Inquiries about this report may be addressed to the Office of Public Affairs at Public.Affairs@oig.hhs.gov.
Office of Inspector General
https://oig.hhs.gov/

The mission of the Office of Inspector General (OIG), as mandated by Public Law 95-452, as amended, is to protect the integrity of the Department of Health and Human Services (HHS) programs, as well as the health and welfare of beneficiaries served by those programs. This statutory mission is carried out through a nationwide network of audits, investigations, and inspections conducted by the following operating components:

Office of Audit Services

The Office of Audit Services (OAS) provides auditing services for HHS, either by conducting audits with its own audit resources or by overseeing audit work done by others. Audits examine the performance of HHS programs and/or its grantees and contractors in carrying out their respective responsibilities and are intended to provide independent assessments of HHS programs and operations. These assessments help reduce waste, abuse, and mismanagement and promote economy and efficiency throughout HHS.

Office of Evaluation and Inspections

The Office of Evaluation and Inspections (OEI) conducts national evaluations to provide HHS, Congress, and the public with timely, useful, and reliable information on significant issues. These evaluations focus on preventing fraud, waste, or abuse and promoting economy, efficiency, and effectiveness of departmental programs. To promote impact, OEI reports also present practical recommendations for improving program operations.

Office of Investigations

The Office of Investigations (OI) conducts criminal, civil, and administrative investigations of fraud and misconduct related to HHS programs, operations, and beneficiaries. With investigators working in all 50 States and the District of Columbia, OI utilizes its resources by actively coordinating with the Department of Justice and other Federal, State, and local law enforcement authorities. The investigative efforts of OI often lead to criminal convictions, administrative sanctions, and/or civil monetary penalties.

Office of Counsel to the Inspector General

The Office of Counsel to the Inspector General (OCIG) provides general legal services to OIG, rendering advice and opinions on HHS programs and operations and providing all legal support for OIG’s internal operations. OCIG represents OIG in all civil and administrative fraud and abuse cases involving HHS programs, including False Claims Act, program exclusion, and civil monetary penalty cases. In connection with these cases, OCIG also negotiates and monitors corporate integrity agreements. OCIG renders advisory opinions, issues compliance program guidance, publishes fraud alerts, and provides other guidance to the health care industry concerning the anti-kickback statute and other OIG enforcement authorities.
NOTICES

THIS REPORT IS AVAILABLE TO THE PUBLIC at https://oig.hhs.gov

Section 8M of the Inspector General Act, 5 U.S.C. App., requires that OIG post its publicly available reports on the OIG website.

OFFICE OF AUDIT SERVICES FINDINGS AND OPINIONS

The designation of financial or management practices as questionable, a recommendation for the disallowance of costs incurred or claimed, and any other conclusions and recommendations in this report represent the findings and opinions of OAS. Authorized officials of the HHS operating divisions will make final determination on these matters.
Why OIG Did This Review
For a covered outpatient drug to be eligible for Federal reimbursement under the Medicaid program’s drug rebate requirements, manufacturers must pay rebates to the States for the drugs. However, a prior OIG review found that States did not always invoice and collect all rebates due for drugs administered by physicians.

Our objective was to determine whether Connecticut complied with Federal Medicaid requirements for invoicing manufacturers for rebates for physician-administered drugs.

How OIG Did This Review
We reviewed claims for physician-administered drugs paid between January 2012 and December 2016.

We used the Centers for Medicare & Medicaid Services’ (CMS’s) Medicare Part B crosswalk and the CMS Medicaid Drug File to identify single-source and multiple-source drugs. Additionally, we determined whether the Healthcare Common Procedure Coding System codes were published in CMS’s top-20 multiple-source drug listing.

Connecticut Claimed Unallowable Federal Reimbursement for Medicaid Physician-Administered Drugs That Were Not Invoiced to Manufacturers for Rebates

What OIG Found
Connecticut did not always comply with Federal Medicaid requirements for invoicing manufacturers for rebates for physician-administered drugs. Connecticut did not invoice manufacturers for rebates associated with $1.1 million (Federal share) in physician-administered drugs. Of this amount, $1.07 million was for single-source drugs, and $46,210 was for top-20 multiple-source drugs. Further, Connecticut did not submit the utilization data necessary to secure rebates for all other physician-administered drug claims totaling $2.8 million (Federal share).

What OIG Recommends and Connecticut Comments
We recommend that Connecticut refund to the Federal Government $1.07 million (Federal share) for claims for single-source physician-administered drugs, and $46,210 for claims for top-20 multiple-source physician-administered drugs, and work with CMS to determine the unallowable portion of the $2.8 million (Federal share) for other claims for outpatient physician-administered drugs that were at issue. We also make procedural recommendations to Connecticut.

Connecticut concurred with our first two recommendations and, for our other recommendations, described corrective actions that it had taken or planned to take.

The full report can be found at https://oig.hhs.gov/oas/reports/region7/71806078.asp.
# TABLE OF CONTENTS

## INTRODUCTION
- Why We Did This Review ................................................................. 1
- Objective ......................................................................................... 1
- Background ..................................................................................... 1
  - Medicaid Drug Rebate Program .................................................. 1
  - Physician-Administered Drugs ...................................................... 2
  - The State Agency’s Medicaid Drug Rebate Program ................. 2
- How We Conducted This Review ..................................................... 3

## FINDINGS
- Federal and State Requirements and State Agency Guidance ........ 4
  - The State Agency Did Not Invoice Manufacturers for Rebates on
    Some Single-Source Physician-Administered Drugs ..................... 5
  - The State Agency Did Not Invoice Manufacturers for Rebates on
    Some Top-20 Multiple-Source Physician-Administered Drugs ....... 5
  - The State Agency Did Not Invoice Manufacturers for Rebates on
    Other Physician-Administered Drugs ............................................ 5

## RECOMMENDATIONS ........................................................................ 6

## STATE AGENCY COMMENTS ............................................................ 6

## APPENDICES
- A: Audit Scope and Methodology ..................................................... 7
- B: Related Office of Inspector General Reports ............................... 9
- C: Federal and State Requirements and State Agency Guidance
  Related to Physician-Administered Drugs ....................................... 12
- D: State Agency Comments ............................................................ 14
INTRODUCTION

WHY WE DID THIS REVIEW

For a covered outpatient drug to be eligible for Federal reimbursement under the Medicaid program’s drug rebate requirements, manufacturers must pay rebates to the States for the drugs. States generally offset their Federal share of these rebates against their Medicaid expenditures. States invoice the manufacturers for rebates to reduce the cost of drugs to the program. However, a prior Office of Inspector General review found that States did not always invoice and collect all rebates due for drugs administered by physicians.¹ (Appendix B lists previous reviews of the Medicaid drug rebate program.) For this audit, we reviewed the Connecticut Department of Social Services’ (State agency’s) invoicing for rebates for physician-administered drugs for the period January 1, 2012, through December 31, 2016.

OBJECTIVE

Our objective was to determine whether the State agency complied with Federal Medicaid requirements for invoicing manufacturers for rebates for physician-administered drugs.

BACKGROUND

Medicaid Drug Rebate Program

The Medicaid drug rebate program became effective in 1991 (the Social Security Act (the Act) § 1927). For a covered outpatient drug to be eligible for Federal reimbursement under the program, the drug’s manufacturer must enter into a rebate agreement with the Centers for Medicare & Medicaid Services (CMS) and pay quarterly rebates to the States. CMS, the States, and drug manufacturers each have specific functions under the program.

Manufacturers are required to submit a list to CMS of all covered outpatient drugs and to report each drug’s average manufacturer price and, where applicable, best price.² On the basis of this information, CMS calculates a unit rebate amount for each drug and provides the information to the States each quarter. Covered outpatient drugs reported by participating drug manufacturers are listed in the CMS Medicaid Drug File, which identifies drugs with such fields as National Drug Code (NDC), unit type, units per package size, and product name.

Section 1903(i)(10) of the Act prohibits Federal reimbursement for States that do not capture the information necessary for invoicing manufacturers for rebates as described in section 1927 of the Act. To invoice for rebates, States capture drug utilization data that identifies, by NDC, the number of units of each drug for which the States reimbursed Medicaid providers and

¹ States’ Collection of Medicaid Rebates for Physician-Administered Drugs (OEI-03-09-00410), issued June 24, 2011.
² Section 1927(b) of the Act and section II of the Medicaid rebate agreement.
report the information to the manufacturers (the Act § 1927(b)(2)(A)). The number of units is multiplied by the unit rebate amount to determine the actual rebate amount due from each manufacturer.

States report drug rebate accounts receivable data to CMS on the Medicaid Drug Rebate Schedule. This schedule is part of the Quarterly Medicaid Statement of Expenditures for the Medical Assistance Program report, which contains a summary of actual Medicaid expenditures for each quarter and is used by CMS to reimburse States for the Federal share of Medicaid expenditures.

**Physician-Administered Drugs**

Drugs administered by a physician are typically invoiced to the Medicaid program on a claim form using Healthcare Common Procedure Coding System (HCPCS) codes. For purposes of the Medicaid drug rebate program, physician-administered drugs are classified as either single-source or multiple-source.

The Deficit Reduction Act of 2005 (DRA) amended section 1927 of the Act to specifically address the collection of rebates on physician-administered drugs for all single-source physician-administered drugs and for the top 20 multiple-source physician-administered drugs. Beginning on January 1, 2007, CMS was responsible for publishing annually the list of the top 20 multiple-source drugs by HCPCS codes that had the highest dollar volume dispensed. Before the DRA, many States did not collect rebates on physician-administered drugs if the drug claims did not contain NDCs. NDCs enable States to identify the drugs and their manufacturers and facilitate the collection of rebates for the drugs.

**The State Agency’s Medicaid Drug Rebate Program**

The State agency is responsible for paying claims, submitting invoices to manufacturers, and collecting Medicaid drug rebates for physician-administered drugs.

The State agency also requires “the submission of National Drug Codes (NDCs) on all claims with procedure codes for physician-administered drugs.”

---

3 HCPCS codes (sometimes referred to as J-Codes) are used throughout the health care industry to standardize coding for medical procedures, services, products, and supplies.

4 See, e.g., the Act § 1927(a)(7). In general terms, multiple-source drugs are covered outpatient drugs for which there are two or more drug products that are rated therapeutically equivalent by the Food and Drug Administration. See, e.g., the Act § 1927(k)(7). Multiple-source drugs stand in contrast to single-source drugs, which do not have therapeutic equivalents.

5 The term “top-20 multiple-source drugs” is drawn from a CMS classification and describes these drugs in terms of highest dollar volume of physician-administered drugs in Medicaid. The Act § 1927(a)(7)(B)(i).

The State agency uses its claim utilization data for physician-administered drugs, which it derives from claims submitted by providers, to invoice manufacturers quarterly and to maintain a record of rebate accounts receivable due from the manufacturers. The manufacturers then pay the rebates directly to the State agency.

**HOW WE CONDUCTED THIS REVIEW**


We used the CMS Medicaid Drug File to determine whether the NDCs listed on the claims were classified as single-source drugs or multiple-source drugs. For claims submitted without an NDC, we matched the HCPCS code on the drug claim to the HCPCS code on CMS’s Medicare Part B crosswalk to identify the drug classification. Additionally, we determined whether the HCPCS codes were published in CMS’s top-20 multiple-source drug listing.

We conducted this performance audit 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.

Appendix A contains the details of our audit scope and methodology.

**FINDINGS**

The State agency did not always comply with Federal Medicaid requirements for invoicing manufacturers for rebates for physician-administered drugs. The State agency did not invoice for, and collect from manufacturers, rebates associated with $2.2 million ($1.1 million Federal share) in physician-administered drugs.8 Of this amount, $2.13 million ($1.07 million Federal share) was for single-source drugs and $92,000 ($46,000 Federal share) was for top-20 multiple-source drugs.9 Because the State agency’s internal controls did not always ensure that

---

7 The Medicare Part B crosswalk is published quarterly by CMS and is based on drug and biological information submitted to CMS by manufacturers. CMS uses this information along with pricing data submitted by manufacturers to calculate a volume-weighted sales price for each HCPCS code, which becomes the basis for the reimbursement rate the States pay to providers for the following quarter. CMS instructed States that they could use the crosswalk as a reference because HCPCS codes and NDCs are standardized codes used across health care programs.

8 Specifically, the State agency did not invoice manufacturers for rebates associated with drug expenditures that totaled $2,223,194 ($1,111,597 Federal share).

9 Specifically, $2,130,774 ($1,065,387 Federal share) was for single-source drugs and $92,420 ($46,210 Federal share) was for top-20 multiple-source drugs.
it invoiced manufacturers to secure rebates, the State agency improperly claimed Federal reimbursement for these single-source drugs and top-20 multiple-source drugs.

Further, the State agency did not submit the utilization data necessary to secure rebates for all other physician-administered drugs. Although the State agency generally collected the drug utilization data necessary to invoice manufacturers for rebates associated with these claims, providers submitted claims totaling $4.8 million ($2.4 million Federal share) that did not have NDCs. We were unable to determine whether the State agency was required to invoice for rebates for these other physician-administered drug claims that did not have NDCs in the utilization data. Furthermore, under the Medicaid drug rebate program, claims totaling $776,000 ($388,000 Federal share), which contained NDCs, could have been eligible for rebates. Accordingly, we set aside these amounts and are recommending that the State agency work with CMS to determine (1) the unallowable portion of the $4.8 million ($2.4 million Federal share) of claims that were submitted without NDCs and (2) whether the remaining $776,000 ($388,000 Federal share) of claims could have been invoiced to the manufacturers for rebates.

FEDERAL AND STATE REQUIREMENTS AND STATE AGENCY GUIDANCE

The DRA amended section 1927 of the Act to specifically address the collection of rebates on physician-administered drugs. States must capture NDCs for single-source and top-20 multiple-source drugs (the Act § 1927(a)(7)). To secure rebates, States are required to report certain information to manufacturers within 60 days after the end of each rebate period (the Act § 1927(b)(2)(A)). Federal regulations prohibit Federal reimbursement for physician-administered drugs for which a State has not required the submission of claims containing the NDCs (42 CFR § 447.520).

The State agency publishes provider bulletins to clarify and explain new and existing programs and policies for providers and other interested parties. The Connecticut Department of Social Services, Medical Assistance Program, Provider Bulletin, number 2008-35 (June 2008), states that for “claims for dates of service on or after July 1, 2008, the [State agency] will implement new billing requirements to support the Federal Deficit Reduction Act of 2005, which mandates the submission of National Drug Codes (NDCs). . . . This mandate requires the submission of NDCs on all claims with procedure codes for physician administered drugs.” This Provider Bulletin adds that outpatient claims associated with physician-administered drugs require a HCPCS code and corresponding NDC information; claims that are missing the NDC will be denied.

Appendix C contains Federal and State requirements and guidance related to physician-administered drugs.

---

10 Specifically, claims submitted without NDCs totaled $4,760,710 ($2,380,355 Federal share).

11 Specifically, claims that could have been eligible for rebates totaled $775,522 ($387,761 Federal share).
THE STATE AGENCY DID NOT INVOICE MANUFACTURERS FOR REBATES ON SOME SINGLE-SOURCE PHYSICIAN-ADMINISTERED DRUGS

The State agency improperly claimed Federal reimbursement of $2.13 million ($1.07 million Federal share) for single-source physician-administered drug claims for which it did not invoice manufacturers for rebates.

Because the State agency did not invoice for rebates for all single-source physician-administered drugs, these claims were not eligible for Federal reimbursement.

THE STATE AGENCY DID NOT INVOICE MANUFACTURERS FOR REBATES ON SOME TOP-20 MULTIPLE-SOURCE PHYSICIAN-ADMINISTERED DRUGS

The State agency improperly claimed Federal reimbursement of $92,000 ($46,000 Federal share) for top-20 multiple-source physician-administered drug claims for which it did not invoice manufacturers for rebates.

Before 2012, CMS provided the State agency with an annual listing of top-20 multiple-source HCPCS codes and their respective NDCs. However, the State agency did not always submit the utilization data to the drug manufacturers for rebate purposes.

Because the State agency did not invoice for rebates for all top-20 multiple-source physician-administered drugs, the claims that were not invoiced for rebates were not eligible for Federal reimbursement.

THE STATE AGENCY DID NOT INVOICE MANUFACTURERS FOR REBATES ON OTHER PHYSICIAN-ADMINISTERED DRUGS

We were unable to determine whether, in some cases, the State agency was required to invoice for rebates for other physician-administered drug claims.

Although the State agency generally collected the drug utilization data necessary to invoice the manufacturers for rebates associated with other physician-administered drug claims, providers submitted some claims, totaling $4.8 million ($2.4 million Federal share), that did not have NDCs. Without the NDCs for those claims, we were unable to determine whether the State agency was required to invoice for rebates for these other physician-administered drug claims. Furthermore, under the Medicaid drug rebate program, claims totaling $776,000 ($388,000 Federal share), which contained NDCs, could have been eligible for rebates.

Accordingly, we set aside these amounts and are recommending that the State agency work with CMS to determine (1) the unallowable portion of the $4.8 million ($2.4 million Federal share) of claims that were submitted without NDCs and (2) whether the remaining $776,000 ($388,000 Federal share) of other physician-administered drug claims could have been invoiced.
to the manufacturers to receive rebates and, if so, upon receipt of the rebates, refund the Federal share of the manufacturers’ rebates for those claims.

RECOMMENDATIONS

We recommend that the Connecticut Department of Social Services:

• refund to the Federal Government $1,065,387 (Federal share) for claims for single-source physician-administered drugs that were ineligible for Federal reimbursement;

• refund to the Federal Government $46,210 (Federal share) for claims for top-20 multiple-source physician-administered drugs that were ineligible for Federal reimbursement;

• work with CMS to determine:
  o the unallowable portion of $2,380,355 (Federal share) for other claims for outpatient physician-administered drugs that were submitted without NDCs or with invalid NDCs and that may have been ineligible for Federal reimbursement and refund that amount, and
  o whether the remaining $387,761 (Federal share) of other physician-administered drug claims could have been invoiced to the manufacturers to receive rebates and, if so, upon receipt of the rebates, refund the Federal share of the manufacturers’ rebates for those claims;

• work with CMS to determine and refund the unallowable portion of Federal reimbursement for physician-administered drugs that were not invoiced for rebates after December 31, 2016; and

• strengthen its internal controls to ensure that all physician-administered drugs eligible for rebates are invoiced.

STATE AGENCY COMMENTS

In written comments on our draft report, the State agency concurred with our first two recommendations and, for our other recommendations, described corrective actions that it had taken or planned to take.

The State agency’s comments appear in their entirety as Appendix D.
APPENDIX A: AUDIT SCOPE AND METHODOLOGY

SCOPE


Our audit objective did not require an understanding or assessment of the complete internal control structure of the State agency. We limited our internal control review to obtaining an understanding of the State agency’s processes for reimbursing physician-administered drug claims and its process for claiming and obtaining Medicaid drug rebates for physician-administered drugs.

We conducted our audit work, which included contacting the State agency in Hartford, Connecticut, from February 2018 to May 2019.

METHODOLOGY

To accomplish our objective, we took the following steps:

- We reviewed applicable Federal laws, regulations, and guidance pertaining to the Medicaid drug rebate program and physician-administered drugs.

- We reviewed State agency regulations and guidance to providers, including invoicing instructions for physician-administered drugs.

- We reviewed State agency policies and procedures for rebates for physician-administered drugs.

- We interviewed State agency personnel to gain an understanding of the administration of and controls over the Medicaid invoicing and rebate process for physician-administered drugs.

- We obtained listings of the CMS top-20 multiple-source physician-administered drugs, the Medicare Part B crosswalk, and the CMS Medicaid Drug File for our audit period.

- We obtained claim details from the State agency for all physician-administered drugs for the period January 1, 2012, through December 31, 2016.
• We obtained the listing of 340B entities from the State agency.¹²

• We removed drug claims totaling $186,277,248 ($93,138,624 Federal share) that either were not eligible for a drug rebate (including the drug claims submitted by 340B entities) or were invoiced for rebate.

• We reviewed the remaining drug claims totaling $7,759,426 ($3,879,713 Federal share) to determine whether the State agency complied with Federal Medicaid requirements for invoicing manufacturers for rebates for physician-administered drugs. Specifically:
  
  o We used the CMS Medicaid Drug File to determine whether the NDCs listed on the claims were classified as single-source drugs or multiple-source drugs. For claims submitted without an NDC, we matched the HCPCS code on the drug claim to the HCPCS code on CMS’s Medicare Part B crosswalk to identify the drug classification. Additionally, we determined whether the HCPCS codes were published in CMS’s top-20 multiple-source drug listing.
  
  o We identified the top-20 multiple-source drugs by matching the HCPCS code on the drug claim to the HCPCS code on CMS’s top-20 multiple-source drug listing.

  o We identified the other multiple-source drugs (those not identified as single-source or as top-20 multiple-source drugs) as other outpatient physician-administered drugs.

• We discussed the results of our review with State agency officials on March 21, 2019.

We conducted this performance audit 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.

¹² Under the 340B drug pricing program (set forth in 42 U.S.C. § 256b), a 340B entity may purchase reduced-price covered outpatient drugs from manufacturers; examples of 340B entities are disproportionate share hospitals, which generally serve large numbers of low-income and/or uninsured patients, and State AIDS drug assistance programs. Drugs subject to discounts under the 340B drug pricing program are not subject to rebates under the Medicaid drug rebate program. Section 1927(j) of the Act and 42 U.S.C. § 256(a)(5)(A).
<table>
<thead>
<tr>
<th>Report Title</th>
<th>Report Number</th>
<th>Date Issued</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arizona Did Not Bill Manufacturers for Some Rebates for Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-09-16-02031</td>
<td>2/16/2018</td>
</tr>
<tr>
<td>Arkansas Claimed Unallowable Federal Reimbursement for Some Medicaid Physician-Administered Drugs</td>
<td>A-06-16-00018</td>
<td>2/7/2018</td>
</tr>
<tr>
<td>Nebraska Did Not Invoice Rebates to Manufacturers for Physician-Administered Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-07-13-06046</td>
<td>12/12/2017</td>
</tr>
<tr>
<td>Texas Did Not Bill Manufacturers for Some Rebates for Pharmacy Drugs of Medicaid Managed-Care Organizations</td>
<td>A-06-16-00004</td>
<td>12/12/2017</td>
</tr>
<tr>
<td>Washington State Did Not Bill Manufacturers for Some Rebates for Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-09-16-02028</td>
<td>9/26/2017</td>
</tr>
<tr>
<td>Hawaii Did Not Bill Manufacturers for Some Rebates for Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-09-16-02029</td>
<td>9/26/2017</td>
</tr>
<tr>
<td>Nevada Did Not Bill Manufacturers for Some Rebates for Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-09-16-02027</td>
<td>9/12/2017</td>
</tr>
<tr>
<td>Iowa Did Not Invoice Rebates to Manufacturers for Physician-Administered Drugs of Medicaid Managed-Care Organizations</td>
<td>A-07-16-06065</td>
<td>5/5/2017</td>
</tr>
<tr>
<td>Colorado Claimed Unallowable Federal Reimbursement for Some Medicaid Physician-Administered Drugs</td>
<td>A-07-14-06050</td>
<td>1/5/2017</td>
</tr>
<tr>
<td>Report Title</td>
<td>Report Number</td>
<td>Date Issued</td>
</tr>
<tr>
<td>------------------------------------------------------------------------------</td>
<td>-----------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Delaware Did Not Bill Manufacturers for Some Rebates for Physician-Administered Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-03-15-00202</td>
<td>12/30/2016</td>
</tr>
<tr>
<td>Virginia Did Not Bill Manufacturers for Some Rebates for Physician-Administered Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-03-15-00201</td>
<td>12/22/2016</td>
</tr>
<tr>
<td>California Did Not Bill Manufacturers for Rebates For Physician-Administered Drugs Dispensed to Enrollees of Some Medicaid Managed-Care Organizations</td>
<td>A-09-15-02035</td>
<td>12/8/2016</td>
</tr>
<tr>
<td>Iowa Claimed Unallowable Federal Reimbursement for Some Medicaid Physician-Administered Drugs</td>
<td>A-07-14-06049</td>
<td>7/22/2015</td>
</tr>
<tr>
<td>Report Title</td>
<td>Report Number</td>
<td>Date Issued</td>
</tr>
<tr>
<td>------------------------------------------------------------------------------</td>
<td>-----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Oregon Did Not Bill Manufacturers for Rebates for Physician-Administered Drugs Dispensed to Enrollees of Medicaid Managed-Care Organizations</td>
<td>A-09-13-02037</td>
<td>3/4/2015</td>
</tr>
<tr>
<td>Louisiana Complied With the Federal Medicaid Requirements for Billing Manufacturers for Rebates for Physician-Administered Drugs</td>
<td>A-06-14-00031</td>
<td>2/10/2015</td>
</tr>
<tr>
<td>The District of Columbia Claimed Unallowable Federal Reimbursement for Some Medicaid Physician-Administered Drugs</td>
<td>A-03-12-00205</td>
<td>8/21/2014</td>
</tr>
<tr>
<td>Idaho Did Not Bill Manufacturers for Rebates for Some Medicaid Physician-Administered Drugs</td>
<td>A-09-12-02079</td>
<td>4/30/2014</td>
</tr>
<tr>
<td>Maryland Claimed Unallowable Federal Reimbursement for Some Medicaid Physician-Administered Drugs</td>
<td>A-03-12-00200</td>
<td>11/26/2013</td>
</tr>
<tr>
<td>Oklahoma Complied With the Federal Medicaid Requirements for Billing Manufacturers for Rebates for Physician-Administered Drugs</td>
<td>A-06-12-00059</td>
<td>9/19/2013</td>
</tr>
<tr>
<td>Nationwide Rollup Report for Medicaid Drug Rebate Collections</td>
<td>A-06-10-00011</td>
<td>8/12/2011</td>
</tr>
<tr>
<td>States’ Collection of Medicaid Rebates for Physician-Administered Drugs</td>
<td>OEI-03-09-00410</td>
<td>6/24/2011</td>
</tr>
</tbody>
</table>
APPENDIX C: FEDERAL AND STATE REQUIREMENTS AND STATE AGENCY GUIDANCE RELATED TO PHYSICIAN-ADMINISTERED DRUGS

FEDERAL LAWS

Under the Medicaid program, States may provide coverage for outpatient drugs as an optional service (the Act § 1905(a)(12)). Section 1903(a) of the Act provides for Federal financial participation (Federal share) in State expenditures for these drugs. The Medicaid drug rebate program, created by the Omnibus Budget Reconciliation Act of 1990 that added section 1927 to the Act, became effective on January 1, 1991. Manufacturers must enter into a rebate agreement with the Secretary of Health and Human Services (HHS) and pay rebates for States to receive Federal funding for the manufacturer’s covered outpatient drugs dispensed to Medicaid patients (the Act § 1927(a)). Responsibility for the drug rebate program is shared among the drug manufacturers, CMS, and the States.

Section 6002 of the DRA added section 1927(a)(7) to the Act to require that States capture information necessary to secure rebates from manufacturers for certain covered outpatient drugs administered by a physician. In addition, section 6002 of the DRA amended section 1903(i)(10) of the Act to prohibit a Medicaid Federal share for covered outpatient drugs administered by a physician unless the States collect the utilization and coding data described in section 1927(a)(7) of the Act.

Section 1927(a)(7) of the Act requires that States shall provide for the collection and submission of such utilization data and coding (such as J-codes and NDCs) for each such drug as the Secretary may specify as necessary to identify the manufacturer of the drug in order to secure rebates for all single-source physician-administered drugs effective January 1, 2006, and for the top 20 multiple-source drugs effective January 1, 2008. Section 1927(a)(7)(C) of the Act stated that, effective January 1, 2007, the utilization data must be submitted using the NDC. To secure rebates, States are required to report certain information to manufacturers within 60 days after the end of each rebate period (the Act § 1927(b)(2)(A)).

Section 1927(a)(7)(D) of the Act allowed HHS to delay any of the above requirements to prevent hardship to States that required additional time to implement the physician-administered drug reporting requirements.

FEDERAL REGULATIONS

Federal regulations set conditions for States to obtain a Federal share for covered outpatient drugs administered by a physician and specifically state that no Federal share is available for physician-administered drugs for which a State has not required the submission of claims using codes that identify the drugs sufficiently for the State to bill a manufacturer for rebates (42 CFR § 447.520).
STATE REGULATIONS AND GUIDANCE

The State agency publishes provider bulletins to clarify and explain new and existing programs and policies for providers and other interested parties. The Connecticut Department of Social Services, Medical Assistance Program, Provider Bulletin, number 2008-35 (June 2008), states that for “claims for dates of service on or after July 1, 2008, the [State agency] will implement new billing requirements to support the Federal Deficit Reduction Act of 2005, which mandates the submission of National Drug Codes (NDCs). . . . This mandate requires the submission of NDCs on all claims with procedure codes for physician administered drugs.” This Provider Bulletin adds that outpatient claims associated with physician-administered drugs will require a HCPCS code and corresponding NDC information; claims that are missing the NDC will be denied.

This requirement became effective July 1, 2008, and was in effect for our entire audit period.
July 18, 2019

Department of Health and Human Services
Office of Audit Service, Region VII
Attn: Patrick J. Cogley, Regional Inspector General for Audit Services
601 East 12th Street, Room 0429
Kansas City, MO 64106

Re: Draft Audit Report A-07-18-06078

Dear Mr. Cogley,

The Connecticut Department of Social Services (DSS) is in receipt of the draft audit report issued by the Office of Inspector General (OIG) entitled 'Connecticut Claimed Unallowable Federal Reimbursement for Medicaid Physician-Administered Drugs That Were Not Invoiced to Manufacturers For Rebate' for the period of January 1, 2012 through December 31, 2016. Thank you for the opportunity to respond.

DSS concurs with the recommendations 1 and 2 as noted in the draft audit report to refund the Federal Government $1,065,387 (Federal Share) for claims for single source and $46,210 (Federal Share) for claims for top-20 multiple-source physician administered drugs that were ineligible for Federal reimbursement.

In regards to recommendation 3 and 4, DSS agrees to work with CMS to determine the unallowable portion for drugs that either had no NDC, or invalid NDCs, and provide the appropriate Federal reimbursement. We will also continue to work with CMS to determine the amount of CMS reimbursement for drugs that were not invoiced for rebates after December 31, 2016.

Finally, in regards to recommendation 5, the Department has begun reviewing and recommending changes to strengthen internal controls in this area to ensure that all eligible physician administered drugs are billed, reimbursed, and invoiced for rebate appropriately.

If you have any questions or require additional information, please contact Sandi Ouellette, Medical Operations Manager, by phone at 860-424-5216 or email at Sandra.ouellette@ct.gov.

Sincerely,

Kate McEvoy, Esq.
Director, Division of Health Services

cc: Francis McCullough