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/S/

FROM: Stuart Wright
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SUBJECT: Memorandum Report: Comparison of First-Quarter 2012 Average Sales Prices and Average Manufacturer Prices: Impact on Medicare Reimbursement for Third Quarter 2012, OEI-03-12-00730

This review was conducted in accordance with the statutory mandate for the Office of Inspector General (OIG) to identify Medicare Part B prescription drugs with average sales prices (ASP) that exceed average manufacturer prices (AMP) by at least 5 percent. This review estimated the financial impact of lowering reimbursement amounts for drugs that met the 5-percent threshold to 103 percent of the AMPs, and also examined the potential effect of a November 2012 final rule that, among other things, specifies the circumstances under which the Centers for Medicare & Medicaid Services (CMS) will make AMP-based price substitutions.

SUMMARY

When Congress established ASP as the primary basis for Medicare Part B drug reimbursement, it also mandated that OIG compare ASPs with AMPs and directed CMS to lower reimbursement for drugs with ASPs that exceed AMPs by a threshold of 5 percent. Since the implementation of the ASP payment methodology in 2005, OIG has fulfilled its responsibility by issuing 26 reports comparing ASPs and AMPs. However, CMS has yet to lower reimbursement in response to OIG's findings and recommendations. This latest comparison examines drugs that exceeded the 5-percent threshold based on either complete or partial AMP data in the first quarter of 2012. Of the 385 drug codes with complete AMP data, 22 exceeded the 5-percent threshold. If reimbursement amounts for all 22 codes had been based on 103 percent of the AMPs in the third quarter of 2012, Medicare would have saved an estimated $739,000 in that quarter alone. Under CMS's price substitution policy, reimbursement amounts for 15 of the 22 drugs would have been reduced, saving an estimated $606,000 in the quarter. Of the 64 drug codes with partial AMP data, 6 exceeded the 5-percent threshold. CMS has

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expressed concern that partial AMP data may not adequately reflect market trends and therefore will not apply its price substitution policy to drugs with partial AMP data. However, we found that pricing comparisons for two of the six codes with partial AMP data seemed to accurately capture market trends; therefore, price reductions may be appropriate in these cases. We could not perform pricing comparisons for an additional 52 drug codes because none of the associated drug products had corresponding AMP data. Manufacturers for 9 percent of the associated drug products had Medicaid drug rebate agreements and were therefore generally required to submit AMPs.

BACKGROUND

The Social Security Act (the Act) mandates that OIG compare ASPs to AMPs.\(^1\) If OIG finds that the ASP for a drug exceeds the AMP by a certain percentage (currently 5 percent), the Act states that the Secretary of Health and Human Services (the Secretary) may disregard the ASP for the drug when setting reimbursement amounts.\(^2\) The Act further states that “... the Inspector General shall inform the Secretary (at such times as the Secretary may specify to carry out this subparagraph) and the Secretary shall, effective as of the next quarter, substitute for the amount of payment ... the lesser of (i) the widely available market price ... (if any); or (ii) 103 percent of the average manufacturer price....”\(^4\)

MEDICARE PART B COVERAGE OF PRESCRIPTION DRUGS

Medicare Part B covers only a limited number of outpatient prescription drugs. Covered drugs include injectable drugs administered by a physician; certain self-administered drugs, such as oral anticancer drugs and immunosuppressive drugs; drugs used in conjunction with durable medical equipment; and some vaccines.

MEDICARE PART B PAYMENTS FOR PRESCRIPTION DRUGS

CMS contracts with private companies to process and pay Medicare Part B claims, including those for prescription drugs. To obtain reimbursement for covered outpatient prescription drugs, health care providers submit claims to Medicare contractors using procedure codes. CMS established the Healthcare Common Procedure Coding System (HCPCS) to provide a standardized coding system for describing the specific items and services provided in the delivery of health care. In the case of prescription drugs, each HCPCS code defines the drug name and the amount of the drug represented by the HCPCS code but does not specify manufacturer or package size information.

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\(^1\) Section 1847A (d)(2)(B) of the Act.
\(^2\) Section 1847A (d)(3)(A) of the Act.
\(^3\) Section 1847A (d)(3)(B)(ii) of the Act provides the Secretary with authority to adjust the applicable threshold percentage in 2006 and subsequent years; however, the threshold percentage has been maintained at 5 percent.
\(^4\) Section 1847A (d)(3)(C) of the Act.
Medicare and its beneficiaries spent over $12 billion for Part B drugs in 2011. Although Medicare paid for more than 500 outpatient prescription drug HCPCS codes that year, most of the spending for Part B drugs was concentrated on a relatively small subset of those codes. In 2011, 62 HCPCS codes accounted for 90 percent of the expenditures for Part B drugs, with only 13 of these codes representing the majority of total Part B drug expenditures.

**Reimbursement Methodology for Part B Drugs and Biologicals**

Medicare Part B pays for most covered drugs using a reimbursement methodology based on ASPs. As defined by law, an ASP is a manufacturer’s sales of a drug to all purchasers in the United States in a calendar quarter divided by the total number of units of the drug sold by the manufacturer in that same quarter. The ASP is net of any price concessions, such as volume discounts, prompt pay discounts, cash discounts, free goods contingent on purchase requirements, chargebacks, and rebates other than those obtained through the Medicaid drug rebate program. Sales that are nominal in amount are exempted from the ASP calculation, as are sales excluded from the determination of “best price” in Medicaid’s drug rebate program.

Manufacturers that participate in the Medicaid drug rebate program must provide CMS with the ASP and volume of sales for each of their national drug codes (NDC) on a quarterly basis, with submissions due 30 days after the close of each quarter. An NDC is an 11-digit identifier that represents a specific manufacturer, product, and package size.

Because Medicare Part B reimbursement for outpatient drugs is based on HCPCS codes rather than NDCs and more than one NDC may meet the definition of a particular HCPCS code, CMS has developed a file that “crosswalks” manufacturers’ NDCs to HCPCS codes. CMS uses information in this crosswalk file to calculate volume-weighted ASPs for covered HCPCS codes.

**Calculation of Volume-Weighted ASPs**

Third-quarter 2012 Medicare payments for most covered drug codes were based on first-quarter 2012 ASP submissions from manufacturers, which were volume weighted

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5 Medicare expenditures for Part B drugs in 2011 were calculated using CMS’s Part B Analytics and Reports (PBAR). The PBAR data for 2011 were 98-percent complete when the data were downloaded in May 2012.

6 Several Part B drugs, including certain vaccines and blood products, are not paid for under the ASP methodology.

7 Section 1847A(c) of the Act, as added by the Medicare Prescription Drug, Improvement, and Modernization Act of 2003, P.L. 108-173.

8 Section 1847A(c)(3) of the Act.

9 Section 1847A(c)(2) of the Act.

10 Pursuant to § 1927(c)(1)(C)(i) of the Act, “best price” is the lowest price available from the manufacturer during the rebate period to any wholesaler, retailer, provider, health maintenance organization, nonprofit entity, or governmental entity within the United States, with certain exceptions.

11 Section 1927(b)(3) of the Act.
using an equation that involves the following variables: the ASP for the 11-digit NDC as reported by the manufacturer, the volume of sales for the NDC as reported by the manufacturer, and the number of billing units in the NDC as determined by CMS.\textsuperscript{12} The amount of the drug contained in an NDC may differ from the amount of the drug specified by the HCPCS code that providers use to bill Medicare. Therefore, the number of billing units in an NDC describes the number of HCPCS code units that are in that NDC. For instance, an NDC may contain 10 milliliters of Drug A, but the corresponding HCPCS code may be defined as only 5 milliliters of Drug A. In this case, there are two billing units in the NDC. CMS calculates the number of billing units in each NDC when developing its crosswalk files.

Under the ASP pricing methodology, the Medicare allowance for most Part B drugs is equal to 106 percent of the volume-weighted ASP for the HCPCS code.\textsuperscript{13} Medicare beneficiaries are generally responsible for 20 percent of this amount in the form of coinsurance.

**The Medicaid Drug Rebate Program and AMPs**

In general, for Federal payment to be available for covered outpatient drugs provided under Medicaid, the Act mandates that drug manufacturers enter into rebate agreements with the Secretary and pay quarterly rebates to State Medicaid agencies.\textsuperscript{14} Under these rebate agreements and pursuant to the Act, manufacturers must provide CMS with the AMPs for each of their NDCs.\textsuperscript{15} As further explained in regulation, manufacturers are required to submit AMPs within 30 days after the end of each quarter.\textsuperscript{16}

The AMP is generally calculated as a weighted average of prices for all of a manufacturer’s package sizes of a drug and is reported for the lowest identifiable quantity of the drug (e.g., 1 milliliter, one tablet, one capsule). By law, AMP is defined as the average price paid to the manufacturer for the drug in the United States by (1) wholesalers for drugs distributed to retail community pharmacies and (2) retail community pharmacies that purchase drugs directly from the manufacturer.\textsuperscript{17, 18}

\textsuperscript{12} The equation that CMS currently uses to calculate volume-weighted ASPs is described in § 1847A(b)(6) of the Act. It is also provided in Appendix A.
\textsuperscript{13} Section 1847A(b)(1) of the Act.
\textsuperscript{14} Sections 1927(a)(1) and (b)(1) of the Act.
\textsuperscript{15} Section 1927(b)(3) of the Act.
\textsuperscript{16} 42 CFR § 447.510.
\textsuperscript{17} Section 1927(k)(1) of the Act, as amended by § 2503 of the Patient Protection and Affordable Care Act, P.L. 111-148.
\textsuperscript{18} Pursuant to § 1927(k)(10) of the Act, “retail community pharmacy” means an independent, chain, supermarket, or mass merchandiser pharmacy that is licensed as a pharmacy by the State and that dispenses medications to the general public at retail prices. Such term does not include a pharmacy that dispenses prescription medications to patients primarily through the mail; nursing home, long-term-care, or hospital pharmacies; clinics; charitable or not-for-profit pharmacies; government pharmacies; or pharmacy benefit managers.
Penalties for Failure To Report Timely Drug Pricing Data

Pursuant to the Act, manufacturers that fail to provide ASP and AMP data on a timely basis may be subject to civil money penalties and/or termination from the drug rebate program. Accordingly, CMS has terminated rebate agreements with a number of manufacturers for failure to report AMPs and, for the purposes of evaluating potential civil money penalties, has referred to OIG manufacturers that failed to submit timely ASPs and AMPs. In accordance with an enforcement initiative announced in September 2010, OIG has imposed civil monetary penalties on certain manufacturers that failed to report timely ASPs and/or AMPs.

OIG’s Monitoring of ASPs and AMPs

To comply with its statutory mandate, OIG has issued 22 quarterly pricing comparisons since the ASP reimbursement methodology for Part B drugs was implemented in January 2005. In addition, OIG has completed four annual overviews of ASPs and AMPs, which examined data across all four quarters of 2007, 2008, 2009, and 2010, respectively.

OIG has consistently recommended that CMS develop a price substitution policy and lower the reimbursement amounts for drugs that exceed the 5-percent threshold as directed by the Act. Although CMS has yet to make any changes to Part B drug reimbursement as a result of OIG’s studies, the agency published a final rule in November 2012 that, among other things, specifies the circumstances under which AMP-based price substitutions will occur beginning in 2013.

CM S’s Price Substitution Policy

According to its November 2012 final rule, CMS will substitute 103 percent of the AMP for the ASP-based reimbursement amount when OIG identifies a HCPCS code that exceeds the 5-percent threshold in two consecutive quarters or three of four quarters. Because CMS believes that comparisons based on partial AMP data may not adequately reflect market trends, the agency will lower reimbursement amounts only when ASP

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19 Sections 1927(b)(3)(C)(i) and (4)(B)(i) of the Act.
20 The Secretary delegated to OIG the responsibility to impose civil money penalties for violations of § 1927(b)(3)(C) of the Act in 59 Fed. Reg. 52967 (Oct. 20, 1994).
23 This is the third time that CMS has pursued rulemaking on AMP-based price substitutions. In July 2010, CMS published a proposed rule that specified the circumstances under which AMP-based price substitutions would occur, effective January 2011; however, the agency opted not to finalize this proposed rule, in part, on impending changes to the definition of AMP (75 Fed. Reg. 73170, 73471 (Nov. 29, 2010)). In November 2011, CMS published a final rule that again specified circumstances under which price substitutions would occur (76 Fed. Reg. 73062, 73473 (Nov. 28, 2011)). Although that final rule took effect in January 2012, CMS did not implement that policy in light of access concerns related to drug shortages.
and AMP comparisons are based on the same set of NDCs (i.e., based on complete AMP data).\textsuperscript{26} Price substitutions will take effect in the quarter after OIG shares the results of its most recent pricing comparison and remain in effect for one quarter.\textsuperscript{27, 28} Drugs identified by the Food and Drug Administration (FDA) as being in short supply will not be eligible for price substitution.\textsuperscript{29}

**METHODOLOGY**

We obtained a file from CMS containing NDC-level ASP data from the first quarter of 2012, which were used to establish Part B drug reimbursement for the third quarter of 2012. This file also includes information that crosswalks NDCs to their corresponding HCPCS codes. Both the ASP data and the crosswalk data were current as of June 21, 2012. We also obtained AMP data from CMS for the first quarter of 2012, which were current as of May 7, 2012.

**Analyzing ASP Data From the First Quarter of 2012**

As mentioned previously, Medicare does not base reimbursement for covered drugs on NDCs; instead, it uses HCPCS codes. Therefore, CMS uses ASP information submitted by manufacturers for each NDC to calculate a volume-weighted ASP for each covered HCPCS code. When calculating these volume-weighted ASPs, CMS includes only NDCs with ASP submissions that are deemed valid. As of June 2012, CMS had established prices for 514 HCPCS codes based on the ASP reimbursement methodology mandated by the Act.\textsuperscript{30} Reimbursement amounts for the 514 HCPCS codes were based on ASP data for 3,197 NDCs.

**Analyzing AMP Data From the First Quarter of 2012**

To ensure that the broadest range of drug codes is subject to OIG’s pricing comparisons, we divided HCPCS codes into the following three groups:

1. **HCPCS codes with complete AMP data**—i.e., HCPCS codes with AMP data for every NDC that CMS used in its calculation of volume-weighted ASPs;

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\textsuperscript{26} 77 Fed. Reg. 68892, 69368 (Nov. 16, 2012).
\textsuperscript{27} See, e.g., 76 Fed. Reg. 73026, 73473 (Nov. 28, 2011). After one quarter, the reimbursement amount will be either 106 percent of the volume-weighted ASP for the current quarter or, if the HCPCS code continues to meet CMS’s price substitution criteria, 103 percent of the volume-weighted AMP for the current quarter.
\textsuperscript{28} To prevent CMS’s policy from inadvertently raising the Medicare reimbursement amount, a price substitution will not occur when the substituted amount is greater than the ASP-based payment amount calculated for the quarter in which the price substitution would take effect (77 Fed. Reg. 68892, 69368 (Nov. 16, 2012)). For example, if the AMP-based substitution amount were $5 and the ASP-based reimbursement amount were $4 for the quarter in which the substitution would take place, CMS would not make the price substitution.
\textsuperscript{29} 77 Fed. Reg. 68892, 69368 (Nov. 16, 2012).
\textsuperscript{30} Section 1847A(b)(6) of the Act.
(2) HCPCS codes with partial AMP data—i.e., HCPCS codes with AMP data for only some of the NDCs that CMS used in its calculation of volume-weighted ASPs; and

(3) HCPCS codes with no AMP data—i.e., HCPCS codes with no AMP data for any of the NDCs that CMS used in its calculation of volume-weighted ASPs.

As previously noted, the AMP for each NDC is reported for the lowest identifiable quantity of the drug contained in that NDC (e.g., 1 milliliter, one tablet, one capsule). In contrast, the ASP is reported for the entire amount of the drug contained in the NDC (e.g., 50 milliliters, 100 tablets). To ensure that the AMP would be comparable to the ASP, it was necessary to convert the AMP for each NDC so that it represented the total amount of the drug contained in that NDC.

To calculate “converted AMPs” for NDCs in the first and second groups, we multiplied the AMP by the total amount of the drug contained in each NDC, as identified by sources such as the CMS crosswalk file, manufacturer Web sites, drug labels, Thomson Reuters’ Red Book, and FDA’s NDC directory. For certain NDCs, we were unable to identify the amount of the drug reflected by the ASP or AMP and therefore could not calculate a converted AMP. Because of these unsuccessful AMP conversions, 13 HCPCS codes were removed from our analysis.

Using NDCs with successful AMP conversions, we then calculated a volume-weighted AMP for each of the corresponding HCPCS codes, consistent with CMS’s methodology for calculating volume-weighted ASPs. When calculating the volume-weighted AMP for a HCPCS code with partial AMP data, we excluded any NDCs without AMPs; however, we did not exclude those NDCs from the corresponding volume-weighted ASP. This means that the volume-weighted AMP for a HCPCS code with partial AMP data is based on fewer NDCs than the volume-weighted ASP for that same code. Appendix B provides a more detailed description of the methods we used to both convert AMPs and calculate volume-weighted AMPs. Table 1 provides the final number of HCPCS codes and NDCs included in our analysis after we removed NDCs with either no AMP data or unsuccessful AMP conversions.

Table 1: Number of Drug Codes and NDCs Included in OIG’s Pricing Comparison

<table>
<thead>
<tr>
<th>Availability of AMP Data for HCPCS Codes</th>
<th>Number of HCPCS Codes</th>
<th>Number of NDCs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complete AMP Data</td>
<td>385</td>
<td>1,713</td>
</tr>
<tr>
<td>Partial AMP Data</td>
<td>64</td>
<td>728</td>
</tr>
<tr>
<td>No AMP Data</td>
<td>52</td>
<td>208</td>
</tr>
</tbody>
</table>


31 We did not calculate converted AMPs for NDCs in the third group because they had no AMP data.
Comparing First-Quarter 2012 Volume-Weighted ASPs and AMPs for HCPCS Codes With Complete AMP Data
For each of the 385 HCPCS codes with complete AMP data, we compared the volume-weighted ASP and AMP and determined whether the ASP for the code exceeded the AMP by at least 5 percent. For HCPCS codes that exceeded the 5-percent threshold, we reviewed the associated NDCs to verify the accuracy of the billing unit information. According to our review, none of the HCPCS codes that exceeded the threshold based on complete AMP data was associated with questionable billing units.

For each of the HCPCS codes that exceeded the 5-percent threshold, we estimated the monetary impact of lowering reimbursement to 103 percent of the AMP. First, we calculated 103 percent of the volume-weighted AMP and subtracted this amount from the third-quarter 2012 reimbursement amount for the HCPCS code. To estimate the financial effect for the third quarter of 2012, we then multiplied the difference by one-fourth of the number of services that were allowed by Medicare for each HCPCS code in 2011, as reported in the PBAR.

To determine which HCPCS codes would have been subject to CMS’s price substitution policy, we identified codes with complete AMP data that met the 5-percent threshold in two consecutive or three of four quarters and were not identified by FDA as being in short supply. We then totaled the estimated third-quarter 2012 savings for that subset of codes.

Comparing First-Quarter 2012 Volume-Weighted ASPs and AMPs for HCPCS Codes With Partial AMP Data
For each of the 64 HCPCS codes with partial AMP data, we compared the volume-weighted ASP and AMP and determined whether the ASP for the code exceeded the AMP by at least 5 percent. For HCPCS codes that exceeded the 5-percent threshold, we reviewed the associated NDCs to verify the accuracy of the billing unit information. According to our review, one of the HCPCS codes that exceeded the threshold based on partial AMP data was associated with questionable billing units. Because volume-weighted ASPs and AMPs are calculated using this billing unit information, we could not be certain that the results for this code was correct. Therefore, we excluded this HCPCS code from our count of codes with partial AMP data that exceeded the 5-percent threshold.

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32 Section 1847A(d)(3)(C) of the Act directs the Secretary to replace payment amounts for drugs that exceed the 5-percent threshold with the lesser of the widely available market price for the drug (if any) or 103 percent of the AMP. For the purposes of this study, we used 103 percent of the AMP to estimate the impact of lowering reimbursement amounts. If widely available market prices had been available for these drugs and lower than 103 percent of the AMP, the savings estimate presented in this report would have been greater.
33 The PBAR data for 2011 were 98-percent complete when the data were downloaded in May 2012. This estimate assumes that the number of services allowed by Medicare in 2011 remained consistent from one quarter to the next and that there were no significant changes in utilization between 2011 and 2012.
For each of the remaining HCPCS codes that exceeded the 5-percent threshold based on partial AMP data, we determined whether missing AMPs unduly influenced the results of our pricing comparison. As mentioned previously, the volume-weighted AMP for a HCPCS code with partial AMP data is based on fewer NDCs than the volume-weighted ASP for that same code. Therefore, there may be a disparity between the volume-weighted ASP and AMP that would not exist if AMP data were available for the full set of NDCs. In other words, the volume-weighted ASP for the HCPCS code could exceed the volume-weighted AMP by at least 5 percent only because AMPs for certain NDCs were not represented.

CMS has expressed concern that partial AMP data may not adequately reflect market trends. Therefore, to identify HCPCS codes with partial AMP data that exceeded the 5-percent threshold only because AMP data were missing, we reanalyzed pricing data after accounting for the missing values. Specifically, we replaced each missing AMP with its corresponding ASP and recalculated the volume-weighted AMPs using those imputed prices. We then compared those new volume-weighted AMPs to the volume-weighted ASPs originally calculated by CMS.

If a HCPCS code no longer exceeded the 5-percent threshold, we concluded that the missing AMPs likely caused the HCPCS code to initially exceed the threshold, as opposed to an actual disparity between ASPs and AMPs in the marketplace.

If a HCPCS code continued to exceed the 5-percent threshold, we concluded that missing AMPs had little impact on the results of our pricing comparison. These HCPCS codes likely exceeded the threshold as a result of actual pricing differences between ASPs and AMPs. Because price substitutions for these HCPCS codes may be warranted, we also identified which of these HCPCS codes met the threshold in two consecutive or three of four quarters.

**Limitations**

We did not verify the accuracy of manufacturer-reported ASP and AMP data, nor did we verify the underlying methodology used by manufacturers to calculate ASPs and AMPs. Furthermore, we did not verify the accuracy of CMS's crosswalk files or examine NDCs that CMS opted to exclude from its calculation of Part B drug reimbursement amounts.

Manufacturers are required to submit their quarterly ASP and AMP data to CMS 30 days after the close of the quarter. Our analyses were performed on ASP and AMP data

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36 Although an NDC’s ASP is not usually the same as its AMP, ASPs were within 3 percent of the AMPs at the median during the last three quarters of 2011 and the first quarter of 2012. Therefore, we believe that ASP acts as a reasonable proxy for AMP, ensuring that the NDC is represented in both the volume-weighted ASP and the volume-weighted AMP for the HCPCS code.

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We generally did not determine whether manufacturers provided additional or revised pricing data to CMS at a later date.

Standards
This inspection was conducted in accordance with the Quality Standards for Inspection and Evaluation approved by the Council of the Inspectors General on Integrity and Efficiency.

RESULTS

Of the 385 Drug Codes With Complete AMP Data, Volume-Weighted ASPs for 22 Exceeded the Volume-Weighted AMPs by at Least 5 Percent

As mandated by the Act, OIG compared ASPs to AMPs to identify instances in which the ASP for a particular drug exceeded the AMP by a threshold of 5 percent. In the first quarter of 2012, 22 of the 385 HCPCS codes with complete AMP data (6 percent) exceeded this 5-percent threshold. Table 2 describes the extent to which ASPs exceeded AMPs for the 22 HCPCS codes. For half of the codes, the volume-weighted ASP exceeded the volume-weighted AMP by more than 20 percent. A list of all 22 HCPCS codes, including their descriptions and HCPCS dosage amounts, is presented in Appendix C.

Table 2: Extent to Which ASPs Exceeded AMPs for 22 HCPCS Codes With Complete AMP Data

<table>
<thead>
<tr>
<th>Percentage</th>
<th>Number of Codes</th>
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<tbody>
<tr>
<td>5.00–9.99%</td>
<td>8</td>
</tr>
<tr>
<td>10.00–19.99%</td>
<td>3</td>
</tr>
<tr>
<td>20.00–29.99%</td>
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<tr>
<td>30.00–39.99%</td>
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<tr>
<td>40.00–49.99%</td>
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<tr>
<td>90.00–99.99%</td>
<td>0</td>
</tr>
<tr>
<td>100% and above</td>
<td>2</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>22</strong></td>
</tr>
</tbody>
</table>


Pursuant to section 1847A(d)(3) of the Act, the Secretary may disregard the ASP for a drug that exceeds the 5-percent threshold and shall substitute the payment amount with the lesser of either the widely available market price or 103 percent of the AMP. If reimbursement amounts for all 22 codes with complete AMP data had been based on

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103 percent of the AMPs during the third quarter of 2012, Medicare expenditures would have been reduced by an estimated $739,000.\textsuperscript{37}

If CMS's price substitution policy had been in effect, reimbursement amounts for 15 of the 22 HCPCS codes would have been reduced. These 15 HCPCS codes had complete AMP data, exceeded the 5-percent threshold in either two consecutive quarters or three of four quarters, and were not identified by FDA as being in short supply as of September 2012 (see Table 3).\textsuperscript{38} If reimbursement amounts for the 15 codes had been based on 103 percent of the AMPs during the third quarter of 2012, Medicare expenditures would have been reduced by an estimated $606,000.

### Table 3: Fifteen HCPCS Codes With Complete AMP Data in the First Quarter of 2012 That Would Have Met CMS's Criteria for Price Substitution

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Source: OIG analysis of ASP and AMP data from the second through fourth quarters of 2011 and the first quarter of 2012.

\textsuperscript{37} All savings estimates in this report assume that the number of services allowed by Medicare in 2011 remained consistent from one quarter to the next and that there were no significant changes in utilization between 2011 and 2012.

\textsuperscript{38} Two additional HCPCS codes had complete AMP data and exceeded the 5-percent threshold in two consecutive or three of four quarters; however, the drugs represented by these HCPCS codes were identified by FDA as being in short supply at the time of our analysis.
Comparison of First-Quarter 2012 ASPs and AMPs (OEI-03-12-00730)

Of the 64 Drug Codes With Partial AMP Data, Volume-Weighted ASPs for 6 Exceeded the Volume-Weighted AMPs by at Least 5 Percent

In addition to examining HCPCS codes with complete AMP data, we examined 64 HCPCS codes for which only partial AMP data were available. ASPs for 6 of these 64 HCPCS codes (9 percent) exceeded the AMPs by at least 5 percent in the first quarter of 2012. A list of the six HCPCS codes, including their descriptions and HCPCS dosage amounts, is presented in Appendix D.

For one-third of the HCPCS codes, missing AMPs likely had little influence on the outcome of the pricing comparisons. Two of the six HCPCS codes with partial AMP data continued to exceed the threshold when we accounted for missing AMPs, suggesting that the pricing comparisons for these codes were accurately capturing underlying market trends even though AMP data were not available for the full set of NDCs. Because missing AMPs likely had little influence on the pricing comparison results for these two HCPCS codes, price substitutions may be legitimately warranted in these cases. Both of these codes exceeded the 5-percent threshold in either two consecutive or three of four quarters.

For the remaining four of six HCPCS codes, ASPs no longer exceeded the AMPs by at least 5 percent in the first quarter of 2012, suggesting that these codes initially exceeded the threshold because of missing AMPs rather than a genuine pricing disparity between the ASPs and AMPs.

Pricing Comparisons Could Not Be Performed on 52 Drug Codes Because No AMP Data Were Available

For 52 HCPCS codes, OIG could not compare ASPs and AMPs because there were no AMP data for any of the 208 NDCs that CMS used when calculating drug reimbursement amounts for these codes. In 2011, Medicare allowances for these 52 codes totaled $338 million.39

Manufacturers for 9 percent of the NDCs without AMP data (19 of 208) participated in the Medicaid drug rebate program as of the first quarter of 2011 and were therefore generally required to submit AMP data for their covered outpatient drugs.40, 41, 42

39 Of the 52 HCPCS codes with no associated AMP data, three were not listed in the 2011 PBAR file. As a result, these codes were not included in the total Medicare allowances for the year.
40 To determine whether a manufacturer participated in the Medicaid drug rebate program, we consulted CMS’s labeler contact file for the first quarter of 2012, which was obtained from CMS on October 2, 2012.
41 Although manufacturers with rebate agreements are required to submit AMP data for their covered outpatient drugs, there may be valid reasons why an AMP was not provided for a specific NDC in a given quarter. For example, a manufacturer may not have been required to submit an AMP if the drug product had been terminated and there was no drug utilization during the quarter.
42 These 19 NDCs were crosswalked to 13 HCPCS codes.
Manufacturers for the remaining 189 of 208 NDCs did not participate in the Medicaid drug rebate program and therefore were not required to submit AMP data.

CONCLUSION

When Congress established ASP as the basis for Medicare Part B drug reimbursement, it also provided a mechanism for monitoring market prices and limiting potentially excessive payment amounts based on ASPs. Specifically, the ASP statute mandates that OIG monitor ASPs by comparing them with AMPs and widely available market prices, and directs CMS to lower reimbursement for certain drugs based on OIG’s findings. Since the ASP payment methodology took effect in January 2005, OIG has fulfilled its responsibility to monitor ASP-based payment amounts by issuing 26 comparisons between ASPs and AMPs, each of which identified Part B drugs that would have been eligible for price reductions under the law. However, CMS has yet to lower reimbursement in response to OIG’s findings and recommendations.

In this current report, we identified 28 HCPCS codes that exceeded the threshold for price adjustment in the first quarter of 2012. Of these, 22 had complete AMP data (i.e., AMP data for every drug product that CMS used to establish reimbursement amounts). If reimbursement amounts for all 22 codes had been based on 103 percent of the AMPs in the third quarter of 2012, Medicare would have saved an estimated $739,000 in that quarter alone. Under CMS’s price substitution policy, reimbursement amounts for 15 of the 22 HCPCS codes would have been lowered to 103 percent of the AMP, thereby saving Medicare and its beneficiaries $606,000. The remaining 6 of 28 HCPCS codes also exceeded the threshold for price adjustment in the first quarter of 2012 but did so based on partial AMP data. Although CMS’s price substitution policy will not apply to codes with partial AMP data, price reductions may be legitimately warranted for two of the six codes because missing AMPs likely had little influence on the pricing comparison results. We could not compare ASPs and AMPs for 52 HCPCS codes because AMP data were not submitted for any of the NDCs that CMS used to calculate reimbursement.

 Manufacturers for 9 percent of these NDCs had Medicaid drug rebate agreements and were therefore generally required to submit AMPs.

Although we do not make recommendations in this report, some of OIG’s previous pricing comparisons have contained recommendations, which we continue to support. In response to OIG’s most recent report with recommendations, CMS expressed uncertainty about the payoff associated with quarterly pricing comparisons, stating that

43 For example, OIG, Comparison of Average Sales Prices and Average Manufacturer Prices: An Overview of 2007, OEI-03-08-00450, December 2008; OIG, Comparison of Average Sales Prices and Average Manufacturer Prices: An Overview of 2008, OEI-03-09-00350, February 2010; OIG, Comparison of Average Sales Prices and Average Manufacturer Prices: An Overview of 2009, OEI-03-10-00380, April 2011; and OIG, Comparison of Average Sales Prices and Average Manufacturer Prices: An Overview of 2010, OEI-03-11-00410, November 2011. These reports are available online at https://www.oig.hhs.gov.

Comparison of First-Quarter 2012 ASPs and AMPs (OEI-03-12-00730)
the price substitution policy will generate minor savings for the program. Although we acknowledge that the savings from any single OIG report may be modest relative to total expenditures for Part B drugs, significant savings would have accrued had CMS taken action immediately after OIG issued its first pricing comparison. In the long term, savings achieved through price substitution could reduce waste and conserve taxpayer funds at a time when increased focus has been placed on rising health care costs and fiscal responsibility.

This report is being issued directly in final form because it contains no recommendations. If you have comments or questions about this report, please provide them within 60 days. Please refer to report number OEI-03-12-00730 in all correspondence.
APPENDIX A

The Equation Used by the Centers for Medicare & Medicaid Services To Calculate Volume-Weighted Average Sales Prices on or After April 1, 2008

A volume-weighted average sales price (ASP) is calculated for the dosage amount associated with the Healthcare Common Procedure Coding System (HCPCS) code. In the following equation, the “number of billing units” represents the number of HCPCS code doses that are contained in a national drug code (NDC).

\[
\text{Volume-Weighted ASP for Dosage Amount of HCPCS Code} = \frac{\text{Sum of (ASP for NDC} \times \text{Number of NDCs Sold)}}{\text{Sum of (Number of NDCs Sold} \times \text{Number of Billing Units in NDC)}}
\]
APPENDIX B

Detailed Methodology for Converting and Volume-Weighting Average Manufacturer Prices for the First Quarter of 2012

Healthcare Common Procedure Coding System codes with complete average manufacturer price data. Of the 514 Healthcare Common Procedure Coding System (HCPCS) codes with reimbursement amounts based on average sales prices (ASP), 397 had average manufacturer prices (AMP) for every national drug code (NDC) that the Centers for Medicare & Medicaid Services (CMS) used to calculate volume-weighted ASPs. These 397 HCPCS codes represented 1,969 NDCs. For 19 NDCs, we could not successfully identify the amount of the drug reflected by the ASP and therefore could not calculate a converted AMP. These 19 NDCs were crosswalked to 12 HCPCS codes. We did not include these 12 HCPCS codes (256 NDCs) in our final analysis.

Using the converted AMPs for the remaining 1,713 NDCs, we calculated a volume-weighted AMP for each of the remaining 385 HCPCS codes consistent with CMS’s methodology for calculating volume-weighted ASPs.

HCPCS codes with partial AMP data. There were 65 HCPCS codes with AMP data for only some of the NDCs that CMS used in its calculation of volume-weighted ASPs. These 65 HCPCS codes represented a total of 1,020 NDCs. AMP data were either missing or unavailable for 285 of these NDCs, which were then excluded from our calculation of volume-weighted AMPs.

We calculated converted AMPs for each of the remaining 735 NDCs. For 7 of the 735 NDCs, we could not identify the amount of the drug reflected by the ASP and therefore could not calculate a converted AMP. We removed these 7 NDCs from our analysis. As a result, one HCPCS code no longer had any NDCs with AMP data. Therefore, this HCPCS code was removed from our analysis.

Using the converted AMPs for the remaining 728 NDCs, we then calculated a volume-weighted AMP for each of the remaining 64 HCPCS codes consistent with CMS’s methodology for calculating volume-weighted ASPs.

44 Although AMP data for these 285 NDCs were excluded from our calculation of volume-weighted AMPs, the corresponding ASPs were not excluded from the volume-weighted ASPs as determined by CMS. Volume-weighted ASPs remained the same, regardless of the availability of AMP data.

45 Although we removed NDCs with problematic AMP conversions, we did not remove the corresponding HCPCS codes, provided that other NDCs for those drug codes had usable AMP data. This differs from our analysis of HCPCS codes with complete AMP data, in which we removed not only the NDCs with problematic AMP conversions, but also the corresponding HCPCS codes.
HCPCS codes with no AMP data. For 52 HCPCS codes, there were no AMP data for any of the NDCs that CMS used in its calculation of volume-weighted ASPs. These 52 HCPCS codes represented 208 NDCs.
## APPENDIX C

### Twenty-Two Drug Codes With Complete Average Manufacturer Price Data That Exceeded the 5-Percent Threshold in the First Quarter of 2012

<table>
<thead>
<tr>
<th>Drug Code</th>
<th>Short Description</th>
<th>Drug Code Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>J0500</td>
<td>Dicyclomine injection</td>
<td>20 mg</td>
</tr>
<tr>
<td>J0595</td>
<td>Butorphanol tartrate injection</td>
<td>1 mg</td>
</tr>
<tr>
<td>J0610</td>
<td>Calcium gluconate injection</td>
<td>10 mL</td>
</tr>
<tr>
<td>J0636</td>
<td>Calcitriol injection</td>
<td>0.1 mcg</td>
</tr>
<tr>
<td>J0670</td>
<td>Mepivacaine HCl injection</td>
<td>10 mL</td>
</tr>
<tr>
<td>J1205</td>
<td>Chlorothiazide sodium injection</td>
<td>500 mg</td>
</tr>
<tr>
<td>J1270</td>
<td>Doxercalciferol injection</td>
<td>1 mcg</td>
</tr>
<tr>
<td>J1742</td>
<td>Ibutidine fumarate injection</td>
<td>1 mg</td>
</tr>
<tr>
<td>J1756</td>
<td>Iron sucrose injection</td>
<td>1 mg</td>
</tr>
<tr>
<td>J1955</td>
<td>Levocarnitine injection</td>
<td>1 g</td>
</tr>
<tr>
<td>J2501</td>
<td>Paricalcitol injection</td>
<td>1 mcg</td>
</tr>
<tr>
<td>J2675</td>
<td>Progesterone injection</td>
<td>50 mg</td>
</tr>
<tr>
<td>J2780</td>
<td>Rantidine HCl injection</td>
<td>25 mg</td>
</tr>
<tr>
<td>J7500</td>
<td>Azathioprine, oral</td>
<td>50 mg</td>
</tr>
<tr>
<td>J9045</td>
<td>Carboplatin injection</td>
<td>50 mg</td>
</tr>
<tr>
<td>J9065</td>
<td>Cladribine injection</td>
<td>1 mg</td>
</tr>
<tr>
<td>J9208</td>
<td>Ifosfamide injection</td>
<td>1 g</td>
</tr>
<tr>
<td>J9211</td>
<td>Idarubicin HCl injection</td>
<td>5 mg</td>
</tr>
<tr>
<td>J9214</td>
<td>Interferon alfa-2b injection</td>
<td>1 million units</td>
</tr>
<tr>
<td>J9280</td>
<td>Mitomycin injection</td>
<td>5 mg</td>
</tr>
<tr>
<td>J9360</td>
<td>Vinblastine sulfate injection</td>
<td>1 mg</td>
</tr>
<tr>
<td>Q0166</td>
<td>Granisetron HCl, oral</td>
<td>1 mg</td>
</tr>
</tbody>
</table>

### APPENDIX D

**Six Drug Codes With Partial Average Manufacturer Price Data That Exceeded the 5-Percent Threshold in the First Quarter of 2012**

<table>
<thead>
<tr>
<th>Drug Code</th>
<th>Short Description</th>
<th>Drug Code Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>90586</td>
<td>Bcg vaccine, for intravesical use</td>
<td>1 each</td>
</tr>
<tr>
<td>J0171</td>
<td>Adrenalin epinephrine injection</td>
<td>0.1 mg</td>
</tr>
<tr>
<td>J1650*</td>
<td>Enoxaparin sodium injection</td>
<td>10 mg</td>
</tr>
<tr>
<td>J2700</td>
<td>Oxacillin sodium injection</td>
<td>250 mg</td>
</tr>
<tr>
<td>J9031</td>
<td>Bcg live intravesical vaccine</td>
<td>1 each</td>
</tr>
<tr>
<td>Q0163*</td>
<td>Diphenhydramine HCl injection</td>
<td>50 mg</td>
</tr>
</tbody>
</table>

*These drug codes continued to exceed the 5-percent threshold after the Office of Inspector General (OIG) accounted for missing average manufacturer prices (AMP).