1	HOUSE OF REPRESENTATIVES - FLOOR VERSION
2	STATE OF OKLAHOMA
3	1st Session of the 60th Legislature (2025)
4	COMMITTEE SUBSTITUTE FOR ENGROSSED
5	SENATE BILL NO. 993 By: Gollihare and Jech of the Senate
6	and
7	Stinson of the House
8	
9	
10	COMMITTEE SUBSTITUTE
11	An Act relating to pharmacy benefits managers; amending 59 O.S. 2021, Sections 356.1, 356.2, 356.3,
12	as amended by Sections 1, 2, and 3, Chapter 332, O.S.L. 2024, and 356.4 (59 O.S. Supp. 2024, Sections
13	356.1, 356.2, and 356.3), which relate to definitions, pharmacy audit requirements, appeals
14	process, and prohibited extrapolation audit; modifying notice contents; prohibiting assessment of
15	certain fines under certain circumstances; expanding certain claim limits; establishing requirements for
16	preliminary audit findings reports; requiring provision of certain final audit results within a
17	certain time period; updating statutory reference; requiring certain notification to Attorney General in
18	certain circumstances; expanding requirement for initiation of certain audit; lengthening time period
19	for certain preliminary report; allowing certain extension request; shortening certain time period for
20	certain final report; establishing requirements for audit findings report; modifying definition; defining
21	terms; requiring certain tolling in certain declared disaster; providing certain exceptions; amending 59
22	O.S. 2021, Sections 357, 358, and 360, as amended by Sections 4, 5, and 6, Chapter 332, O.S.L. 2024 (59
23	O.S. Supp. 2024, Sections 357, 358, and 360), which relate to definitions, pharmacy benefits management
24	licensure, and pharmacy benefits manager contractual

1	duties; modifying notice contents; defining terms; updating statutory references; requiring certain time
2	period of tolling in certain declared disaster; requiring certain documented proof by certain
3	pharmacy benefits managers; establishing certain denial for certain appeals; prohibiting certain
4	collection of additional monies by certain pharmacy
5	benefits managers; establishing certain filing period after lifting of disaster declaration; prohibiting
6	certain denials; updating statutory language; providing for codification; and declaring an
7	emergency.
8	
9	BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA:
10	SECTION 1. AMENDATORY 59 O.S. 2021, Section 356.1, as
11	amended by Section 1, Chapter 332, O.S.L. 2024 (59 O.S. Supp. 2024,
12	Section 356.1), is amended to read as follows:
13	Section 356.1. A. For purposes of the Pharmacy Audit Integrity
14	Act, "pharmacy benefits manager":
15	1. "Audit" means any review, inspection, or analysis conducted
16	by a pharmacy benefits manager (PBM) or its representative of a
17	pharmacy's records, practices, or compliance with contractual
18	obligations;
19	2. "Disaster declaration" and "declared disaster" mean a
20	declaration issued by the Governor or the President of the United
21	States for an event that qualifies as a disaster including, but not
22	limited to, a flood, tornado, earthquake, wildfire, terrorist
23	attack, or other catastrophic event; and

<u>3. "Pharmacy benefits manager"</u> or "PBM" shall have the same
 meaning as in Section 6960 of Title 36 of the Oklahoma Statutes.

B. The purpose of the Pharmacy Audit Integrity Act is to
establish minimum and uniform standards and criteria for the audit
of pharmacy records by or on behalf of certain entities.

C. The Pharmacy Audit Integrity Act shall apply to any audit of
the records of a pharmacy conducted by a managed care company,
nonprofit hospital, medical service organization, insurance company,
third-party payor, pharmacy benefits manager, a health program
administered by a department of this state, or any entity that
represents these companies, groups, or departments.

D. The Attorney General may promulgate rules to implement theprovisions of the Pharmacy Audit Integrity Act.

14SECTION 2.AMENDATORY59 O.S. 2021, Section 356.2, as15amended by Section 2, Chapter 332, O.S.L. 2024 (59 O.S. Supp. 2024,16Section 356.2), is amended to read as follows:

17 Section 356.2. A. The entity conducting an audit of a pharmacy 18 shall:

Identify and specifically describe the audit and appeal
 procedures in the pharmacy contract. Prescription claim
 documentation and record-keeping recordkeeping requirements shall
 not exceed the requirements set forth by the Oklahoma Pharmacy Act
 or other applicable state or federal laws or regulations;

1 2. Give the pharmacy written notice by certified letter to the 2 pharmacy and the pharmacy's contracting agent, including identification of specific prescription numbers and, fill dates, 3 drug names, and National Drug Code (NDC) numbers to be audited, at 4 5 least fourteen (14) calendar days prior to conducting the audit, including, but not limited to, an on-site audit, a desk audit, or a 6 wholesale purchase audit, request for documentation related to the 7 dispensing of a prescription drug, or any reimbursed activity by a 8 9 pharmacy provider; provided, however, that wholesale purchase audits shall require a minimum of thirty (30) calendar days' written 10 notice. For an on-site audit, the audit date shall be the date the 11 12 on-site audit occurs. For all other audit types, the audit date shall be the date the pharmacy provides the documentation requested 13 in the audit notice. The pharmacy shall have the opportunity to 14 reschedule the audit no more than seven (7) calendar days from the 15 date designated on the original audit notification; 16

Not interfere with the delivery of pharmacist services to a
 patient and shall utilize every reasonable effort to minimize
 inconvenience and disruption to pharmacy operations during the audit
 process;

4. Conduct any audit involving clinical or professional
 judgment by means of or in consultation with a licensed pharmacist;
 5. Not consider as fraud any clerical or record-keeping
 recordkeeping error, such as a typographical error, scrivener's

1 error or computer error, including, but not limited to, a 2 miscalculated day supply, incorrectly billed prescription written date or prescription origin code, and such errors shall not be 3 subject to recoupment. The pharmacy shall have the right to submit 4 5 amended claims electronically to correct clerical or record-keeping recordkeeping errors in lieu of recoupment. To the extent that an 6 audit results in the identification of any clerical or record-7 keeping recordkeeping errors such as typographical errors, 8 9 scrivener's errors or computer errors in a required document or 10 record, the pharmacy shall not be subject to recoupment of funds by the pharmacy benefits manager unless the pharmacy benefits manager 11 can provide proof of intent to commit fraud. A person shall not be 12 13 subject to criminal penalties for errors provided for in this paragraph without proof of intent to commit fraud; 14

6. Permit a pharmacy to use the records of a hospital,
physician, or other authorized practitioner of the healing arts for
drugs or medicinal supplies written or transmitted by any means of
communication for purposes of validating the pharmacy record with
respect to orders or refills of a legend or narcotic drug;

20 7. Not include the dispensing fee amount or the actual invoice 21 cost of the prescription dispensed in a finding of an audit 22 recoupment unless a prescription was not actually dispensed or a 23 physician denied authorization of a dispensing order;

8. Audit each pharmacy under identical standards, regularity
 and parameters as other similarly situated pharmacies and all
 pharmacies owned or managed by the pharmacy benefits manager
 conducting or having conducted the audit;

9. Not exceed one (1) year from the date the claim was
submitted to or adjudicated by a managed care company, nonprofit
hospital or medical service organization, insurance company, thirdparty payor, pharmacy benefits manager, a health program
administered by a department of this state, or any entity that
represents the companies, groups, or departments for the period
covered by an audit;

12 10. Not schedule or initiate an audit during the first seven 13 (7) calendar days of any month unless otherwise consented to by the 14 pharmacy;

15 11. Disclose to any plan sponsor whose claims were included in 16 the audit any money recouped in the audit;

17 12. Not require pharmacists to break open packaging labeled 18 "for single-patient-use only". Packaging labeled "for single-19 patient-use only" shall be deemed to be the smallest package size 20 available; and

21 13. Upon recoupment of funds from a pharmacy, refund first to 22 the patient the portion of the recovered funds that were originally 23 paid by the patient, provided such funds were part of the

24 recoupment; and

Page 6

<u>14. Not assess a fine, penalty, or any other financial</u>
 <u>requirement on the pharmacy or pharmacist for any prescription</u>
 <u>audited unless there is a valid recoupment under the Pharmacy Audit</u>
 Integrity Act.

5 B. 1. Any entity that conducts wholesale purchase review 6 during an audit of a pharmacist or pharmacy shall not require the 7 pharmacist or pharmacy to provide a full dispensing report. 8 Wholesaler invoice reviews shall be limited to verification of 9 purchase inventory specific to the pharmacy claims paid by the 10 health benefits plan or pharmacy benefits manager conducting the 11 audit.

12 2. Any entity conducting an audit shall not identify or label a13 prescription claim as an audit discrepancy when:

- a. the National Drug Code for the dispensed drug is in a
 quantity that is a subunit or multiple of the drug
 purchased by the pharmacist or pharmacy as supported
 by a wholesale invoice,
- b. the pharmacist or pharmacy dispensed the correct
 quantity of the drug according to the prescription,
 and
- c. the drug dispensed by the pharmacist or pharmacy
 shares all but the last two digits of the National
 Drug Code of the drug reflected on the supplier
 invoice.

3. An entity conducting an audit shall accept as evidence,
 subject to validation, to support the validity of a pharmacy claim
 related to a dispensed drug:

a. redacted copies of supplier invoices in the
pharmacist's or pharmacy's possession, or
b. invoices and any supporting documents from any
supplier as authorized by federal or state law to
transfer ownership of the drug acquired by the
pharmacist or pharmacy.

10 4. An entity conducting an audit shall provide, no later than 11 five (5) calendar days after the date of a request by the pharmacist 12 or pharmacy, all supporting documents the pharmacist's or pharmacy's 13 purchase suppliers provided to the health benefits plan issuer or 14 pharmacy benefits manager.

15 C. A pharmacy shall be allowed to provide the pharmacy's 16 computerized patterned medical records or the records of a hospital, 17 physician, or other authorized practitioner of the healing arts for 18 drugs or medicinal supplies written or transmitted by any means of 19 communication for purposes of supporting the pharmacy record with 20 respect to orders or refills of a legend or narcotic drug.

D. The entity conducting the audit shall not audit more than
fifty prescriptions, with specific date of service, per calendar
year PBM or its agent shall not exceed an annual limit of fifty
prescription claims with a specific prescription number and date of

fill per calendar year. The annual limit to the number of prescription claims audited shall be inclusive of all audits by a <u>PBM or its agent</u>, including any prescription-related documentation requests from the health insurer, pharmacy benefits manager or any third-party company conducting audits on behalf of any health insurer or pharmacy benefits manager during a calendar year.

E. If paper copies of records are requested by the entity
conducting the audit, the entity shall pay twenty-five cents (\$0.25)
per page to cover the costs incurred by the pharmacy. The entity
conducting the audit shall provide the pharmacy with accurate
instructions, including any required form for obtaining
reimbursement for the copied records.

13 F. The entity conducting the audit shall:

Deliver a preliminary audit findings report to the pharmacy
 and the pharmacy's contracting agent within forty-five (45) calendar
 days of conducting the audit;

17 2. Allow the pharmacy at least ninety (90) calendar days 18 following receipt of the preliminary audit findings report in which 19 to produce documentation to address any discrepancy found during the 20 audit; provided, however, a pharmacy may request an extension, not 21 to exceed an additional forty-five (45) calendar days;

3. Deliver a final audit findings report to the pharmacy and the pharmacy's contracting agent signed by the auditor within ten (10) calendar days after receipt of additional documentation 1 provided by the pharmacy, as provided for in Section 356.3 of this
2 title;

4. Allow the pharmacy to reverse and resubmit claims
electronically within thirty (30) calendar days of receipt of the
final audit report in lieu of the auditing entity recouping
discrepant claim amounts from the pharmacy;

5. Not recoup any disputed funds until after final disposition
of the audit findings, including the appeals process as provided for
in Section 356.3 of this title; and

10 6. Not accrue interest during the audit and appeal period;

11 <u>7. Ensure that each preliminary audit findings report required</u> 12 <u>by this section includes:</u>

a. specific prescription numbers, fill dates, drug names,
 and NDC numbers, and

15 b. the date of receipt of documents from the pharmacy,

16 the pharmacy's contracting agent, or any other source 17 associated with the audit.

18 G. Each entity conducting an audit shall provide a copy of the

19 final audit results, and a final audit report upon request, after

20 completion of any review process to the plan sponsor

21 <u>In addition to the requirements for a preliminary audit findings</u>
22 <u>report in this paragraph, the final audit findings report shall</u>

23 include any additional documentation that was submitted to the

24 auditing entity;

1 <u>8. Provide the plan sponsor a copy of the final audit results</u>
2 within thirty (30) calendar days of the final disposition of the
3 <u>audit; and</u>

<u>9. At the request of the plan sponsor, provide a copy of the</u>
<u>final audit findings report within thirty (30) calendar days of the</u>
request.

H. G. 1. The full amount of any recoupment on an audit shall
be refunded to the plan sponsor. Except as provided for in
paragraph 2 of this subsection, a charge or assessment for an audit
shall not be based, directly or indirectly, on amounts recouped.

11 2. This subsection does not prevent the entity conducting the 12 audit from charging or assessing the responsible party, directly or 13 indirectly, based on amounts recouped if both of the following 14 conditions are met:

a. the plan sponsor and the entity conducting the audit
have a contract that explicitly states the percentage
charge or assessment to the plan sponsor, and
a commission to an agent or employee of the entity
conducting the audit is not based, directly or
indirectly, on amounts recouped.

21 I. H. Unless superseded by state or federal law, auditors shall 22 only have access to previous audit reports on a particular pharmacy 23 conducted by the auditing entity for the same pharmacy benefits 24 manager, health plan or insurer. An auditing vendor contracting with multiple pharmacy benefits managers or health insurance plans shall not use audit reports or other information gained from an audit on a pharmacy to conduct another audit for a different pharmacy benefits manager or health insurance plan.

5

J. Sections A through I

I. Paragraph 2 of subsection A of this section through
subsection D of this section, and paragraph 1 through paragraph 7 of
<u>subsection F</u> of this section shall not apply to any audit initiated
based on or that involves suspicion of fraud, willful

10 misrepresentation, or abuse.

11 K. J. If the Attorney General, after notice and opportunity for 12 hearing, finds that the entity conducting the audit failed to follow any of the requirements pursuant to the Pharmacy Audit Integrity 13 Act, the audit shall be considered null and void. Any monies 14 recouped from a null and void audit shall be returned to the 15 affected pharmacy within fourteen (14) calendar days. Any violation 16 of this section by a pharmacy benefits manager or auditing entity 17 shall be deemed a violation of the Pharmacy Audit Integrity Act. 18 SECTION 3. AMENDATORY 59 O.S. 2021, Section 356.3, as 19 amended by Section 3, Chapter 332, O.S.L. 2024 (59 O.S. Supp. 2024, 20 Section 356.3), is amended to read as follows: 21 Section 356.3. A. Each entity conducting an audit shall 22

23 establish a written appeals process under which a pharmacy may

1 appeal an unfavorable preliminary audit report and/or final audit
2 report to the entity.

B. Following an appeal, if the entity finds that an unfavorable audit report or any portion thereof is unsubstantiated, the entity shall dismiss the audit report or the unsubstantiated portion of the audit report without any further action.

Any final audit report, following the final audit appeal 7 С. period, with a finding of fraud or willful misrepresentation shall 8 9 be referred to the district attorney having proper jurisdiction or 10 the Attorney General for prosecution upon completion of the appeals process. If a finding of fraud or willful misrepresentation is 11 12 referred to a district attorney under this subsection, the auditing 13 entity shall notify the Attorney General as to whom the referral was made and the date the referral was made. 14

D. For any audit initiated based on or that involves suspicion 15 of fraud, willful misrepresentation, or abuse, the auditing entity 16 shall provide, in writing, at the time of the audit, a clear and 17 conspicuous declaration to the pharmacy being audited that the audit 18 is being conducted under suspicion of fraud, willful 19 misrepresentation, or abuse and a statement of facts that supports 20 the reasonable suspicion. The entity conducting an audit based on 21 suspicion of fraud, willful misrepresentation, or abuse shall 22 provide a copy of the clear and conspicuous declaration required by 23 this subsection to the pharmacy's contracting agent by certified 24

1	mail within five (5) business days of notifying the pharmacy of an
2	audit pursuant to this section.
3	E. The entity conducting an audit based on suspicion of fraud,
4	willful misrepresentation, or abuse shall:
5	1. Deliver a preliminary findings report to the pharmacy and
6	the pharmacy's contracting agent within ninety (90) calendar days of
7	notification of the audit;
8	2. Allow the pharmacy at least ninety (90) calendar days
9	following the receipt of the preliminary audit findings report in
10	which to produce documentation to address any discrepancy found
11	during the audit. A pharmacy may request an extension, not to
12	exceed an additional forty-five (45) calendar days;
13	3. Deliver a final audit findings report to the pharmacy and
14	the pharmacy's contracting agent signed by the auditor within thirty
15	(30) calendar days after receipt of additional documentation
16	provided by the pharmacy;
17	4. Allow the pharmacy to reverse and resubmit claims
18	electronically within thirty (30) calendar days of receipt of the
19	final audit report in lieu of the auditing entity recouping
20	discrepant claim amounts from the pharmacy;
21	5. Not recoup any disputed funds until after the final
22	disposition of the audit findings, including the appeals process
23	pursuant to this section;
24	6. Not accrue interest during the audit and appeal period;

1	7. Ensure that each preliminary audit findings report submitted
2	pursuant to this section includes:
3	a. specific prescription numbers, fill dates, drug names,
4	and NDC numbers, and
5	b. the date of receipt of documents from the pharmacy,
6	the pharmacy's contracting agent, or any other source
7	associated with the audit;
8	8. Ensure that each final audit findings report includes any
9	additional documentation that was submitted to the auditing entity;
10	9. Provide the plan sponsor a copy of the final audit results
11	within thirty (30) calendar days of the final disposition of the
12	audit; and
13	10. At the request of the plan sponsor, provide a copy of the
14	final audit report within thirty (30) calendar days of the request.
15	<u>F.</u> Any entity conducting an audit that is based on or involves
16	suspicion of fraud, willful misrepresentation, or abuse shall
17	provide to the Office of the Attorney General:
18	1. Notice at least two (2) calendar days prior to beginning
19	performance of an audit pursuant to this section;
20	2. A preliminary report within thirty (30) calendar days of
21	performing the audit five (5) business days of providing a copy of
22	the preliminary report to the pharmacy and the pharmacy's
23	contracting agent pursuant to this section. The auditing entity may
24	

1 request an extension from the Attorney General, not to exceed an 2 additional ninety (90) calendar days; and

3 3. A final report within thirty (30) ten (10) calendar days 4 following the closure of the final appeal period for an audit 5 performed pursuant to this section.

- The final report for the Office of the Attorney 6 a. General shall include the name of each plan sponsor 7 whose claims were included in the audit recover, the 8 9 amount of funds recouped on behalf of the plan, the 10 date the plan sponsor was notified of the recoupment, 11 the date the plan sponsor was paid any recoupment, and 12 the name and contact information for the 13 representative of the plan sponsor who was notified of the recoupment at issue in an audit pursuant to this 14 15 section. The auditing entity may request an extension from the 16 b. Attorney General, not to exceed an additional ten (10) 17
- 18

calendar days.

19 F. G. The Attorney General, authorized employees, and examiners 20 shall have access to any pharmacy benefits manager's files and 21 records that may relate to an <u>any</u> audit <u>including</u>, <u>but not limited</u> 22 <u>to, an audit</u> that is based on or involves <u>suspicion of</u> fraud, 23 willful misrepresentation, or abuse.

G. H. The Attorney General may levy a civil or administrative
 fine of not less than One Hundred Dollars (\$100.00) and not greater
 than Ten Thousand Dollars (\$10,000.00) for each violation of this
 section and assess any other penalty or remedy authorized by law.
 SECTION 4. AMENDATORY 59 O.S. 2021, Section 356.4, is
 amended to read as follows:

7 Section 356.4. A. For the purposes of the Pharmacy Audit Integrity Act, "extrapolation audit" means an audit of a sample of 8 9 prescription drug benefit claims submitted by a pharmacy to the 10 entity conducting the audit that is then used to estimate audit 11 results for a larger batch or group of claims not reviewed by the 12 auditor, including refills not listed in the written notification in accordance with paragraph 2 of subsection A of Section 356.2 of this 13 14 title.

B. The entity conducting the audit shall not use the accounting practice of extrapolation in calculating recoupments or penalties for audits.

18 SECTION 5. NEW LAW A new section of law to be codified 19 in the Oklahoma Statutes as Section 356.6 of Title 59, unless there 20 is created a duplication in numbering, reads as follows:

A. Notwithstanding any other provision of law, the ability of a pharmacy benefits manager (PBM) to initiate, continue, or conclude an audit of a pharmacy shall be tolled for the duration of a

declared disaster and for an additional period of thirty (30)
 calendar days following the termination of a declared disaster.

Such requirement shall apply only to the pharmacies located within the geographical boundaries of the county or counties affected by the declared disaster.

B. The provisions of this section shall apply to all PBMs
operating within this state, and to all audits conducted pursuant to
contracts between PBMs and pharmacies.

9 C. This section shall not apply to:

Audits conducted for suspected fraudulent activity if
 documented evidence of such activity exists; or

12 2. Audits required to comply with federal or state law 13 unrelated to the contractual relationship between a PBM and a 14 pharmacy.

D. Nothing in this section shall be construed to prohibit a pharmacy from voluntarily agreeing to continue or complete an audit during the tolling period, provided such agreement is documented in writing and signed by both parties.

E. A PBM may submit a request to the Attorney General to
continue or complete an audit during the tolling period, which the
Attorney General may grant at his or her sole discretion. Any PBM
granted such permission by the Attorney General shall do so pursuant
to the requirements of this act.

SECTION 6. AMENDATORY 59 O.S. 2021, Section 357, as
 amended by Section 4, Chapter 332, O.S.L. 2024 (59 O.S. Supp. 2024,
 Section 357), is amended to read as follows:

4 Section 357. A. As used in Sections 357 through 360 of this 5 title and Section 9 of this act:

1. "Covered entity" means a nonprofit hospital or medical 6 service organization, for-profit hospital or medical service 7 organization, insurer, health benefit plan, health maintenance 8 9 organization, health program administered by the state in the 10 capacity of providing health coverage, or an employer, labor union, or other group of persons that provides health coverage to persons 11 in this state. This term does not include a health benefit plan 12 that provides coverage only for accidental injury, specified 13 disease, hospital indemnity, disability income, or other limited 14 benefit health insurance policies and contracts that do not include 15 prescription drug coverage; 16

17 2. "Covered individual" means a member, participant, enrollee, 18 contract holder or policy holder or beneficiary of a covered entity 19 who is provided health coverage by the covered entity. A covered 20 individual includes any dependent or other person provided health 21 coverage through a policy, contract or plan for a covered 22 individual;

23 3. "Department" means the Insurance Department;

4. "Maximum allowable cost", "MAC", or "MAC list" means the
 list of drug products delineating the maximum per-unit reimbursement
 for multiple-source prescription drugs, medical product, or device;

5. "Multisource drug product reimbursement" (reimbursement)
means the total amount paid to a pharmacy inclusive of any reduction
in payment to the pharmacy, excluding prescription dispense fees;

7 6. "Office" means the Office of the Attorney General;

8 7. "Pharmacy benefits management" means a service provided to 9 covered entities to facilitate the provision of prescription drug 10 benefits to covered individuals within the state, including 11 negotiating pricing and other terms with drug manufacturers and 12 providers. Pharmacy benefits management may include any or all of 13 the following services:

a. claims processing, retail network management and
 payment of claims to pharmacies for prescription drugs
 dispensed to covered individuals,

b. clinical formulary development and management
 services, or

19 c. rebate contracting and administration;

8. "Pharmacy benefits manager" or "PBM" means a person,
 business, or other entity that performs pharmacy benefits
 management. The term shall include <u>any business or entity licensed</u>
 <u>by the Insurance Department to perform PBM services, or</u> a person or
 entity acting on behalf of a PBM in a contractual or employment

1 relationship in the performance of pharmacy benefits management for a managed care company, nonprofit hospital, medical service 2 organization, insurance company, third-party payor, or a health 3 program administered by an agency or department of this state; 4 5 9. "Plan sponsor" means the employers, insurance companies, unions and health maintenance organizations or any other entity 6 responsible for establishing, maintaining, or administering a health 7 benefit plan on behalf of covered individuals; and 8

9 10. "Provider" means a pharmacy licensed by the State Board of
10 Pharmacy, or an agent or representative of a pharmacy, including,
11 but not limited to, the pharmacy's contracting agent, which
12 dispenses prescription drugs or devices to covered individuals.

Β. Nothing in the definition of pharmacy benefits management or 13 pharmacy benefits manager in the Patient's Right to Pharmacy Choice 14 Act, Pharmacy Audit Integrity Act, or Sections 357 through 360 of 15 this title, or Section 9 of this act shall deem an employer a 16 "pharmacy benefits manager" pharmacy benefits manager of its own 17 self-funded health benefit plan, except, to the extent permitted by 18 applicable law, where the employer, without the utilization of a 19 third party and unrelated to the employer's own pharmacy: 20

21

a. negotiates

22 <u>1. Negotiates</u> directly with drug manufacturers,

- 23 b. processes
- 24 2. Processes claims on behalf of its members τ ; or

1

c. manages

2 3. Manages its own retail network of pharmacies.

3 SECTION 7. AMENDATORY 59 O.S. 2021, Section 358, as 4 amended by Section 5, Chapter 332, O.S.L. 2024 (59 O.S. Supp. 2024, 5 Section 358), is amended to read as follows:

6 Section 358. A. In order to provide pharmacy benefits 7 management or any of the services included under the definition of 8 pharmacy benefits management in this state, a pharmacy benefits 9 manager or any entity acting as one in a contractual or employment 10 relationship for a covered entity shall first obtain a license from 11 the Insurance Department, and the Department may charge a fee for 12 such licensure.

The Department shall establish, by regulation, licensure 13 Β. procedures, required disclosures for pharmacy benefits managers 14 (PBMs) and other rules as may be necessary for carrying out and 15 enforcing the provisions of this title. The licensure procedures 16 17 shall, at a minimum, include the completion of an application form that shall include the name and address of an agent for service of 18 process, the payment of a requisite fee, and evidence of the 19 procurement of a surety bond. 20

C. The Department or the Office of the Attorney General may
subpoena witnesses and information. Its compliance officers may
take and copy records for investigative use and prosecutions.
Nothing in this subsection shall limit the Office of the Attorney

General from using its investigative demand authority to investigate
 and prosecute violations of the law.

The Department may suspend, revoke or refuse to issue or 3 D. renew a license for noncompliance with any of the provisions hereby 4 5 established or with the rules promulgated by the Department; for conduct likely to mislead, deceive or defraud the public or the 6 Department; for unfair or deceptive business practices or for 7 nonpayment of an application or renewal fee or fine. The Department 8 9 may also levy administrative fines for each count of which a PBM has 10 been convicted in a Department hearing.

The Office of the Attorney General, after notice and 11 Ε. 1. 12 opportunity for hearing, may instruct the Insurance Commissioner that the PBM's license be censured, suspended, or revoked for 13 conduct likely to mislead, deceive, or defraud the public or the 14 State of Oklahoma; or for unfair or deceptive business practices, or 15 for any violation of the Patient's Right to Pharmacy Choice Act, the 16 Pharmacy Audit Integrity Act, or Sections 357 through 360 of this 17 title, or Section 9 of this act. The Office of the Attorney General 18 may also levy administrative fines for each count of which a PBM has 19 been convicted following a hearing before the Attorney General. 20 Ιf the Attorney General makes such instruction, the Commissioner shall 21 enforce the instructed action within thirty (30) calendar days. 22

23 2. In addition to or in lieu of any censure, suspension, or24 revocation of a license by the Commissioner, the Attorney General

may levy a civil or administrative fine of not less than One Hundred Dollars (\$100.00) and not greater than Ten Thousand Dollars (\$10,000.00) for each violation of this subsection and/or assess any other penalty or remedy authorized by this section. For purposes of this section, each day a PBM fails to comply with an investigation or inquiry may be considered a separate violation.

F. The Attorney General may promulgate rules to implement the
provisions of Sections 357 through 360 of this title and Section 9
of this act.

10 SECTION 8. AMENDATORY 59 O.S. 2021, Section 360, as 11 amended by Section 6, Chapter 332, O.S.L. 2024 (59 O.S. Supp. 2024, 12 Section 360), is amended to read as follows:

Section 360. A. The pharmacy benefits manager shall, with respect to contracts between a pharmacy benefits manager and a provider, including a pharmacy service administrative organization:

Include in such contracts the specific sources utilized to
 determine the maximum allowable cost (MAC) pricing of the pharmacy,
 update MAC pricing at least every seven (7) calendar days, and
 establish a process for providers to readily access the MAC list
 specific to that provider;

21 2. In order to place a drug on the MAC list, ensure that the 22 drug is listed as "A" or "B" rated in the most recent version of the 23 FDA's United States Food and Drug Administration (FDA) Approved Drug 24 Products with Therapeutic Equivalence Evaluations, also known as the Orange Book, and the drug is generally available for purchase by
 pharmacies in the state from national or regional wholesalers and is
 not obsolete;

4 3. Ensure dispensing fees are not included in the calculation5 of MAC price reimbursement to pharmacy providers;

6 4. Provide a reasonable administration appeals procedure to allow a provider, a provider's representative and a pharmacy service 7 administrative organization to contest reimbursement amounts within 8 9 fourteen (14) calendar days of the final adjusted payment date. The 10 pharmacy benefits manager shall not prevent the pharmacy or the pharmacy service administrative organization from filing 11 12 reimbursement appeals in an electronic batch format. The pharmacy 13 benefits manager must respond to a provider, a provider's representative and a pharmacy service administrative organization 14 who have contested a reimbursement amount through this procedure 15 within ten (10) calendar days. The pharmacy benefits manager must 16 respond in an electronic batch format to reimbursement appeals filed 17 in an electronic batch format. The pharmacy benefits manager shall 18 not require a pharmacy or pharmacy services administrative 19 organization to log into a system to upload individual claim appeals 20 or to download individual appeal responses. If a price update is 21 warranted, the pharmacy benefits manager shall make the change in 22 the reimbursement amount, permit the dispensing pharmacy to reverse 23

and rebill the claim in question, and make the reimbursement amount
 change retroactive and effective for all contracted providers; and

3 5. If a below-cost reimbursement appeal is denied, the PBM shall provide the reason for the denial, including the National Drug 4 5 Code (NDC) number from, and the name of, the specific national or regional wholesalers doing business in this state where the drug is 6 currently in stock and available for purchase by the dispensing 7 pharmacy at a price below the PBM's reimbursement price. 8 The PBM 9 shall include documented proof from the specific national or 10 regional wholesalers doing business in this state showing that the 11 drug is currently in stock and available for purchase by the 12 dispensing pharmacy at a price below the PBM's reimbursement price. If the NDC number provided by the pharmacy benefits manager is not 13 available below the acquisition cost obtained from the 14 pharmaceutical wholesaler from whom the dispensing pharmacy 15 purchases the majority of the prescription drugs that are dispensed, 16 the pharmacy benefits manager shall immediately adjust the 17 reimbursement amount, permit the dispensing pharmacy to reverse and 18 rebill the claim in question, and make the reimbursement amount 19 20 adjustment retroactive and effective for all contracted providers; 6. Any appeal that results in an increase in the reimbursement 21 from the PBM that continues to be below the pharmacy's acquisition 22 cost shall be considered a denial under this section. Any denial of 23

1 an appeal shall follow the requirements of paragraph 5 of this
2 subsection; and

The PBM shall not require a pharmacy to collect additional 3 7. 4 monies following a successful below-cost reimbursement appeal from 5 any person or entity other than the PBM who adjudicated the drug claim, including the patient or plan sponsor. 6 The reimbursement appeal requirements in this section shall 7 в. apply to all drugs, medical products, or devices reimbursed 8 9 according to any payment methodology, including, but not limited to: 10 1. Average acquisition cost, including the National Average Drug Acquisition Cost; 11 12 2. Average manufacturer price; 3. Average wholesale price; 13 Brand effective rate or generic effective rate; 4. 14 Discount indexing; 15 5. Federal upper limits; 16 6. 7. Wholesale acquisition cost; and 17 Any other term that a pharmacy benefits manager or an 18 8. insurer of a health benefit plan may use to establish reimbursement 19 rates to a pharmacist or pharmacy for pharmacist services. 20 С. The pharmacy benefits manager shall not place a drug on a 21 MAC list, unless there are at least two therapeutically equivalent, 22 multiple-source drugs, generally available for purchase by 23 dispensing retail pharmacies from national or regional wholesalers. 24

D. In the event that a drug is placed on the FDA Drug Shortages
 Database, pharmacy benefits managers shall reimburse claims to
 pharmacies at no less than the wholesale acquisition cost for the
 specific NDC number being dispensed.

E. The pharmacy benefits manager shall not require
accreditation or licensing of providers, or any entity licensed or
regulated by the State Board of Pharmacy, other than by the State
Board of Pharmacy or federal government entity as a condition for
participation as a network provider.

F. A pharmacy or pharmacist may decline to provide the pharmacist clinical or dispensing services to a patient or pharmacy benefits manager if the pharmacy or pharmacist is to be paid less than the pharmacy's cost for providing the pharmacist clinical or dispensing services.

G. The pharmacy benefits manager shall provide a dedicated telephone number, email address and names of the personnel with decision-making authority regarding MAC appeals and pricing.

18 SECTION 9. NEW LAW A new section of law to be codified 19 in the Oklahoma Statutes as Section 360.1 of Title 59, unless there 20 is created a duplication in numbering, reads as follows:

A. If a disaster declaration is issued for a county in this
state, the time period for a provider, a provider's representative,
or a pharmacy service administrative organization to file a belowcost reimbursement appeal pursuant to Section 360 of Title 59 of the

Oklahoma Statutes shall be tolled for the duration of the disaster
 declaration.

B. Upon the expiration of the disaster declaration, the tolling
of the filing period for below-cost reimbursement appeals shall
continue for an additional thirty (30) calendar days. Afterward,
the time period for filing a below-cost reimbursement appeal, as
otherwise provided under state law, shall resume.

8 C. The tolling provisions of this section shall apply only to 9 continuing counties included in the declared disaster area and to 10 below-cost reimbursement appeals arising from claims impacted during 11 the time period of the declared disaster.

D. A pharmacy benefits manager (PBM) shall not deny a belowcost reimbursement appeal on timeliness if such appeal is filed during the tolled period provided in this section.

15 E. The Attorney General may promulgate rules to implement the16 provisions of this act.

17 SECTION 10. It being immediately necessary for the preservation 18 of the public peace, health or safety, an emergency is hereby 19 declared to exist, by reason whereof this act shall take effect and 20 be in full force from and after its passage and approval.

21

22 COMMITTEE REPORT BY: COMMITTEE ON HEALTH AND HUMAN SERVICES OVERSIGHT, dated 04/16/2025 - DO PASS, As Amended.

23

24

SB993 HFLR BOLD FACE denotes Committee Amendments.