Affordability and availability of off-patent drugs in the United States—the case for importing from abroad: observational study

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ABSTRACT

OBJECTIVES
To evaluate whether off-patent prescription drugs at risk of sudden price increases or shortages in the United States are available from independent manufacturers approved in other well regulated settings around the world.

DESIGN
Observational study.

SETTING
Off-patent drugs in the USA and approved by the Food and Drug Administration, up to 10 April 2017.

STUDY COHORT
Novel tablet or capsule prescription drugs approved by the FDA since 1999 that were no longer protected by patents or other market exclusivity and had up to three generic versions.

MAIN OUTCOME MEASURES
Number of additional manufacturers that had obtained approval from any of seven non-US regulators with similar standards (European Medicines Agency (European Union), HealthCanada (Canada), Therapeutic Goods Association (Australia), Medsafe (New Zealand), Swissmedic (Switzerland), Medicines Control Council (South Africa), and the Israel Health Ministry). Association with drug characteristics including US orphan drug designation for drugs treating rare diseases, World Health Organization (WHO) designated essential medicines; only 12 (11%) were complex products that might be more complicated to manufacture, and total Medicaid spending in 2015.

RESULTS
Of 170 eligible study drugs, more than half (109, 64%) had at least one manufacturer approved by a non-US regulator and 32 (19%) had four or more. Among 44 (26%) drugs with no FDA approved generic versions, 21 (68%) were available from at least one manufacturer approved by one of the seven non-US regulators, and two (5%) by four or more manufacturers. Across all drugs and regulators (including the FDA), 66 (39%) drugs were available from four or more total manufacturers. Of 109 drugs with at least one non-US regulator approved manufacturer, 12 (11%) were approved for patients with rare diseases and 29 (27%) were WHO designated essential medicines; only 12 (11%) were complex products that might be more complicated to import. The highest numbers of drugs were indicated for treating cardiovascular diseases, diabetes, or hyperlipidemia (19, 17%); psychiatric disease (16, 15%); and infectious diseases (15, 14%). In 2015, Medicaid alone spent near US$700m (£508m; €570m) on generic drugs without adequate US competition that could have had a manufacturer approved by non-US peer regulatory agencies.

CONCLUSION
In this study, more than half the off-patent drugs with no generic competition in the USA had at least one independent manufacturer approved by a non-US peer regulatory agency; slightly fewer than half had four or more total manufacturers. Facilitating US patient access to such manufacturers could help sustain affordable access to essential off-patent drugs.

Introduction
In recent years, shortages and dramatic price increases have been observed among some generic drugs in the United States.1,3 The value of generic medicines lies in the fact that, when effectively overseen by drug regulators, these drug treatments are widely used and less expensive while still being equally safe and efficacious alternatives to brand name drugs that are no longer protected by patents.4,7 In 2016, generic drugs constituted 89% of dispensed medicines in the USA and only 27% of overall drug spending.8 Appropriate use of generic drugs increases patient adherence,4,9 improves health outcomes,10 and generates substantial savings for US taxpayers and patients.11 Generic drug treatments are inexpensive partly because of competition among manufacturers with regulatory approval to market the drugs.12 Broad competition from multiple independent competitors exists for most off-patent drugs that are in large markets and easy to manufacture, driving down prices. But in recent years, a problem has emerged with some off-patent drugs that are no longer protected by patents, making these products susceptible to shortages and dramatic price increases.13 As more time passes after expiration of a brand name drug’s market exclusivity,
more companies complete the regulatory process to sell their versions of that drug in the US market and offer discounts to gain market share, which eventually drives the price of that drug closer to its marginal cost. At that point, any manufacturers of the drug will only continue to make that product if the cost of its production remains low or if there are strong synergies with that firm’s other product lines. In addition, with profit margins so thin, prospective competitors have little incentive to incur the start-up costs and time required to gain regulatory approval and manufacture that drug. This dynamic is even more acute with drugs used in only small patient populations, since the potential revenues may have already been modest at the outset.14

Between 2007 and 2012, the number of drugs with shortages in the USA tripled from 154 to 456 drugs.15 The problem has been more acute among parenteral drugs, which can be more expensive to manufacture, but oral drug treatments such as abendazole, bumetanide, and ethambutol have also faced shortages in recent years.16 Increasing consolidation of generic manufacturers,17 wait times for regulatory approval,18 and small markets19 are some factors that have contributed to limited generic competition and ensuing shortages. Increasingly, companies have noted this trend and have begun purchasing the rights to manufacture older drugs with few or no competitors and substantially raising their prices. Most notably, Turing Pharmaceuticals purchased the rights to pyrithamine (Daraprim), a decades old treatment for toxoplasmosis, and raised its price overnight from US$13 to $750 per tablet.19 A 2016 General Accountability Office report identified more than 300 generic drugs sold in the USA that had price increases of 100% or more from 2010 to 2015, many of which were older, small market drugs.20

One strategy to counteract these shortages and price increases would be to rapidly increase the number of competitors by expediting Food and Drug Administration (FDA) approval for US marketing of manufacturers of the same drugs already approved by peer drug regulatory bodies, such as the European Medicines Agency (EMA) and Health Canada. It is likely to be faster and more efficient for current manufacturers of an already approved drug to expand production to serve the US market than for a manufacturer to newly enter the market by obtaining FDA approval and building new manufacturing capabilities. Such a system would also allow patients in the USA to access drugs manufactured by previously non-FDA approved companies that may have otherwise been deterred by the cost of an additional approval from the FDA. Thus, the possibility of international competition for US generic drugs would reduce the risk and duration of shortages, and diminish the ability of manufacturers to impose large price increases of generic drugs with limited US competition.

Imported drugs are already widely used in the USA. Drugs manufactured outside the USA represent about a quarter of the US pharmaceutical market, totaling US$86bn (£62.5bn; €70bn) in 2015.21 According to the FDA, 80% of active pharmaceutical ingredients and 60% of finished drugs are manufactured abroad.22 However, in most cases, these imported drugs and ingredients are subject to the same FDA oversight as domestic products manufactured in accordance with good manufacturing practices and pursuant to an FDA approval, which includes meeting packaging, labeling, and dosage requirements. There are a few notable exceptions. A modest proportion of US citizens travel to Canada and Mexico to purchase lower priced prescription drugs.23 The US Food, Drug, and Cosmetics Act officially prohibits such self-importation, but the FDA allows for personal use of non-FDA approved drugs in certain cases, such as when the treatment is unavailable domestically or a person has initiated treatment abroad with a non-FDA approved drug.24 The FDA has also facilitated temporary importation of drugs approved only by peer drug regulatory bodies when a shortage has occurred of an essential medicine that could not be resolved by manufacturers of FDA approved drugs.25 Some experts have also suggested use of temporary importation to deal with substantial price increases of US generic drugs.26

One way that the FDA could expedite the importation of generic drug treatments already approved abroad is through a variation of reciprocal approval.27,28 For example, the FDA could still issue its own approval of each drug (as required under US law), but it would be issuing that approval on the basis of evidence of the drug’s previous approval by another stringent national regulatory authority. FDA could limit the use of this reciprocal drug approval process to countries with stringent national regulatory authorities and strong safety records, and which the FDA has assessed as having equivalent generic drug approval processes. The few areas in which there remain significant differences among comparable regulators, such as approval of generic narrow therapeutic index drugs, could be excluded from the reciprocal process. The reciprocal approval could also be made contingent on the manufacturers starting the process for standard FDA approval.

This approach is consistent with US law and practice.29,31 The 2012 FDA Safety and Innovation Act gave the FDA authority to enter into agreements to recognize drug inspections conducted by foreign regulatory authorities if the FDA determined that those authorities are capable of conducting inspections that meet US requirements.32 Pursuant to that authority, the FDA and EMA concluded an agreement on mutual recognition of inspection reports, which was added as an amendment to the existing 1998 US-EU Mutual Recognition Agreement.33 Limiting the use of reciprocal approval to generic versions of drugs already approved and long used in the USA minimizes the safety risks that might otherwise arise from relying on the approval of a novel medication by another regulatory authority. To justify the resources required to establish and maintain a reciprocal approval pathway, it must be useful. This pathway would work only if there are
sufficient numbers of generic manufacturers approved to supply drugs outside the USA and that could be called on to export their products. Thus, we assessed manufacturer approvals from non-US peer regulators of drugs approved by the FDA since 1939 that are no longer protected by patents, but lack adequate generic competition. We also examined specific drug characteristics that describe the potential importance of reciprocal drug approval.

**Methods**

**Sample construction**

We used the Drugs@FDA database to identify novel drugs in tablet or capsule form approved by the FDA since 1939, the first full year after the agency began requiring safety information for drug approval. We excluded tentative approvals, biological treatments, over-the-counter drugs, and duplicate listings. If there were multiple currently marketed dosage strengths of a novel drug, we randomly selected one dosage strength for inclusion. If a novel drug was discontinued but had at least one active generic version available, it was included in the sample. We excluded discontinued novel drugs for which there were also no active generic versions, because we assumed that the drug was no longer used. We excluded drugs with current patent protections or other market exclusivity, identified from the Orange Book in April 2017. We focused on drugs in tablet or capsule formulation partly because injections and infused drugs are predominantly purchased by hospitals and physician facilities. Finally, we excluded one drug because it had only one approved generic version that was protected from generic competitors by 180 day exclusivity at the time of data extraction.

For each drug in our final cohort, we used Drugs@FDA to identify FDA approved bioequivalent generic drugs by matching active ingredient and formulation. We excluded discontinued generic drugs, tentative approvals, and duplicate applications filed by the same generic manufacturer. We disregarded multiple dosage strengths and only counted the generic drug once.

We determined the number of different generic manufacturers that had received FDA approval for each novel drug and only included those drugs with up to three approved generic versions (web appendix shows full list of drugs included). A previous study of generic competition used three manufacturers as the threshold for considering a marketplace that could lack sufficient competition to maintain low prices. The FDA subsequently issued a policy with a similar threshold, prioritizing review of drugs without existing patents or exclusivity and fewer than three approved generic versions.

**Characteristics of drugs lacking generic competition in the USA**

Firstly, we determined whether each FDA approved drug was designated under the US Orphan Drug Act of 1983, for the indication for which it was initially FDA approved. The Orphan Drug Act status signifies a drug indicated for a rare disease affecting fewer than 200,000 people in the USA. Secondly, we identified whether each drug in our sample was included in the most recent World Health Organization’s Model List of Essential Medicines, which includes drugs considered fundamental for any healthcare system. One drug, ferric hexacyanoferrate (Prussian blue), was included in the WHO list as a powder but approved by the FDA as a capsule.

Thirdly, using WHO’s Anatomic Therapeutic Classification system, we categorized all drugs in the sample into one of nine treatment classes: autoimmune or musculoskeletal; cancer; cardiovascular, diabetes, or hyperlipidemia; gastrointestinal; genitourinary; infectious disease; neurology; psychiatry; and other. If a drug was listed in more than one treatment class, we selected the class within which the drug was initially approved by the FDA.

We also determined the complexity of drugs lacking in US generic competition. A drug is considered a complex product if specific attributes make it difficult to manufacture the drug or establish bioequivalence, such as complex active ingredients (eg, peptides, complex mixtures, naturally sourced) or complex formulations (eg, colloids or liposomes). One author (RG) reviewed the chemical structure of each novel drug in our cohort to determine whether it was a complex product. In addition, a list of pharmaceutical related citizen petitions and https://www.regulations.gov were used to identify any citizen petitions that were filed for issues of bioequivalence or generic interchangeability. These drugs were also considered complex. Drugs with unclear status were independently classified by a second author (ASK) and disagreements resolved through consensus (n=2).

Finally, we used the Medicaid state drug use database to determine spending by that program in 2015 for each drug in our sample. These data provide insights into US spending on generic drugs without adequate competition and the drugs’ market sizes. We focused on Medicaid spending because the data source separates spending by different formulations of each drug, while data sources for Medicare Part D spending do not. Medicaid is a federal and state funded health insurance program for people on low incomes that represents about 20% of US prescription drug spending and serves 70 million patients. This data source includes every US state’s data for outpatient drugs covered under the Medicaid Drug Rebate Program, but does not incorporate rebates. For each drug, we matched the National Drug Code (NDC) number with the generic drug name, and totaled spending across all FDA approved brand name and generic versions of the identified formulation.

**Availability of drug competition from outside the USA**

For all drugs in our sample, we obtained data from seven non-US peer drug regulatory agencies: EMA (European Union), HealthCanada (Canada), Therapeutic Goods Association (Australia), Medsafe (New Zealand), Swissmedic (Switzerland), Medicines Control Council
(South Africa), and the Israel Health Ministry. These regulatory bodies were selected because they were included in a list of high quality regulatory authorities in a recent US bill that proposed to create a system of international drug reciprocity, and their publicly accessible databases offered information in English.

Two authors (RG and MC) independently cataloged manufacturers with approval from each of these seven regulators. Manufacturers were counted if their drug shared its active ingredient and formulation with the FDA approved drug. Through searches of company websites and other internet sources, the reviewers identified the manufacturers approved by each regulatory body that were different from the FDA approved manufacturers, accounting for mergers, acquisitions, and transfers of drug licenses between manufacturers. If the manufacturers were the same as those approved in the USA, they were excluded. Disagreements were resolved through consensus (n=3).

For each drug, we determined the number of manufacturers that were approved by the seven non-US drug regulators but not approved by the FDA. We also identified each drug the total number of manufacturers approved across all regulators, including the FDA and seven non-US regulators. We converted these results into ordinal variables (0, 1, 2, 3, and ≥4 manufacturers). We then determined the proportion of drugs with at least one manufacturer and those with four or more manufacturers, because a previous study determined that four manufacturers ensured sufficient competition to prevent substantial price increases. We did this assessment for the seven non-US regulators and for all eight regulators, including the FDA.

Statistical analysis
We used χ² tests to compare the likelihood of having at least one non-US regulator approved manufacturer based on four novel drug characteristics: US orphan status designation, WHO essential medicine designation, treatment class, and complexity. We used the Wilcoxon-Mann-Whitney test to compare the difference in 2015 Medicaid spending between drugs with and without at least one non-US regulator approved manufacturer of FDA approved generic drugs. Among these same five characteristics, we repeated the analyses to compare the likelihood of having four or more different manufacturers across all regulators, including the FDA and non-US regulators. We used Microsoft Excel 2016 (Microsoft Corp) and Stata 12.0 (StatCorp) to conduct all analyses.

Patient involvement
No patients were involved in setting the research question or the outcome measures; nor were they involved in the design and implementation of the study. We have no plans to involve patients in dissemination.

Results
We found 170 novel drugs in tablet or capsule formulation approved by the FDA since 1939 that were off-patent and had up to three FDA approved generic versions, as of 10 April 2017 (fig 1). Three quarters of drugs in our cohort (126, 74%) had at least one FDA approved generic version (table 1). Seventy (41%) were approved after 1983, of which 23 (33%) were designated as having orphan status. About one fifth (38, 22%) were considered essential medicines by WHO. Drugs indicated for the treatment of cardiovascular disease, diabetes, or hyperlipidemia were most common (33, 19%), followed by those for infectious disease (22, 13%) and psychiatric disease (22, 13%). A small number of drugs (22, 13%) were complex products. Among the 151 drugs covered by Medicaid, the median 2015 Medicaid spending was US$1m (interquartile range 0.2m-4.3m; table 1); total spending was $1bn, which accounted for about 2% of total Medicaid spending in 2015 ($57bn).

Potential competitor manufacturers of off-patent US drugs, approved by non-US regulators
More than half the drugs (109, 64%) had at least one manufacturer with a version approved by a non-US drug regulator, while about a fifth (32, 19%) had four or more manufacturers. Of 44 (26%) drugs in our sample with no generic versions approved by the FDA, 21 (48%) had at least one manufacturer with a version approved by a non-US drug regulator (fig 2), and 19 (43%) drugs were produced by one to three non-US manufacturers. For example, albendazole, a broad spectrum antiparasitic drug, had no generic versions approved by the FDA but three versions approved by non-US drug regulators. Nitisinone, which treats hereditary tyrosinemia type 1, and nabilone, which treats chemotherapy induced nausea and vomiting, were the only drugs with no generic versions approved by the FDA and four or more versions approved by non-US regulators.

Of 170 drugs with up to three FDA approved generic versions, 147 (86%) had at least one approved manufacturer across both FDA and non-US regulators. Furthermore, 66 (39%) could reach the threshold of four or more approved manufacturers if the FDA permitted reciprocal approval of drugs approved by any of the seven non-US regulators. For instance, cefixime, a commonly used antibiotic, had one FDA approved version and seven non-US approved versions.

**Table 1**

<table>
<thead>
<tr>
<th>Drug Characteristics</th>
<th>Number of Drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>170</td>
</tr>
<tr>
<td>FDA approved</td>
<td>170</td>
</tr>
<tr>
<td>FDA generic</td>
<td>152</td>
</tr>
<tr>
<td>Non-US regulator</td>
<td>126</td>
</tr>
<tr>
<td>Four or more</td>
<td>23 (33%)</td>
</tr>
<tr>
<td>&gt; Four</td>
<td>77 (45%)</td>
</tr>
<tr>
<td>Highest number</td>
<td>11 of 4</td>
</tr>
</tbody>
</table>

**Fig 1** Study sample definition flowchart

New drug applications listed in Drugs@FDA (n=8906)

Excluded (n=8736):
- Biological treatments (n=202)
- Tentative approvals (n=106)
- Over the counter (n=242)
- Non-novel drugs (n=5735)
- Non-tablets or non-capsules (n=966)
- Duplicate drugs (n=107)
- Multiple strengths (n=949)
- Drugs with existing patents and/or market exclusivity (n=136)
- All versions (brand name and generic) discontinued (n=83)
- Drugs with ≥4 FDA approved generic versions (n=210)

Novel drugs with 0–3 generic versions (n=170)
approved generic version but a total of four versions when including those approved by non-US regulators. Similarly, diltiazem, a calcium channel blocker used to treat cardiovascular conditions such as angina and hypertension, had two FDA approved generic versions but 10 approved versions across all regulators.

Factors associated with additional competition from non-US regulator approved manufacturers
Among drugs with up to three generic versions approved by the FDA, the number of manufacturers approved by non-US regulators did not vary significantly by drug characteristics (table 2). However, of the 109 drugs with at least one manufacturer approved by a non-US regulator, 12 (11%) were approved under the 1983 Orphan Drug Act for patients with rare diseases and 29 (27%) were WHO designated essential medicines, suggesting that reciprocal drug approval could be useful for some vulnerable populations. The highest numbers of drugs were indicated for the treatment of cardiovascular diseases, diabetes, or hyperlipidemia (19, 17%); psychiatric disease (16, 15%); and infectious diseases (15, 14%). Only 12 (11%) were complex products, which may need product specific testing. The total 2015 Medicaid spending for the 109 drugs with at least one manufacturer approved by a non-US regulator manufacturer was nearly US$700m.

Of the 66 drugs with four or more manufacturers across all eight regulators, including the FDA and non-US regulators, eight (12%) were designated to have orphan status and treat rare diseases and 18 (27%) were WHO designated essential medicines, suggesting that reciprocal drug approval could be useful for some vulnerable populations. The highest numbers of drugs were indicated for the treatment of cardiovascular diseases, diabetes, or hyperlipidemia (19, 17%); psychiatric disease (16, 15%); and infectious diseases (15, 14%). Only 12 (11%) were complex products, which may need product specific testing. The total 2015 Medicaid spending for the 109 drugs with at least one manufacturer approved by a non-US regulator manufacturer was nearly US$700m.

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Discussion
Principal findings
International sources could improve the supply and increase the competition for off-patent drugs in the USA with insufficient competition, about two thirds of

RESEARCH

Table 1 | Characteristics of 170 novel off-patent drugs approved by the Food and Drug Administration (FDA) and lacking generic competition in the USA

<table>
<thead>
<tr>
<th>Drug characteristic</th>
<th>FDA approved drugs*</th>
<th>0 generic versions (n=44)</th>
<th>1 generic version (n=47)</th>
<th>2 generic versions (n=41)</th>
<th>3 generic versions (n=38)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Approved for rare disease</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orphan status designation</td>
<td>23 (33)</td>
<td>12 (60)</td>
<td>4 (22)</td>
<td>1 (8)</td>
<td>6 (32)</td>
</tr>
<tr>
<td>No designation</td>
<td>47 (67)</td>
<td>8 (40)</td>
<td>14 (78)</td>
<td>12 (92)</td>
<td>13 (68)</td>
</tr>
<tr>
<td>WHO Essential Medicine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Included</td>
<td>38 (22)</td>
<td>14 (32)</td>
<td>8 (17)</td>
<td>6 (15)</td>
<td>10 (26)</td>
</tr>
<tr>
<td>Not included</td>
<td>132 (78)</td>
<td>30 (68)</td>
<td>39 (83)</td>
<td>35 (85)</td>
<td>28 (74)</td>
</tr>
<tr>
<td>Treatment class</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Autoimmune or musculoskeletal</td>
<td>13 (8)</td>
<td>5 (11)</td>
<td>2 (4)</td>
<td>2 (5)</td>
<td>4 (11)</td>
</tr>
<tr>
<td>Cancer</td>
<td>16 (9)</td>
<td>9 (20)</td>
<td>3 (6)</td>
<td>1 (2)</td>
<td>3 (8)</td>
</tr>
<tr>
<td>Cardiovascular, diabetes, or hyperlipidemia</td>
<td>33 (19)</td>
<td>3 (7)</td>
<td>10 (21)</td>
<td>11 (27)</td>
<td>9 (24)</td>
</tr>
<tr>
<td>Gastrointestinal</td>
<td>19 (11)</td>
<td>7 (16)</td>
<td>4 (9)</td>
<td>6 (15)</td>
<td>2 (5)</td>
</tr>
<tr>
<td>Genitourinary</td>
<td>12 (7)</td>
<td>4 (9)</td>
<td>4 (9)</td>
<td>3 (7)</td>
<td>1 (3)</td>
</tr>
<tr>
<td>Infectious disease</td>
<td>22 (13)</td>
<td>8 (18)</td>
<td>7 (15)</td>
<td>3 (7)</td>
<td>4 (11)</td>
</tr>
<tr>
<td>Neurology</td>
<td>16 (9)</td>
<td>2 (5)</td>
<td>3 (6)</td>
<td>4 (10)</td>
<td>2 (8)</td>
</tr>
<tr>
<td>Psychiatry</td>
<td>22 (13)</td>
<td>1 (2)</td>
<td>8 (17)</td>
<td>7 (17)</td>
<td>6 (16)</td>
</tr>
<tr>
<td>Other</td>
<td>17 (10)</td>
<td>5 (11)</td>
<td>6 (13)</td>
<td>4 (10)</td>
<td>2 (5)</td>
</tr>
<tr>
<td>Product complexity‡</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Complex</td>
<td>22 (13)</td>
<td>8 (19)</td>
<td>5 (11)</td>
<td>5 (12)</td>
<td>4 (11)</td>
</tr>
<tr>
<td>Non-complex</td>
<td>148 (87)</td>
<td>36 (81)</td>
<td>42 (89)</td>
<td>36 (88)</td>
<td>34 (90)</td>
</tr>
<tr>
<td>Median (IQR) Medicaid spending in 2015 (USD)¶</td>
<td>1 038 995 (229 052-4 307 314)</td>
<td>691 207 5 (229 052-9 469 889)</td>
<td>322 175 (97 865-2 601 182)</td>
<td>1 219 333 (272 251 5-3 152 809)</td>
<td>1 925 303 (649 931-5 261 405)</td>
</tr>
</tbody>
</table>

Data are number (%) of drugs unless stated otherwise. IQR=intertquartile range, WHO=World Health Organization.
*Number of generic versions approved by the FDA for each brand name drug in the sample.
†Orphan status designation (attributed to drugs that treat rare diseases) began in 1983, so the number of orphan versus non-orphan status designated drugs differs from the total number of drugs.
‡A drug is considered a complex product if specific attributes make it difficult to manufacture the drug or establish bioequivalence, such as complex active ingredients (eg, peptides, complex mixtures, naturally sourced) or complex formulations (eg, colloids or liposomes).
¶The Medicaid state drug utilization database was used to determine spending by that program in 2015 for each drug in the study sample.
which had at least one additional manufacturer with a version approved by a non-US regulator. Including the seven non-US peer regulators, nearly 40% of the drugs in our analysis could be served by four or more manufacturers, helping ensure low prices and avoid unexpected shortages. Most of those drugs treat infectious diseases; neurological and psychiatric diseases; or cardiovascular diseases, diabetes, or hyperlipidemia—some of the most prevalent and preventable medical conditions affecting US patients. In addition, many of the drugs in our sample were WHO designated essential medicines or intended for patients with rare diseases.

Policy implications
Several legislative attempts have been made to allow drug imports into the USA, particularly from Canada. For example, the Maine Pharmacy Act was passed in 2013 to allow direct importation of prescription drugs for personal use from pharmacies in Canada, the United Kingdom, New Zealand, and Australia, but was later ruled unconstitutional under the premise that importation is a federal issue. The premise that importation is a federal issue. The 2003 Medicare Modernization Act allowed for drug importation with the certification of the Secretary of Health and Human Services, but it has never been implemented. More recently, Senators Bernie Sanders (Independent, Vermont) and Amy Klobuchar (Democrat, Minnesota) proposed allowing importation from Canada, but it was voted down.

Drug industry advocates and even previous FDA commissioners have argued that importing prescription drugs that were approved outside the US could compromise, or at least raise concerns for, patient safety. Some safety concerns include difficulty tracking imported drugs to their origin, particularly in cases of counterfeit drugs that cause harm to patients, and insufficient resources for screening and verifying drug authenticity. The safety of the US prescription drug supply is of paramount importance.

To support this principle, any reciprocal style approval system should include generic drugs already approved and used in the USA with well established safety and efficacy profiles, as opposed to brand name drugs without previous FDA approval. Importation should occur only from manufacturers approved by non-US peer regulators with strong safety records under a baseline set of requirements for approval, precluding a “regulatory race to the bottom.” In addition, the FDA’s role in drug approval should be.
central. For example, the FDA could reserve the right to require different labeling and the right to deny reciprocal approval of qualifying drugs, along with an opinion explaining its rationale. A reciprocal approval pathway would admittedly place greater demands on the FDA and its existing application backlog, but this has to be balanced with the potential increase in patient access to vital drug treatments. In our study, we focused on reputable non-US regulators that had been proposed as sources of prescription drug imports in prior literature and proposed legislation. Furthermore, as the world’s largest importer of pharmaceuticals, the USA already relies heavily on drugs manufactured outside its borders.21

In addition to safety concerns, critics of importation have taken issue with the quality of non-FDA approved products, particularly in the context of so-called complex products that could require additional testing to demonstrate bioequivalence.48 Among FDA approved drugs with few generic competitors, only 22 drugs (13%) were complex products, of which only 12 had at least one manufacturer with approval from a non-US regulator. If US legislators designed a system to facilitate reciprocal generic drug approval, it could initially exclude this small number of complex drugs to build trust while studying the interchangeability of complex products approved outside the US to determine whether the system could eventually be extended to this class of products.

Our findings suggest that a formalized system of importation of off-patent prescription drugs with insufficient US generic competition would not on its own fully solve the issue of their rising prices and shortages. One way in which such a system may increase competition is by providing incentives for international suppliers to enter US drug markets by lowering the cost of FDA approval. Reciprocal approval could also facilitate regulatory responses to mitigate bad public health outcomes when drugs face shortages or dramatic price increases. Thus, international sources could lead to increased US competition for a meaningful number of drugs and might be worth pursuing in concert with other strategies. These strategies would include continuing the increased resources and capacity at FDA; prioritizing approvals and waiving application fees for drugs with few generic versions; and where medically appropriate, permitting automatic substitution of drugs within treatment classes at the pharmacy level.49

Study limitations
Although we reviewed all FDA approvals since 1939, our results could be conservative because we only included novel drugs approved in tablet or capsule formulation, and did not account for other treatments such as reformulated drugs and combination therapies. In addition, when we categorized non-US regulators’ approvals, we maintained consistency by only considering drugs approved in the same formulation. Our results could also be conservative because our analysis used data on regulatory drug approvals, which does not necessarily indicate that the generic drug was marketed. Thus, there might be many more off-patent products that would qualify for reciprocal approval because some US generic manufacturers have decided to stop manufacturing the product without formally withdrawing their regulatory approval.

Conversely, some of the manufacturers approved in non-US markets may have never launched or may have stopped selling the product in those settings. Furthermore, our analysis describes only the number of potential additional non-US manufacturers, without accounting for any effect on international supply; that is, some non-US approved generic manufacturers might not be able to supply the US market because of limited production capacities.

Finally, our estimates for prescription drug spending could be conservative given that we focused only on Medicaid spending, although the government program covers nearly one quarter of the US population.50 Further studies might offer comparative analyses of Medicaid spending on drugs with limited US generic competition versus overall Medicaid spending on all generic drugs.

Conclusion
More than half the off-patent drugs approved by the FDA since 1939 with limited generic competition in the USA had at least one additional non-US manufacturer. Importation would help slightly less than half reach a threshold number of generic competitors sufficient to address periods of drug shortages and substantial price increases. Reciprocal approval of prescription drugs in the USA from international sources could help with rising prices and shortages of off-patent drugs, but only along with other strategies dealing with the domestic causes of generic drug market failures.

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Data sharing: Data files are available from the authors on reasonable request.

Transparency: The lead authors (the manuscript’s guarantors) affirm that the manuscript is an honest, accurate, and transparent account of the study being reported, that no important aspects of the study have been omitted, and that any discrepancies from the study as planned have been explained.

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Web appendix: Drugs with manufacturers