Testimony
Before the Subcommittee on Oversight and Investigations, Committee on Energy and Commerce, House of Representatives

DRUG DISCOUNT PROGRAM
Update on Agency Efforts to Improve 340B Program Oversight

Statement of Debra A. Draper
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Accessible Version
DRUG DISCOUNT PROGRAM

Update on Agency Efforts to Improve 340B Program Oversight

What GAO Found

The 340B Drug Pricing Program requires drug manufacturers to sell outpatient drugs at discounted prices to covered entities—eligible clinics, hospitals, and others—to have their drugs covered by Medicaid. Covered entities are only allowed to provide 340B drugs to certain eligible patients. Entities dispense 340B drugs through in-house pharmacies or contract pharmacies, which are outside pharmacies entities contract with to dispense drugs on their behalf. The number of contract pharmacies has increased significantly in recent years.

In its September 2011 report, GAO found that the Health Resources and Services Administration’s (HRSA) oversight of the 340B program was inadequate to ensure compliance with program rules, and GAO recommended actions that HRSA should take to improve program integrity, particularly given significant growth in the program in recent years. HRSA has taken steps to address two of GAO’s four recommendations:

- **HRSA initiated audits of covered entities.** GAO found that HRSA’s oversight of the 340B Program was weak because it primarily relied on covered entities and manufacturers to ensure their own compliance with program requirements and HRSA engaged in few oversight activities. GAO recommended that HRSA conduct audits of covered entities and in fiscal year 2012, HRSA implemented a systematic approach to conducting annual audits of covered entities. HRSA now conducts 200 audits a year, which have identified instances of non-compliance with program requirements, including the dispensing of drugs to ineligible patients.

- **HRSA clarified guidance for manufacturers.** GAO found a lack of specificity in guidance for manufacturers for handling cases in which distribution of drugs is restricted, such as when there is a shortage in drug supply. GAO recommended that HRSA refine its guidance. In May 2012, HRSA clarified its policy for when manufacturers restricted distribution of a drug and provided additional detail on the type of information manufacturers should include in their restricted distribution plans.

- **HRSA has not clarified guidance on two issues.** GAO also found that HRSA guidance on (1) the definition of an eligible patient and (2) hospital eligibility criteria for program participation lacked specificity and recommended that HRSA clarify its guidance. HRSA agreed that clearer guidance was necessary and, in 2015, released proposed guidance that addressed both issues. However, earlier this year, the agency withdrew that guidance in accordance with recent directives to freeze, withdraw, or postpone pending federal guidance.

Given particular concerns that the significant escalation in the number of contract pharmacies poses a potential risk to the integrity of the 340B Program, GAO was asked to examine this issue and expects to issue a future report, in which it plans to address the extent to which covered entities use contract pharmacies; financial arrangements between covered entities and pharmacies; the provision of discounts on drugs dispensed by contract pharmacies to low-income, uninsured patients; and how covered entities and HRSA ensure compliance with 340B program requirements at contract pharmacies.

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**Why GAO Did This Study**

According to HRSA, the purpose of the 340B Program, which was created in 1992, is to enable covered entities to stretch scarce federal resources to reach more eligible patients, and provide more comprehensive services. Covered entities can provide 340B drugs to patients regardless of income or insurance status and generate revenue by receiving reimbursement from patients insurance. The program does not specify how this revenue is to be used or whether discounts are to be passed on to patients. The number of participating covered entity sites—currently about 38,000—has almost doubled in the past 5 years and the number of contract pharmacies increased from about 1,300 in 2010 to around 18,700 in 2017. In recent years, questions have been raised regarding oversight of the 340B Program, particularly given the program’s growth over time.

In September 2011, GAO identified inadequacies in HRSA’s oversight of the 340B program and made recommendations for improvement. This statement describes (1) HRSA actions in response to GAO recommendations to improve its program oversight, and (2) ongoing GAO work regarding the 340B program and HRSA oversight.

For this statement, GAO obtained information and documentation from HRSA officials about any significant program updates and steps they have taken to implement the 2011 GAO recommendations. More detailed information on the objectives, scope, and methodology can be found in GAO’s September 2011 report.

View GAO-17-749T. For more information, contact Debra A. Draper at (202) 512-7114 or draperd@gao.gov.
Chairman Murphy, Ranking Member DeGette, and Members of the Subcommittee:

I am pleased to be here today as you examine the 340B Drug Pricing Program (340B Program), including issues concerning its oversight. The program, created in 1992 and named for the statutory provision authorizing it in the Public Health Service Act (PHSA), requires drug manufacturers to sell outpatient drugs at discounted prices to eligible clinics, hospitals, and other entities—commonly referred to as covered entities—in order to have their drugs covered by Medicaid. According to the Health Resources and Services Administration (HRSA), the agency within the Department of Health and Human Services (HHS) responsible for administering and overseeing the 340B Program, the purpose of the program is to enable covered entities to stretch scarce federal resources to reach more eligible patients, and provide more comprehensive services. In recent years, questions have been raised regarding HRSA’s oversight of the 340B Program, particularly given growth in the program over time. According to HRSA, as of January 2017, covered entities had more than 38,000 sites participating in the 340B Program—almost double the number reported just 5 years earlier.

Participation in the 340B Program is voluntary for both covered entities and drug manufacturers, but there are strong incentives to participate:

- Covered entities can realize substantial savings through 340B price discounts—an estimated 20 to 50 percent of the cost of the drugs, according to HRSA. In addition, covered entities can generate 340B revenue. For example, they can purchase drugs at 340B prices for all eligible patients regardless of the patients’ income or insurance status and generate revenue, such as by receiving reimbursement from a patient’s insurance that may exceed the 340B price paid for the drugs.


Data represent both unique covered entities and all their eligible sites, such as satellite clinics. According to HRSA, there were 12,340 unique organizations participating in the program as of January 1, 2017.
The 340B Program does not dictate how covered entities should use this revenue or require that discounts on the drugs be passed on to patients.

- Incentives for participation by drug manufacturers also are strong because they must participate in the 340B Program to receive Medicaid reimbursement for their drugs. According to HRSA, most manufacturers that produce outpatient drugs have participated in the program since its inception.

HRSA also requires program participants to meet certain conditions set forth both in law and agency guidance. For example, covered entities are prohibited from diverting 340B drugs—that is, transferring 340B drugs to individuals who are not eligible patients of the entities. Similarly, to help ensure covered entities receive discounts to which they are entitled, HRSA has issued guidance (referred to as “HRSA’s nondiscrimination guidance” throughout this statement) prohibiting drug manufacturers from distributing drugs in ways that would discriminate against covered entities compared to non-340B health care providers, such as by imposing minimum purchase requirements or other restrictive conditions.

In a September 2011 report, we identified inadequacies in HRSA’s oversight of this program and recommended actions that should be taken to improve oversight and ensure appropriate use of the program. Since then, we have been monitoring HRSA’s progress in addressing our recommendations, including at a March 24, 2015, hearing before your Subcommittee on Health. My statement today will describe HRSA actions in response to GAO recommendations to address (1) weaknesses in oversight of the 340B program and (2) the lack of clarity in program guidance. The statement will also (3) describe ongoing GAO work regarding the 340B program and HRSA oversight.

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For this statement, we obtained information and documentation from HRSA officials about any significant program updates, and steps they have taken to implement our 2011 recommendations. More detailed information on the objectives, scope, and methodology for our 2011 report can be found in that report.8 We conducted our work for the 2011 report from September 2010 to September 2011, and updated this work in February and March 2015 and again in June and July 2017. The work upon which this statement is based was conducted in accordance with generally accepted government auditing standards. Those standards require that we plan and perform the audit to obtain sufficient, appropriate evidence to provide a reasonable basis for our findings and conclusions based on our audit objectives. We believe that the evidence obtained provides a reasonable basis for our findings and conclusions based on our audit objectives.

Background

The 340B Program was created following the enactment of the Medicaid Drug Rebate Program and gives 340B covered entities discounts on outpatient drugs comparable to those made available to state Medicaid agencies.9 HRSA is responsible for administering and overseeing the 340B Program.

Program Participants

Eligibility for the 340B Program, which is defined in the PHSA, has expanded over time, most recently through the Patient Protection and Affordable Care Act, which extended eligibility to additional types of hospitals.10 Entities generally become eligible by receiving certain federal grants or by being one of six hospital types. Eligible grantees include clinics that offer primary and preventive care services, such as Federally

8GAO-11-836.

9The Medicaid Drug Rebate Program was established through the Omnibus Budget Reconciliation Act of 1990 and requires drug manufacturers to pay rebates to states as a condition of having their drugs covered by Medicaid. Pub. L. No. 101-508 § 4401, 104 Stat. 1388, 1388-143 (adding 42 U.S.C. § 1396r-8).

Qualified Health Centers, clinics that target specific conditions or diseases that raise public health concerns or are expensive to treat, and state-operated AIDS Drug Assistance Programs, which serve as a “payer of last resort” to cover the cost of providing HIV-related medications to certain low-income individuals. Eligible hospitals include certain children’s hospitals, free-standing cancer hospitals, rural referral centers, sole community hospitals, critical access hospitals, and general acute care hospitals that serve a disproportionate number of low-income patients, referred to as disproportionate share hospitals (DSH). To become a covered entity and participate in the program, eligible entities must register with HRSA and be approved. Entity participation in the 340B program has grown over time to include more than 38,000 entity sites, including more than 21,000 hospital sites and nearly 17,000 federal grantee sites (see fig. 1).

Figure 1: Growth in Covered Entity Sites, 2013 to 2017

<table>
<thead>
<tr>
<th>Year</th>
<th>Federal grantees</th>
<th>Hospitals</th>
</tr>
</thead>
<tbody>
<tr>
<td>2013</td>
<td>12,377</td>
<td>7,806</td>
</tr>
<tr>
<td>2014</td>
<td>24,682</td>
<td>11,093</td>
</tr>
<tr>
<td>2015</td>
<td>14,517</td>
<td>15,181</td>
</tr>
<tr>
<td>2016</td>
<td>15,708</td>
<td>16,780</td>
</tr>
<tr>
<td>2017</td>
<td>34,488</td>
<td>21,554</td>
</tr>
</tbody>
</table>

Source: GAO analysis of Health Resources and Services Administration data. | GAO-17-749T

Medicare DSH hospitals receive an additional Medicare payment based on their DSH patient percentage, which is a statutory formula created to identify hospitals that treat a significantly disproportionate number of low-income Medicare and Medicaid patients.
To be eligible for the 340B Program hospitals must meet certain requirements intended to ensure that they perform a government function to provide care to the medically underserved. First, hospitals generally must meet specified DSH adjustment percentages to qualify. Additionally, they must be (1) owned or operated by a state or local government, (2) a public or private nonprofit corporation that is formally delegated governmental powers by a unit of state or local government, or (3) a private, nonprofit hospital under contract with a state or local government to provide health care services to low-income individuals who are not eligible for Medicaid or Medicare.

All drug manufacturers that supply outpatient drugs are eligible to participate in the 340B Program and must participate in order to have their drugs covered by Medicaid. To participate, manufacturers are required to sign a pharmaceutical pricing agreement with HHS in which both parties agree to certain terms and conditions.

Program Structure, Operation, and Key Requirements

The 340B price for a drug—often referred to as the 340B ceiling price—is based on a statutory formula and represents the highest price a participating drug manufacturer may charge covered entities. Covered entities must follow certain requirements as a condition of participating in the 340B Program. For example

- covered entities are prohibited from subjecting manufacturers to “duplicate discounts” in which drugs prescribed to Medicaid beneficiaries are subject to both the 340B price and a rebate through the Medicaid Drug Rebate Program.

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12 Critical access hospitals are exempt from this requirement.

13 According to HRSA, a hospital is said to be “formally granted governmental powers” when the state formally delegates to the hospital a type of power(s) usually exercised by the state, for the purpose of providing health care services to the medically indigent population of the state.

14 Manufacturers may sell a drug at a price that is lower than the ceiling price. As such, covered entities may negotiate prices below the ceiling price.

• covered entities are also prohibited from diverting any drug purchased at the 340B price to an individual who does not meet HRSA’s definition of a patient. This definition, issued in 1996, outlines three criteria that generally state that diversion occurs when 340B discounted drugs are given to individuals who are not receiving health care services from covered entities or are only receiving non-covered services, such as inpatient hospital services.16 (See table 1 for more information on HRSA’s definition of an eligible patient.) Covered entities are permitted to use drugs purchased at the 340B price for all individuals who meet the 340B Program definition of a patient regardless of whether they are low-income, uninsured, or underinsured.

Table 1: Health Resources and Services Administration’s (HRSA) Definition of a Patient Eligible for Discounted Drugs under the 340B Program

<table>
<thead>
<tr>
<th>Criteria for patient eligibility:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The covered entity has established a relationship with the individual such that the covered entity maintains records of the individual's health care.</td>
</tr>
<tr>
<td>2. The individual receives health care services from a health care professional who is either employed by the covered entity or provides health care under contractual or other arrangements (e.g. referral for consultation) such that responsibility for the care provided remains with the covered entity.</td>
</tr>
<tr>
<td>3. The individual receives a health care service or range of services from the covered entity which is consistent with the service or range of services for which grant funding or Federally Qualified Health Center look-alike status has been provided.</td>
</tr>
</tbody>
</table>

Source: GAO analysis of HRSA guidance. | GAO-17-749T


aThese criteria do not apply to AIDS Drug Assistance Programs; rather an individual enrolled in an AIDS Drug Assistance Programs will be considered a patient of that program.

bAn individual is not considered a patient if the only health care service received from the covered entity is the dispensing of a drug or drugs for subsequent self-administration or administration in the home setting.

cDisproportionate share hospitals are exempt from this requirement. Not all federally qualified health centers receive federal grants. Providers that meet all of the requirements for the health center program but do not receive federal grants are referred to as look-alikes and are eligible to participate in the 340B program.

A covered entity typically purchases and dispenses 340B drugs through pharmacies—either through an in-house pharmacy, or through the use of a contract pharmacy arrangement, in which the covered entity contracts with an outside pharmacy to dispense drugs on its behalf. The adoption

and use of contract pharmacies is governed by HRSA guidance. HRSA’s original guidance permitting the use of contract pharmacies limited their use to covered entities that did not have in-house pharmacies and allowed each covered entity to contract with only one outside pharmacy.\textsuperscript{17} However, March 2010 guidance lifted the restriction on the number of pharmacies with which a covered entity could contract.\textsuperscript{18} Since that time, the number of unique contract pharmacies has increased significantly, from about 1,300 at the beginning of 2010 to around 18,700 in 2017 (see fig. 2); and, according to HRSA data, in 2017, there were more than 46,000 contract pharmacy arrangements.\textsuperscript{19} HRSA guidance requires a written contract between the covered entity and each contract pharmacy. Covered entities are responsible for overseeing contract pharmacies to ensure compliance with prohibitions of drug diversion and duplicate discounts. HRSA guidance indicates that covered entities are “expected” to conduct annual independent audits of contract pharmacies, leaving the exact method of ensuring compliance up to the covered entity.


\textsuperscript{18}Notice Regarding 340B Drug Pricing Program—Contract Pharmacy Services, 75 Fed. Reg. 10272 (March 5, 2010).

\textsuperscript{19}Contract pharmacies may have arrangements to dispense drugs for more than one entity. HRSA data indicates that there were 46,174 contract pharmacy arrangements—arrangements between a covered entity site and a pharmacy—as of January 1, 2017. However, the total number of contract pharmacy arrangements is likely higher, as HRSA does not require entities to report all arrangements to the agency.
Note: Data represent the number of unique contract pharmacies participating in the program as of January 1 of each year. Contract pharmacies may have arrangements to dispense drugs for more than one entity.

Drug manufacturers also must follow certain 340B Program requirements. For example, HRSA’s nondiscrimination guidance prohibits manufacturers from distributing drugs in ways that discriminate against covered entities compared to other providers. This includes ensuring that drugs are made available to covered entities through the same channels that they are made available to non-340B providers, and not conditioning the sale of drugs to covered entities on restrictive conditions, which would have the effect of discouraging participation in the program.
HRSA Has Implemented GAO’s Recommendation to Improve Its Oversight of the 340B Program by Conducting Audits

In our September 2011 report, we found that HRSA’s oversight of the 340B Program was weak because it primarily relied on covered entities and manufacturers to police themselves and ensure their own compliance with program requirements. Upon enrollment into the program, HRSA requires participants to self-certify that they will comply with applicable 340B Program requirements and any accompanying agency guidance, and expects participants to develop the procedures necessary to ensure and document compliance, informing HRSA if violations occur. HRSA officials told us that covered entities and manufacturers could also monitor each other’s compliance with program requirements, but we found that, in practice, participants could face limitations to such an approach.

Beyond relying on participants’ self-policing, we also found that HRSA engaged in few activities to oversee the 340B Program and ensure its integrity, which agency officials said was primarily due to funding constraints. Further, although HRSA had the authority to conduct audits of program participants to determine whether program violations had occurred, at the time of our 2011 report, the agency had never conducted such an audit.

In our 2011 report, we concluded that changes in the settings where the 340B Program was used may have heightened the concerns about the inadequate oversight we identified. In the years leading up to our report, the settings where the 340B Program was used had shifted to more contract pharmacies and hospitals than in the past, and that trend has continued in recent years. We concluded that increased use of the 340B Program by contract pharmacies and hospitals may have resulted in a greater risk of drug diversion to ineligible patients, in part because these facilities were more likely to serve patients that did not meet the definition of a patient of the program.

20 GAO-11-836.
To address these oversight weaknesses, we recommended that the Secretary of HHS instruct the administrator of HRSA to conduct selective audits of covered entities to deter potential diversion. In response to that recommendation, in fiscal year (FY) 2012, HRSA implemented a systematic approach to conducting annual audits of covered entities that is outlined on its website. Now numbering 200 per year, HRSA audits include entities that are randomly selected based on risk-based criteria (approximately 90 percent of the audits conducted each year), and entities that are targeted based on information from stakeholders (10 percent of the audits conducted). (See table 2 for the number of audits conducted by HRSA from FY 2012-2017.)

<table>
<thead>
<tr>
<th>Fiscal year</th>
<th>Total audits</th>
</tr>
</thead>
<tbody>
<tr>
<td>2012</td>
<td>51</td>
</tr>
<tr>
<td>2013</td>
<td>94</td>
</tr>
<tr>
<td>2014</td>
<td>99</td>
</tr>
<tr>
<td>2015</td>
<td>200</td>
</tr>
<tr>
<td>2016</td>
<td>200</td>
</tr>
<tr>
<td>2017 (planned)</td>
<td>200</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>844</strong></td>
</tr>
</tbody>
</table>

Source: HRSA. | GAO-17-749T.

As a result of the audits already conducted, HRSA has identified instances of non-compliance with program requirements, including violations related to drug diversion and the potential for duplicate discounts. The agency has developed a process to address non-compliance through corrective action plans. The results of each year’s audits are available on HRSA’s website.

HRSA Implemented One of Three GAO Recommendations to Clarify Program Guidance

In our 2011 report, we found that HRSA’s guidance on three key program requirements lacked the necessary level of specificity to provide clear direction, making it difficult for participants to self-police or monitor others’ compliance, and raising concerns that the guidance could be interpreted in ways that were inconsistent with its intent.²²

First, we found that HRSA’s nondiscrimination guidance was not sufficiently specific in detailing practices manufacturers should follow to ensure that drugs were equitably distributed to covered entities and non-340B providers when distribution was restricted.²³ Some stakeholders we interviewed for the 2011 report, such as covered entities, raised concerns about the way certain manufacturers interpreted and complied with the guidance in these cases. We recommended that HRSA further clarify its nondiscrimination guidance for cases in which distribution of drugs is restricted and require reviews of manufacturers’ plans to restrict distribution of drugs at 340B prices in such cases. In response, HRSA issued a program notice in May 2012 that clarified HRSA’s policy for manufacturers that intend to restrict distribution of a drug and provided additional detail on the type of information manufacturers should include in such restricted distribution plans.²⁴

In addition, we found a lack of specificity in HRSA’s guidance on two other issues—the definition of an eligible patient and hospital eligibility for program participation. Specifically, we found that

- HRSA’s guidance on the definition of an eligible patient lacked the necessary specificity to clearly define the various situations under which an individual was considered eligible for discounted drugs through the 340B Program. As a result, covered entities could interpret the definition either too broadly or too narrowly. At the time of

²²GAO-11-836.

²³Restricted distribution may occur when there is a shortage in drug supply or when shortages are anticipated.

our report, agency officials told us they recognized the need to provide additional clarity around the definition of an eligible patient, in part because of concerns that some covered entities may have interpreted the definition too broadly to include non-eligible individuals, such as those seen by providers who were only loosely affiliated with a covered entity.

- HRSA had not issued guidance specifying the criteria under which hospitals that were not publicly owned or operated could qualify for the 340B Program. For example, we found HRSA guidance lacking on one of the ways hospitals could qualify for the program, namely by executing a contract with a state or local government to provide services to low-income individuals who are not eligible for Medicaid or Medicare. Specifically, we found that HRSA did not outline any criteria that must be included in such contracts, such as the amount of care a hospital must provide to these low-income individuals, and did not require the hospitals to submit their contracts for review by HRSA.

As a result, hospitals with contracts that provided a small amount of care to low-income individuals not eligible for Medicaid or Medicare could claim 340B discounts, which may not have been what the agency intended.

Given the lack of specificity in these areas, we recommended that HRSA (1) finalize new, more specific guidance on the definition of an eligible patient, and (2) issue guidance to further specify the criteria that hospitals not publicly owned or operated must meet to be eligible for the 340B program. HRSA agreed with these recommendations and had planned to address them in a comprehensive 340B Program regulation that it submitted to the Office of Management and Budget for review in April 2014. However, HRSA withdrew this proposed regulation in November 2014 following a May 2014 federal district court ruling that the agency had not been granted broad rulemaking authority to carry out all the provisions of the 340B program. After this ruling, the agency issued a proposed omnibus guidance in August 2015 to interpret statutory requirements for

25We use the term “hospitals that are not publicly owned or operated” to refer to public and private nonprofit corporations as well as private, nonprofit hospitals that may be eligible for the 340B Program. The term does not include private, for-profit hospitals as these hospitals are not eligible for the 340B Program.

26HRSA officials we interviewed for the September 2011 report told us that contracts were selectively reviewed if further clarification was necessary.

the 340B program in areas where it did not have explicit rulemaking authority, including further specificity on the definition of a patient of a covered entity and hospital eligibility for 340B program participation.\textsuperscript{28} However, in January 2017, the agency withdrew the guidance following the administration’s January 20 memorandum directing agencies to withdraw or postpone regulations and guidance that had not yet taken effect.\textsuperscript{29} In July 2017, HRSA indicated that it was working with HHS to determine next steps regarding the proposed Omnibus Guidance, which included the patient definition, but that it was unable to further clarify guidance on hospital eligibility without additional authority.

### GAO Has Ongoing Work Examining HRSA Oversight of 340B Contract Pharmacies

Given the increase in the number of contract pharmacies in the 340B Program and concerns that contract pharmacy arrangements present an increased risk to the integrity of the program, we were asked to review contract pharmacy use under the 340B Program. For this review, we are planning to address the following four questions.

- To what extent do the various types of covered entities use contract pharmacies and where are the pharmacies located?
- What, if any, financial arrangements do covered entities have with contract pharmacies and third-party administrators related to the administration and dispensing of 340B drugs, and how, if at all, this varies by entity type?\textsuperscript{30}
- To what extent do covered entities provide low-income, uninsured patients with discounts on drugs dispensed by contract pharmacies?
- How, if at all, do covered entities and HRSA ensure compliance with 340B program requirements at contract pharmacies?


\textsuperscript{30}Third-party administrators are private companies that some covered entities contract with to manage systems for patient eligibility, program finances, and 340B inventory.
We are in the early stages of this work, and we expect to issue a future report on 340B contract pharmacies.

Chairman Murphy, Ranking Member DeGette, and Members of the Committee, this concludes my statement. I would be pleased to respond to any questions you may have.

GAO Contacts and Staff Acknowledgments

For further information about this statement, please contact Debra A. Draper at (202) 512-7114 or draperd@gao.gov. Contact points for our Offices of Congressional Relations and Public Affairs may be found on the last page of this testimony. Key contributors to this statement were Michelle Rosenberg, Assistant Director; Rotimi Adebonojo, Jennie Apter; and Amanda Cherrin.

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