Testimony of David Balto
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Advisory Council on Employee Welfare and Pension Benefit Plans
U.S. Department of Labor

Hearing on
Section 408(b)(2) Regulation
PBM Compensation and Fee Disclosure

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I appreciate the opportunity to come before you today and testify about PBM compensation and fee disclosures to welfare benefits plans under Section 408(b)(2).

As my testimony outlines, extending compensation and fee disclosure provisions to PBMs will benefit ERISA welfare plan beneficiaries by enabling plans to have the full set of information necessary to make PBM markets work effectively and secure benefits at the lowest cost. Greater disclosure will give plans the necessary tools to detect and prevent conflicts of interest and secure all the appropriate compensation, including undisclosed indirect compensation. Plans currently do not fully benefit from PBM cost control efforts because of conflicts of interest and the ability of PBMs to hide undisclosed indirect compensation, such as rebates from drug manufacturers. Extending the fee disclosure provisions will enable plans to secure the full benefit of compensation received on their behalf.

I am a public interest and antitrust attorney, and have practiced law for over 30 years, both in the government and in private practice. Prior to entering private practice, I was the Policy Director of the Office of Policy and Evaluation for the Bureau of Competition of the Federal Trade Commission and attorney advisor to Chairman Robert Pitofsky and helped direct the first antitrust cases against PBMs. I have counseled health and welfare benefit plans, PBMs, pharmacies, and consumers on PBM competition and consumer protection issues. My comments are based on those decades of enforcement and real world experience.

Today’s hearing comes as a result of EBSA’s efforts to thoroughly understand PBM compensation and fee disclosures and how it affects the provision of health care to plan participants and the costs of plan administration to welfare benefit plans under Section 408(b)(2).

I testified before EBSA in 2010 when it was considering expansion of compensation and fee disclosure requirements under 408(b)(2) leading up to release of an interim final rule on disclosure fees and conflicts of interest affecting retirement plans. My testimony focused on the appropriate level of disclosure by PBMs. I have also testified before Congress on how transparency can improve competition in PBM markets. The PBM market is highly concentrated, and has become even more so over the last few years with a number of mergers recently consummated. PBM contracting practices are complex and the markets are opaque. This provides a fertile environment for deceptive and fraudulent practices – in recent years the two major PBMs have settled 4 major cases brought by state attorneys generals resulting in over $370 million in penalties and fines. Much of the concern raised in these cases involved undisclosed indirect compensation of the type the Council is concerned with today. Thus, I argue compensation and fee disclosure requirements should be applied to welfare plans and PBM services. Greater disclosure is needed in the PBM industry to protect plans, consumers and reduce costs.

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I. PBMs no longer serve as “honest brokers” and engage in a wide range of anticompetitive conduct.

Although PBMs offer a great deal of promise in terms of the potential to control pharmaceutical costs, there is a pattern of conflicts of interest, self-dealing and anticompetitive conduct, all of which ultimately means that consumers pay far more for drugs than necessary. The two dominant PBMs (i.e., CVS Caremark and Express Scripts), as well as other large PBMs, including Optum Rx and Catamaran, who substantially increased its size and market power over the last few years through a series of acquisitions, have been plagued with opaque business practices, limited market competition and widespread allegations of fraud. The facts are clear: while PBMs may well prove a necessary expedient in lowering the cost of healthcare, measures must be taken to ensure that they operate as they are supposed to.

The fundamental elements for a competitive market are transparency, choice and a lack of conflicts of interest. This is especially true when dealing with health care intermediaries such as PBMs and health insurers where information may be difficult to access, there are agency relationships and securing adequate information may be difficult.

Why are choice, transparency, and a lack of conflicts of interest important? It should seem obvious. Consumers need meaningful alternatives to force competitors to vie for their loyalty by offering fair prices and better services. Transparency is necessary for consumers to evaluate products carefully, to make informed choices, and to secure the full range of services they desire.

When dealing with intermediaries, it is particularly critical that there are no conflicts of interest. A PBM is fundamentally acting as a fiduciary to the plan it serves. In the PBM market, the service a PBM is supposed to provide is that of being an “honest broker” bargaining to secure the lowest price for drugs and drug dispensing services. When a PBM has an ownership interest in a drug company or a pharmacy chain, or has its own pharmacy dispensing operations, it is effectively serving two masters.

PBMs entered the health care market as “honest brokers” or intermediaries between health care entities. Health plans and plan sponsors agree to a negotiated fee and contract with PBMs to administer drug claims and serve as a third-party broker with pharmaceutical manufacturers. PBMs can play an important function in health care markets by setting up pharmaceutical benefit networks and adjudicating pharmaceutical claims. However, the role of the PBM has evolved over time and increasingly PBMs have found sources of indirect compensation, and by failing to adequately disclose the compensation (typically from manufacturers), or engaging in misleading disclosures they are able to “play the spread” and pocket the indirect compensation. As a result PBM profits have skyrocketed. Over the last 10 years, the two largest PBMs—Caremark and Express Scripts—nearly more than quadrupled their annual profits from $966 million to over $4 billion. CVS Caremark generated $126.8 billion in revenues in 2013, while Express Scripts

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4 Medco Health Solutions, the largest PBM in the United States, was acquired by Express Scripts in 2012.
generated $104.6 billion in revenues in 2013.\(^5\) CVS Caremark and Express Scripts rank as number 12 and 20, respectively, on the 2014 Fortune 500 list.\(^6\) And both CVS Caremark and Express Scripts’ 2013 revenues exceed that of the largest U.S. drug manufacturer, Johnson and Johnson, by over $30 billion. In addition, there has been tremendous consolidation among PBMs, so the largest PBMs (CVS/Caremark, Express Scripts and OptumRx) now have over 70% of the national PBM market.

Facing weak transparency standards, PBMs frequently engage in a wide range of deceptive and anticompetitive conduct that ultimately harms and denies benefits to consumers. Some PBMs secure rebates and kickbacks in exchange for exclusivity arrangements that may keep lower priced drugs off the market. PBMs may switch patients from prescribed drugs to an often more expensive drug to take advantage of rebates that the PBM receives from drug manufacturers. In addition, PBMs derive their enormous profits from the ability to “play the spread” between pharmaceutical manufacturers, pharmacies, and health care plans. (All of these qualify as undisclosed indirect compensation.) Later in my testimony, I will go into mechanics of these deceptive pricing practices, but it is important to note that these pricing tactics ultimately lead to higher prices paid by plans and patients.

In the past decade, a coalition of over 30 state attorneys general have brought several cases attacking unfair, fraudulent and deceptive conduct. The major PBMs have been the subject of six major federal or multidistrict cases over allegations of fraud; misrepresentation to plan sponsors, patients, and providers; unjust enrichment through secret kickback schemes; and failure to meet ethical and safety standards. These cases listed below, resulted in over $371.9 million in damages to states, plans, and patients so far.

- United States v. AdvancePCS (now part of CVS/Caremark) – $137.5 million in damages for kickbacks, submission of false claims, and other rebate issues.
- State Attorneys General v. Caremark, Inc. – $41 million in damages for deceptive trade practices, drug switching, and repacking.
- State Attorneys General v. Express Scripts – $9.5 million for drug switching and illegally retaining rebates and spread profits and discounts from plans.

And most recently, earlier this year Express Scripts was served with two subpoenas from the attorneys general of New Jersey and Rhode Island concerning its relationship with drug makers who are accused of false claims and kickbacks in marketing of several drugs.\(^7\)

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II. A Lack of Transparency allows PBMs to “play the spread,” leading to higher costs for plan sponsors and patients.

PBMs earn enormous profits by negotiating rebates and discounts with drug manufacturers in exchange for promoting certain drugs on their preferred formulary or engaging in drug substitution programs. PBMs also negotiate contracts with pharmacies to determine how much the pharmacists will be paid for dispensing medication and providing services. By paying a lower reimbursement rate to pharmacies, but failing to adequately disclose reimbursement rates PBMs can generate more revenue. In both respects, PBMs can play the spread by failing to disclose these forms of indirect compensation. The failure to disclose these payments denies purchasers important information that impacts their buying decisions. As a result, this lack of information often results in higher costs for consumers, health plans, employers, and other plan sponsors.

PBMs are free to “play the spread” between manufacturers, pharmacists and plans because of a lack of disclosure. Unclear and inadequate disclosure of rebates and discounts undermine the ability of plan sponsors to compare competing proposals. Because rebates, discounts, and other fee structures remain undisclosed, plan sponsors cannot clearly identify and choose PBMs offering the highest value services. PBMs’ promise of controlling pharmaceutical costs has been undercut by a pattern of conflicts of interest, self-dealing, deception, and anticompetitive conduct. The dominant PBMs have been characterized by opaque business practices, limited market competition, and widespread allegations of fraud.

a. Maximum Allowable Cost is the newest profit center for PBMs

As of recently, with the substantial increase in available generic drugs on the market, the PBMs’ biggest profits no longer lie in maximizing rebates on brand-name drugs or shifting patients to higher-cost medication. Instead, they come from maximizing spreads on generics. Generic prices are typically set through lists of maximum allowable cost (“MAC”), which the PBMs establish. The PBMs may use multiple MAC lists to maximize spread, giving one set of prices to pharmacies and another to clients.

MAC lists are PBM-generated list of products that includes the upper limit or maximum amount that a PBM will pay for generic drugs and brand name drugs that have generic versions available. There is no standard methodology for derivation of MAC lists or how the maximum prices are determined. Neither plan sponsors nor retail pharmacies are informed how products are added or removed from a MAC list or the methodology that determines how this so-called “maximum” cost is calculated or adjusted. Moreover, PBMs often change the “MAC” benchmark, or utilize multiple MAC lists to create a spread between what they charge a plan versus the amount they reimburse a pharmacy. This lack of transparency and prevalence of nonstandard MAC list and pricing derivation allows PBMs to utilize an aggressively low MAC price list to reimburse their contracted pharmacies and a different, higher list of prices when they sell to their clients, plan sponsors. Essentially, the PBMs reimburse low and charge high with their MAC price lists, pocketing the significant spread between the two prices. Most plans are unaware that multiple MAC lists are being used and have no real concept of how much revenue the PBM retains.
This can be additionally problematic from a plan sponsor perspective. The lack of transparency surrounding MAC list derivation causes plans worry that they are paying more than they should for some multisource products. Without the knowledge of whether certain generics are included or excluded on MAC lists, a plan does not know whether a member’s copay may increase due to drugs not being available on MAC lists. A member may complain that they cannot get access to a generic that should be available through their benefit and the plan is forced to pay a higher price to the PBM.

**III. Increased disclosures by PBMs have resulted in price decreases and significant savings for health plans.**

Because of the enforcement activity focusing on PBMs, there has been a great deal of attention surrounding transparency. Transparency is a somewhat ambiguous term, but in this context, David Calabrese in *Managed Care Executive* provides a useful definition:

Transparency is a form of business practice involving full disclosure of costs and revenues, allowing the customer to make more well-informed decisions regarding purchases. In the PBM industry, transparency lays the groundwork for more simplified PBM-client business relations, more accurate financial modeling and performance metrics and a greater comfort level among PBM consumers. 'Transparency,' however, is a relative term used freely in the marketing efforts of many PBMs. The genuine commitment to transparency lies in the actual business practices the PBM invokes to support this claim. 'True transparency' is a model in which all PBM revenue streams [drug-level rebates, funding of clinical programs, administrative fees, service fees, management fees, research/educational grants, etc.] are fully disclosed to the payer; the full value of retail and mail-order pharmacy discounts is passed onto the client; data is shared with the client; and the client is given ultimate decision-making control over its drug benefit design and formulary management. It is this commitment to true transparency which has begun to differentiate newer PBMs.  

Responding to the numerous enforcement actions, both a handful of states and Congress have taken measures to enact transparency provisions by requiring some degree of disclosure of rebates and other revenue. Just last month, the Centers for Medicare and Medicaid Services enacted final rules to the Medicare Part D Program which included the requirement that Part D plans and their PBMs make available to all contracted pharmacies the reimbursement rates for drugs under MAC pricing standards. This requirement will be effective for the 2016 contract year. In addition, in the multistate enforcement action against Caremark, 30 state

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8 Calabrese, David. *Managed Care Executive*. May 1, 2006.
9 At least nine states have enacted MAC transparency statutes, with 15 states considering such legislation in 2014 sessions. Moreover the US Senate has introduced S.B. 867 which required PBMs to disclose certain payment methodologies to pharmacies. See MAC Information Center, available at [www.PBMWATCH.com](http://www.PBMWATCH.com).
10 I will discuss the transparency provisions under PPACA. However, Under the MMA, PBMs that serve Part D Plans are also required to disclose to HHS all manufacturer rebates and price concessions.
attorneys generals required rebate disclosure. Finally, some large sophisticated health plans have negotiated for greater transparency.

The most significant disclosure requirements are incorporated in The Patient Protection and Affordable Care Act of 2010. PPACA works to shine light on spread pricing and undisclosed manufacturer agreements by requiring additional data reporting from PBMs that manage contracts under Medicare Part D or the state Exchanges. These PBMs must provide regulators with data on the percentage of all prescriptions that are provided through retail pharmacies compared to mail-order facilities and the generic dispensing rates for each type. PBMs must also submit the aggregate amounts and types of rebates and discounts or price concessions that the PBM negotiates on behalf of a plan. Importantly, PBMs must disclose how much of these rebates and discounts are “passed through” to the plan versus kept as company profits. In addition to this information, PBMs must also supply regulators with the aggregate difference between the amount paid by the plan and the amount the PBM pays the retail and mail-order pharmacy and number of prescriptions dispensed.

In addition to the disclosure provisions established by PPACA, many plans have recognized the importance of transparency, especially plans that represent government entities. Increasingly very powerful plans are negotiating for transparency and securing significant savings. Large plan sponsors, such as universities, states, and federal programs have recently learned that they can achieve substantial cost savings by opting for contacts with transparent PBMs that disclose negotiations with manufacturers or simply managing their own pharmacy benefit. For example, the University of Michigan has saved nearly $55 million by administering its own plan for the past six years. Similarly, New Jersey projects savings of $558.9 million over six years and Texas expects savings of $265 million by switching to a transparent PBM contract. Instead of managing drug benefits through a traditional PBM, the University of Michigan, New Jersey and Texas are able to engage in a more transparent negotiation process and reduce costs.

In the corporate context, a recent report revealed that Meridian Health System discovered that its drug benefit increased by $1.3 million within the first month of contracting with Express Scripts for PBM services. Meridian discovered that they were being billed for generic amoxicillin at $92.53 for every employee prescription; however Express Scripts was paying only $26.91 to the pharmacy to fill these same prescriptions. The result was a spread, also known as the difference between the PBM’s expenditure and the revenue it takes in, of $65.62. Meridian canceled its contract and switched to a transparent PBM which saved Meridian $2 million in the first year of its contract. Each of these examples demonstrates that disclosure can improve competition and reduce costs to plans and consumers.

Some might suggest that if some states and the federal government are regulating and private parties can negotiate for transparency, that further regulation by DOL is unnecessary. They are mistaken. First, less than a handful of states have implemented full transparency

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12 PPACA, Title VI, Subtitle A, Section 6005.
14 Id.
provisions. Second, the PPACA transparency provisions only apply to plans that are in the state Exchanges and the Medicare Part D program, and the information collected must be retained by the government as “confidential.” Third, the fact that some powerful buyers can negotiate for certain levels of transparency does not mean that transparency regulation is unnecessary. These plans can negotiate for transparency because they have clout; but all plans and their subscribers need the protection of transparency. That is why regulation is necessary.

Some may suggest that disclosure provisions may lead to higher costs. The representatives of the PBM industry argue that transparency would increase costs citing a 2003 CBO report based on a proposed amendment to the Medicare Modernization Act. There are several reasons why that argument should be dismissed. First, the CBO estimate is over 10 years old. Since that time there have been numerous multistate actions demonstrating ongoing fraud and deception. Second, in the Caremark case, over 30 state attorneys generals required transparency as part of their consent order. Third, since that time numerous plans have negotiated for transparency and have achieved significant cost savings. Finally, there is no evidence that any additional transparency from these private plans or state regulation have led to collusion or any other conduct to raise costs. Simply, if transparency was bad, why would Congress enact it, state attorneys generals require it, and plans, especially government plans, work so hard to secure it?

For similar reasons, the PBM industry’s reliance on Federal Trade Commission (“FTC”) studies or advocacy is misguided. For example, the PBM industry relies extensively on hearings conducted by the FTC in 2004 and an outdated 2005 PBM study which suggested that the PBM market was competitive. In addition, they rely on other FTC letters to state regulators on PBM transparency provisions. Much of this advocacy including the hearings preceded the numerous attorneys general enforcement actions which uncovered significant evidence of ongoing fraud and deception involving all of the major PBMs. Moreover, the FTC’s suggestion that some PBM clients may be able to secure accurate information on rebates does not discount the need for regulations to protect all purchasers of PBM services. Indeed the FTC notes that “large, sophisticated repeat-purchasers of health care services” can use useful tools to contract with PBMs. But smaller plans lack these tools and are more vulnerable to deception or conflicts of interest by PBMs.

The issue of whether transparency would lead to higher costs was debated during the enactment of PPACA in 2009 and PBM advocates asked CBO to reaffirm that transparency would lead to significantly higher costs. CBO rejected that position. In 2010, the CBO estimated that PBM transparency standards established by the PPACA would result in zero increased costs. The significant reduction in cost estimates represents that CBO recognizes the potential benefits and unlikely risks of greater transparency. Additionally, if concerns over the risks of disclosure still persist, confidentially provisions can be established to protect the flow of information from PBMs to plans and beneficiaries. The exchange of sensitive information between competitors can be reduced through confidentiality agreements and the disclosure of information to only regulatory agencies instead of market participants. Such is the case with the transparency requirements in the PPACA.

15 See supra, note 9.
Conclusion

Disclosure of PBM compensation and fees should broadly apply to welfare benefit plans, which should be removed from the current exemption in 408(b)(2). PBMs operate with little transparency and engage in deceptive practices such as drug switching and spread pricing. Without transparency, PBM profits will continue to rise exponentially at the expense of plans and patients. Broadening compensation and fee disclosures will allow welfare benefit plan administrators better determine if reasonable compensation is being paid for PBM services and better assess the overall costs of plan administration. Applying mandatory PBM disclosures to the plans will produce substantial savings for plans and decrease patient expenditure on premiums and prescription drugs.

We disagree with those who argue that a PBM should not be obligated to disclose specific information regarding its contracts and arrangements with third parties if the information constitutes a trade secret or if the information is not generally known to the public and affords the PBM a competitive advantage. This exemption would basically negate the value of requiring additional disclosure. PBMs claim that they use protected information such as rebates, discounts, and competitive reimbursement rates to achieve savings for plans. These payments which are sometimes considered “indirect compensation” should be subject to disclosure regulations. The savings experienced by health plans utilizing truly transparent PBM models demonstrate that this information can be disclosed without resulting price increases. Expanded compensation and fee disclosures with limited restrictions will ultimately foster competition and cost control within the PBM market, generally, and to welfare benefit plans, specifically.

We recommend the Council:

a. Include welfare benefit plans under Section 408(b)(2) of ERISA; and

b. Apply PBMs as a covered services under Section 408(b)(2) of ERISA.

The establishment of standards to disclose otherwise undisclosed indirect compensation and opaque fees will help restore PBMs to their role as “honest brokers” and facilitate greater competition in health care markets. Thank you for your time.