with that particular patient.

The manufacturer is liable under this inducement theory because it is the arguable cause of the healthcare provider’s billing. The provider presents the false claim, but the off-label promotional efforts of the manufacturer cause the provider to do so. If the manufacturer had not told the provider about the alternative uses for the drug, then the provider would not have prescribed the drug for those alternative, non-medically-accepted indications.

The FCA has proven to be desirable in off-label-promotion cases due to the information problem described earlier; detection of off-label usage is difficult under the present system and there is little day-to-day government oversight of manufacturers’ representatives in the field. The FCA’s whistleblower provisions provide an incentive for those who have information about manufacturer’s behavior to come forward.

D. Off-label promotion cases under the FCA have been contentious for a number of reasons.

1. Off-label promotion does not fit the statutory purpose

First, there is the broader problem of the FCA’s original statutory purpose. The FCA originally targeted wartime-fraud cases in which the government paid for military supplies and received, for example, sawdust instead of gunpowder. It is clear that the government suffered loss in


such a transaction, and courts are most comfortable in assigning liability when the government receives nothing of value in exchange for payment.140

Many modern FCA cases, however, have addressed more difficult problems given the more complex regulatory and administrative state. Courts have been divided as to when civil FCA liability should attach, as it is unclear if any known regulatory violation makes a claim false or fraudulent.141 In Ab-Tech v. United States, for example, the court agreed with the defendant that the government had obtained the benefit of the contracted services, despite a regulatory violation.142 The defendant contractor constructed an automated data-processing facility in accordance with the government’s physical specifications, but it did not comply with the terms of the Small Business Act.143 The court upheld civil FCA liability, but rejected damages in that context. 144 The government paid $1.4 million to Ab-Tech and requested $4.2 million plus interest as treble damages, but the court found there were no damages to treble.145 The court noted that when “viewed strictly as a capital investment, the Government got essentially what it paid for.”146

Generally, courts have attempted to establish some limitations on behavior that could constitute a fraudulent or false claim under the FCA.147 The Second Circuit, for example, expressed discomfort in extending FCA liability in the healthcare context, noting that “the False

143. Id.
144. Id.
145. Id.
146. Id.
Claims Act was not designed for use as a blunt instrument to enforce compliance with all medical regulations.”148 Some courts have focused on whether defendants implicitly or explicitly certified compliance with regulations or contracts in determining whether or not there was a civil FCA violation.149

These limitations reflect a number of different concerns. One problem is that courts may be uncertain about whether harm results from the conduct; if the government feels it is acceptable for a physician to prescribe off-label, this behavior must not be very harmful or may actually be desirable.150 Under such conditions, it may be difficult or inappropriate to sanction a manufacturer.

Another problem is that courts feel that the FCA is punitive in nature.151 Unless the defendant has committed some wrong that is closer to malum in se, courts might feel that a technical regulatory violation does not deserve punishment and would be likely to label the violation as not material.152

On the other hand, if there is concern that sanctions are too great or improperly calculated, it is important to note that the prevalence of off-label promotion cases suggests that manufacturers are not deterred by FCA sanctions.153 In 2013, Pfizer had the distinction of settling its fifth case of off-label promotion since 2002.154 In one of its earlier cases, from 2009, Pfizer paid $2.3 billion to settle healthcare fraud charges arising from improper marketing activities relating to four drugs; it was the largest healthcare-fraud settlement in history at the time.155 Arguably, these

repeated settlements suggest that Pfizer may not be deterred by the present enforcement scheme.

2. Causality

A more specific challenge for FCA liability in the off-label-promotion context is causality. There is a long causal chain between the manufacturer’s promotional efforts and the improper reimbursement from Medicare.\(^{156}\)

The most proximate cause of harm to Medicare is the healthcare provider submitting reimbursement for a drug that has been prescribed for a non-CMS-approved indication.\(^{157}\) If CMS had known the truth about the indication, it would not have provided reimbursement for that drug prescription.

In comparison, the manufacturer’s promotional efforts’ role in causing the improper reimbursement is more attenuated and uncertain. Given the general availability of studies and drug compendia regarding off-label drug uses, it is entirely possible for providers to learn of off-label uses independent of the manufacturer’s paid representatives.\(^{158}\) To the extent that CMS approval does not correspond with the drug compendia recommendations, there is plenty of opportunity for providers to improperly bill Medicare without direct intervention by the manufacturer. Of course, off-label clinical studies may be funded by the manufacturer, but that is also a more attenuated causal inference, and those studies may also offer societal benefit.

Moreover, some commentators have argued that manufacturers should not be held liable because of a specificity problem.\(^{159}\) They suggest that a manufacturer should only be held liable if they have “specific knowledge of the falsity of the claim in question.”\(^{160}\) Thus, while manufacturers might have general knowledge that a number of claims are false, they do not know which specific claims are actually false.

Important for this article’s purposes, though, is that there is no falsity requirement for the manufacturer’s promotional efforts, because off-label promotion claims are under section 3729(a)(1)(A). Claims under section 3729(a)(1)(B) have a “double falsehood” requirement; the statute holds liable any person who “knowingly makes, uses, or causes to be made or


\(^{158}\) See, e.g., Robertson, *supra* note 73, at 550 (noting that the FDCA does not regulate non-manufacturers speech regarding off-label uses and that such independent speech may be more reliable).

\(^{159}\) See Hall & Berlin, *supra* note 156, at 673.

\(^{160}\) *Id.*
used, a false record or statement material to a false or fraudulent claim.”¹⁶¹ In contrast, there is no “double falsehood” requirement under section 3729(a)(1)(A).¹⁶² There is no need to allege that a false statement led to the false claim.¹⁶³ Moreover, these false claims may be filed by innocent third parties.¹⁶⁴

As some critics have noted, the role of the healthcare provider in prescribing a drug is certainly another cause of the eventual submission for off-label reimbursement.¹⁶⁵ The expertise of the healthcare provider may serve to cut causality here.¹⁶⁶ Moreover, critics relatedly argue that if manufacturers face liability for their attenuated causal role in driving the submission for reimbursement, many other parties might also face liability.¹⁶⁷ Nonetheless, the FCA’s broad definition of “knowing” seems to suggest that the attenuated-causality theory under off-label promotion is sufficient to establish liability.¹⁶⁸

E. Expanding FCA liability as a claw-back for excessive reimbursement

If we can tolerate the present concerns about FCA liability for off-label promotion, we can next consider whether expanding liability makes sense. Given that the present reimbursement system does not properly track indications, manufacturers will obtain improper profits through excessive off-label prescriptions and reimbursement. Thus, the remaining interim solution is to claw back those improper profits. Expanding civil FCA liability is the best immediate choice for detecting those problems and bringing back those profits.

1. The Proposal

This proposal suggests that courts hold manufacturers generally liable under the civil FCA for excess profits from improper Medicare Part D off-

¹⁶¹ 31 U.S.C. § 3729 (2012); See also United States ex rel. Franklin, 2003 WL 22048255 at *2-*3.


¹⁶⁵ See Hall & Berlin, supra note 156, at 673.

¹⁶⁶ See id. at 665.

¹⁶⁷ See id. at 673 (“If so, then any person including an independent physician, who discusses off-label uses would be liable under the FCA.”).

¹⁶⁸ See id. at 674 (proposing FCA liability scheme in which manufacturers “would have to have a specific intent to cause a specific treatment reimbursement submission . . . ”). The present FCA explicitly rejects a specific intent requirement. See 31 U.S.C. 3729 (b)(1)(B) (2012).
label reimbursement. Rather than looking at manufacturer’s promotional behavior, courts should view manufacturers as liable under the FCA because they are a cause of the improper reimbursement and they profit from such improper reimbursements. The goal is to pursue the theoretical solution in part IV by clawing back the excessive off-label drug profits from manufacturers.

This is a mild expansion of the existing off-label promotion doctrine. HHS can continue to rely upon whistleblowers to identify egregious off-label promotional behavior. Under this proposal, HHS would also rely upon whistleblowers to identify off-label prescription and the frequency of reimbursement for off-label prescription, independent of egregious manufacturer behavior.

Inferring that manufacturers are knowing, general cause of improper reimbursements is not a large step from the presently accepted inference that egregious manufacturer behavior causes improper reimbursements.169 This one step is sufficient to address cases of completely improper off-label prescriptions—prescriptions that should not be reimbursable at all under Medicare Part D. One example of a completely improper off-label prescription would be a prescription for a drug that does not match indications in any of the specified compendia.170 Another example would be if CMS has already considered the unapproved indication and explicitly rejected it for good reason. One good reason for rejecting the use of a drug for a particular indication would be the existence of sufficient scientific studies to evaluate effectiveness and safety for the unapproved condition that found the drug to be ineffective or unsafe for the unapproved condition.171 Manufacturers would be liable for the entire reimbursement amount for such non-reimbursable off-label prescriptions.

Perhaps the more challenging step is addressing the proper reimbursement rate for off-label prescriptions. The present system is binary; either a drug is reimbursable or it is not. As described in Part IV, I propose a more calibrated approach: implementing a cap on reimbursement rates. Courts or CMS would declare that any

169. See Hall & Berlin, supra note 156, at 673.

170. See SOCIAL SECURITY ACT OF 1935, Pub. L. 74-271, § 1927(g)(1)(B)(ii); Also, following existing case law, illegal kickbacks to physicians would also make such prescriptions sanctionable under the FCA. See, e.g., United States ex rel. Hutcheson v. Blackstone Medical, Inc., 647 F.3d 377, 379 (1st Cir. 2011).

171. This is the rough existing rule regarding Medicare Part A, which indicates that “[a]s long as the FDA has not specified such use as non-approved, coverage is determined taking into consideration the generally accepted medical practice in the community.” CENTERS FOR MEDICAID AND MEDICARE SERVICES, MEDICARE BENEFIT POLICY MANUAL § 1.30 (2014), available at https://www.cms.gov/Regulations-and-Guidance/Guidance/Manuals/Downloads/bp102c01.pdf; 42 U.S.C. § 1395x(t)(2)(B)(II) (2016).
reimbursement rate in excess of the capped competitive rate is subject to FCA liability.

This cap would only be triggered if the off-label condition has patent-protected, FDA-approved treatments available. If one such treatment exists and its cost is less than the price of the off-label drug, then the civil sanction should be the difference in rates.\textsuperscript{172} If there are multiple approved reimbursement rates, the relevant rate should be the lowest reimbursement rate or any of the other options presented earlier in Part IV.A.

If there are no approved treatments for the unapproved indication, then there are a number of possibilities. An aggressive move would be to cap all unapproved treatments at the same reimbursement level. This would level the playing field and reduce costs for Medicare. Here, however, it is unclear whether there is a strong need for intervention, given that no manufacturer has satisfied government standards for approval. The more cautious alternative is to disallow civil liability in this scenario. If there are no other treatments, approved or unapproved, for the unapproved indication, then there is no civil liability.

This strategy will incorporate proper incentives for manufacturers to bring the best products to market. If providers truly believe that a drug is effective for a condition that CMS has not approved condition and CMS has not explicitly rejected the drug, allowing limited reimbursement will provide an incentive for CMS to make a clear determination about the cost-effectiveness of the drug. Basing effective reimbursement on the lowest reimbursement rate will also help ensure that the manufacturer does not have a competitive advantage over competitors who have already obtained CMS approval for the same indication.

Additionally, establishing third-party restitution liability for manufacturers ensures that unapproved CMS reimbursements are not simply a windfall for manufacturers who produce an expensive drug. The fact that a manufacturer has not participated in improper off-label promotion efforts should not be an open door for it to benefit from improper physician-billing practices. Nonetheless, this article’s approach attempts to balance those revenues with the potential good that providers may be accomplishing.

Note also that this proposal provides for sanctions for off-label reimbursements even if those drugs eventually receive FDA approval for the off-label condition. Of course, once those drugs receive FDA approval for the off-label condition, there will be no further FCA liability. Not all off-label treatments will eventually receive FDA approval, and there may be a variety of reasons for such lack of approval. For drugs that do eventually receive approval for the off-label condition, though, note that those

Manufacturers are still civilly liable for excess profits obtained prior to FDA approval. While such manufacturers may enjoy high prices unconstrained by the FCA after they obtain FDA approval, that benefit does not extend to sales made prior to FDA approval for the off-label condition.

2. The statutory basis for the proposal

There are at least two potential bases for manufacturer liability under this proposal. I discuss the most commonly used basis first, section 3729(a)(1)(A). Present actions for off-label promotion typically proceed under this portion of the statute.\(^{173}\)

a) Section 3729(a)(1)(A): “Presentation of a false or fraudulent claim”

As discussed earlier, under section 3729(a)(1)(A), the manufacturer is liable because it is knowingly inducing healthcare providers to bill Medicare for prescription drugs that are not actually reimbursable due to their non-covered off-label usage. Important to note here is that purposeful behavior is not required; the fact that the manufacturer knows or acts in reckless disregard of the improper billing is sufficient.\(^{174}\) Given manufacturer involvement in researching and testing for off-label usage,\(^{175}\) it is difficult to believe that any manufacturer could claim ignorance of such billing.

The challenge, of course, is in the causal inference under this portion of the statute. Did the manufacturer cause the healthcare provider to improperly bill Medicare? I suggest that courts take a broad view of causation here rather than focusing on the manufacturer’s marketing behavior. If the manufacturer conducted or contributed to the research relating the drug to the off-label indication, this alone should be sufficient to demonstrate causation of improper billing.

Thus, courts should set aside their reluctance to impose liability, although their motivation to impose liability likely lies more in the damage done by improper billing and the instrumental usefulness of the FCA.

b) Section 3729(a)(1)(G): “Reverse False Claims”

The expanded version of the reverse-false-claims provision, section 3729(a)(1)(G), was introduced in 2009 as part of the Fraud Enforcement and Recovery Act of 2009 and has yet to generate substantial case law.\(^{176}\) Nonetheless, it provides an alternative route for liability. Section 3729(a)(1)(G) establishes liability for a person who knowingly conceals or knowingly and improperly avoids or decreases an obligation to pay or transmit money or property to the Government. The Affordable Care Act

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173. See SYLVIA, supra note 136.
176. See THE ELEMENTS OF A CAUSE OF ACTION UNDER THE FALSE CLAIMS ACT BEFORE AND AFTER FERA, 1 CIV. FALSE CLAIMS & QUI TAM ACTIONS (CCH)§ 2.01, 2015 WL 4602833 (2016).
(“ACA”) specifies that Medicare overpayments are obligations subject to liability under the FCA.\(^{177}\) Under the ACA, Medicare overpayments must be reported and returned within sixty days of identification; they are otherwise grounds for FCA liability.\(^{178}\)

Unlike section 3729(a)(1)(A), which looks at the cause of the excessive Part D reimbursements, section 3729(a)(1)(G) addresses parties who know of an obligation to pay the government. Given their role in researching off-label drug usage, manufacturers know that they improperly benefit from such off-label reimbursements.\(^{179}\) They therefore have an obligation to pay those excessive profits back.

The difficult part is establishing the element of “identification” of overpayments under this theory. First, as a matter of law, it must be established that reimbursement above the competitive cap for an off-label drug prescription is improper.\(^{180}\) Second, someone must identify such overpayments. This dovetails with the existing specificity problem under section 3729(a)(1)(A) as discussed earlier in section V.C.2. Does “identification” correspond with specific knowledge of a particular claim’s falsity? Such knowledge seems difficult to come by.

As a practical matter, this prong is most likely useful if statistical sampling of aggregate prescription rates with aggregate diagnosis rates is sufficient to establish liability. Given manufacturer’s research into indications for their products, they should know—or at least be aware of—the risk of off-label usage and how much they might benefit. To the extent that they benefit improperly from off-label usage billed to CMS, the reverse-false-claims provision can establish civil FCA liability.

3. Why this proposal works

a) Relators provide the link between indication and prescription.

The FCA allows private litigants to pursue actions against manufacturers in the form of \textit{qui tam} lawsuits. Without extensive overhaul of the present prescription and reimbursement system, it is very costly for CMS and its delegates to determine the eligibility of drugs for reimbursement under Part D, and investing in such improved data acquisition does not appear to be a present priority for CMS.\(^{181}\) Investigating individual doctors and providers is costly. Manufacturers and insiders are best positioned to observe off-label drug issues and to bring them to light.

\(^{177}\) See \textit{Affordable Care Act}, Pub. L. No. 111-148, § 1128I (d) [hereinafter ACA].

\(^{178}\) ACA §1128I(d)(1)-(2).


\(^{181}\) See OIG \textit{Part D Reimbursement Report}, \textit{supra} note 6, at 4; See also Weber et al. \textit{supra} note 7.
Relators can provide a variety of information to make this system work. The present doctrine of off-label promotion has relators focused upon manufacturers’ promotional behavior. The increased scope of manufacturer liability under this proposal allows relators with different types of information to come forward. One possibility is relators who have direct information linking prescriptions together with indications; this would be the most direct linkage between reimbursements and the use of the drug. Another possibility would simply be evidence that manufacturers know of the aggregate rates of off-label prescriptions and reimbursements.

Additionally, this form of litigation provides payment for the cost of detection. The FCA provides for attorneys’ fees, which helps compensate for the role of attorneys in detecting offenses. The percentage bounty for relators similarly compensates for their efforts in uncovering off-label drug usage and reimbursement.

An alternative to FCA liability would be litigation under a theory of unjust enrichment. Civil liability for unjust enrichment incorporates the possibility that the defendant did no wrong but was simply the unknowing recipient of unjust gains. As noted above, though, this form of litigation requires information regarding diagnosis, so this cause of action would be extremely difficult without whistleblower support.

b) Manufacturers are the ones who profit.

Liability under the FCA is fair, given the allocation of revenue from the off-label reimbursement. Unless there are kickbacks or other improper incentives at play, the provider is not a direct beneficiary of the improper billing, except to the extent that the provider generates goodwill and business from the patient. The primary beneficiaries are the patient who receives the drug and the manufacturer. The patient’s benefit is from improved health, which can be difficult to quantify. Moreover, it is politically difficult to go after individual patients who may attract sympathy and may not be aware of the off-label nature of their treatment. As a practical matter, the manufacturer accrues benefit and thus is in a position to pay civilly.

VI. Concerns

A. Stigma and signaling

Expanding FCA liability for manufacturers and off-label drug reimbursement may raise the problem of excessive stigmatic harm by lumping defendants with varying levels of moral culpability together. A healthcare provider committing fraud by collecting Medicare payments for


183. See RESTATEMENT (THIRD) OF RESTITUTION AND UNJUST ENRICHMENT § 41 (AM. LAW INST. 2011).
which no service was provided could be liable under the FCA, as could a manufacturer engaging in truthful off-label promotion of a useful pharmaceutical product.

I suggest that on the balance, though, there is actually insufficient stigmatic sanction under the FCA. As stated in that statute, the FCA addresses false or fraudulent claims. Falsity is a less morally laden description than fraud.\textsuperscript{184} Only certain portions of the FCA actually require fraudulent intent.\textsuperscript{185} Thus, for a defendant committing outright fraud through non-delivery of service, there is probably insufficient stigmatic harm in FCA liability.

Nonetheless, to the extent that there may be excessive stigmatic harm through the aggregation of fraud and falsity in the statute, the DOJ could seek to alleviate this harm by emphasizing the falsity aspect in press releases.

As noted above, another solution would be to pursue civil remedies under the doctrine of unjust enrichment. Civil liability for unjust enrichment incorporates the possibility that the defendant did no wrong but was simply the unknowing recipient of unjust gains.\textsuperscript{186} The combination of litigation and the “unjust” portion of the label may have sufficient stigmatic power against the defendant for those reading the popular press.

As a practical concern, the stigmatic and signaling effects of FCA liability may impact manufacturer behavior. They may further discourage manufacturer investment in off-label drug usage. Note that this proposal deliberately attempts to reduce manufacturer over-investment in off-label drug usage. It is possible that combining the reduced reimbursement rates with the stigma of FCA liability may overly reduce manufacturer investment. As noted in Part IV, the harms from off-label usage may be higher than assumed in Part III’s model, so such increased reduction may actually be desirable.

Stigma could also affect healthcare providers. While this article does not propose litigation against healthcare providers, it is entirely possible that providers would learn about litigation against particular manufacturers and their drugs. A number of problems might result from such knowledge. One might be that healthcare providers might simply be more reluctant to prescribe a manufacturer’s drugs because they interpreted the litigation news as generalized wrongdoing. Professional norms would hopefully prevent healthcare providers from drawing strong negative inferences in such cases. Rather, if they really believed that a drug was particularly risky as a result of hearing of manufacturer litigation,

\textsuperscript{184} Fraud incorporates the intent to deceive for the purposes of causing some loss. See, \textit{e.g.}, United States v. Hawkey, 148 F.3d 920, 924 (8th Cir. 1998).


\textsuperscript{186} See \textit{Restatement (Third) of Restitution and Unjust Enrichment} § 41 (\textit{Am. Law Inst. 2011}).
they would directly investigate the scientific studies concerning the drug. Nonetheless, the stigmatic effects might still subtly reduce healthcare providers’ prescriptions of certain drugs, even for on-label conditions and might reduce provider reliance upon manufacturer-sponsored information. Such reduced reliance might be in society’s interest, depending on society’s beliefs concerning the value of manufacturer-sponsored information.

Healthcare providers might also feel the threat of litigation, even if whistleblowers and assistant U.S. attorneys do not target them directly. The theory of liability under this proposal emphasizes cause-in-fact; the providers likely have at least comparable levels of causal responsibility for off-label drug reimbursements. While providers do not profit in the same way that manufacturers do, they may be concerned about being subject to at least the threat of civil liability. The risk of such fears may be assuaged by continued statements from the FDA and HHS that they do not intend to regulate the practice of medicine.

B. Manufacturers still excessively profit from off-label reimbursement.

This claw-back proposal under the FCA is not a panacea; manufacturers will still profit from off-label reimbursements in a variety of ways.

1. Manufacturers with no knowledge

First, a manufacturer could avoid FCA liability if it had no knowledge of the off-label reimbursements. The FCA diverges from an ideal solution for these profits in that it contains a mens rea component. In theory, the optimal solution would be strict liability for windfall profits; the unintended over-reimbursement by the government for off-label prescriptions is a real loss, regardless of the manufacturer’s subjective awareness of those windfall profits. Nonetheless, given the incentive for manufacturers to study and track the effectiveness and reach of their products, it seems unlikely that manufacturers would be unaware of the general practice of off-label reimbursement.

It is technically possible that a manufacturer might not only have no knowledge of the off-label usage, but it might also have not contributed at all to the research leading to the discovery of the off-label indication. The ideal claw-back solution would need a relatively broad causal theory to claim that manufacturers were liable for the improper off-label drug reimbursements.

Pushing the law to this point may be desirable from a claw-back perspective, but such precedent might cause difficulties in other areas of the FCA. The preferable long-term solution is actual statutory reform of the reimbursement system, rather than acceptance of the more attenuated causality.
2. Profits from private insurance

The improper windfall profits are the result of not only government payments; they may also be the result of private-insurer payments. Implementing this FCA-based claw-back system in the interim will not completely solve the inequitable distribution of profits stemming from off-label reimbursement. To the extent that private health insurance companies also face disparities regarding off-label reimbursement, there may still be excessive expenditures and improper incentives. Rather than the government bearing the costs of such excessive expenditures, though, private health insurance companies likely pass along such excessive costs to their insured.

While excessive costs for private health-insurance companies are not the focus of this paper, it is important to acknowledge that those companies may face challenges that parallel the government’s challenges with off-label drug prescriptions. Some of the systematic changes proposed herein linking prescriptions to indications may similarly help private insurance companies address this shared problem with excessive healthcare costs. Private insurance companies may be useful allies in obtaining reforms of the prescription and reimbursement process; a fair discussion of the potential interaction between the private market for off-label drugs and the government-led marketplace would require a separate paper.

VII. Conclusion

High drug prices are important in motivating pharmaceutical manufacturers to bring safe and effective products to market. At the same time, Medicare should not incur excessive drug costs by paying top dollar for drugs that have not been proven effective for treatment. Unfortunately, Medicare Part D is prone to excessive drug prices due to the practice of off-label drug prescription. The present systemic failure to link indication with reimbursement in the Medicare Part D regime encourages excessive prescription-drug costs from even well-intentioned manufacturers. Moreover, these flawed incentives also increase the risk of spillover effects for non-Medicare patients. Because manufacturers will over-invest in off-label drug usage, even non-Medicare patients may face increased exposure to expensive and ineffective off-label drug usage. This article proposes a theoretically superior Part D reimbursement system that allows for the development of an optimal level of off-label drug usage. In the long-term, such a reimbursement system will allow Medicare beneficiaries to obtain a variety of drugs for treatment while ensuring that Medicare is not paying higher prices for unproven drugs.

The long-term solution requires substantial systemic and regulatory reforms that are not immediately likely. In the short term, civil enforcement through the civil False Claims Act can serve as an interim tool to limit excessive Part D off-label drug costs. Rather than emphasizing
punishment of wrongful behavior, the DOJ can leverage whistleblowers under the FCA to focus on facilitating fair reimbursement for off-label drug prescriptions. Even well-intentioned manufacturers can obtain windfall profits from off-label drug reimbursement. Off-label drug usage may be safe, effective, and desirable for some patients, but those benefits do not automatically justify windfall profits. Litigation under the FCA can claw back the excessive profits and correct the unfairness and improper incentives resulting from the present Medicare Part D reimbursement system.