TOC not required

292.910 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date		
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000		
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000		
00004	GENENTECH, INC.	01/01/1991	01/01/3000		
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000		
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000		
00008	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000		
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000		
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000		
00014	PFIZER, INC	01/01/1991	01/01/3000		
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000		
00023	ALLERGAN INC	01/01/1991	01/01/3000		
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000		
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000		
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000		
00032	ABBVIE INC.	01/01/1991	01/01/3000		

LABELER CODE	LABELER NAME	EFFECTIVE DATE	TERMINATION DATE
00002	ELI LILLY AND COMPANY	1/1/1991	DATE
00003	E.R. SQUIBB & SONS, INC	1/1/1991	
00004	HOFFMANN-LA ROCHE	1/1/1991	
00005	LEDERLE LABORATORIES	1/1/1991	
00006	MERCK & CO., INC.	1/1/1991	
00007	GLAXOSMITHKLINE	1/1/1991	
00008	WYETH LABORATORIES	1/1/1991	
00009	PFIZER, INC.	1/1/1991	
00011	BECTON DICKINSON MICROBIOLOGY SYSTEMS	10/1/1991	7/1/1998
00013	PFIZER, INC.	1/1/1991	

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

Diagram 2

00123	0456	78
LABELER	PRODUCT	PACKAGE
CODE	CODE	CODE
(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

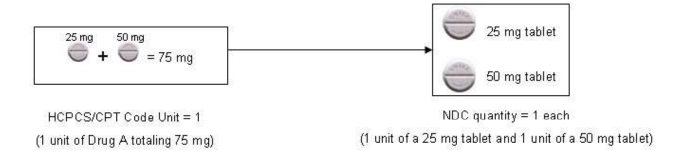
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

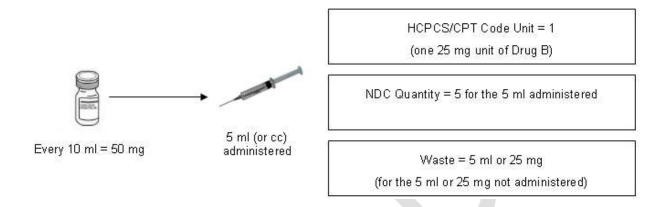
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters, or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



- A. Electronic Claims Filing 837P (Professional) and 837I (Outpatient)
 - Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid requires providers using electronic claims filing through the provider portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500
- Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.

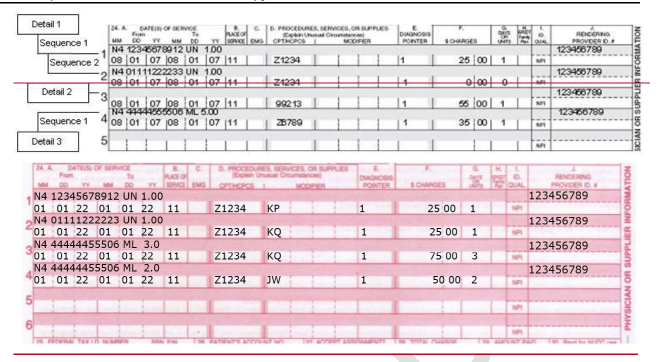
- See Diagram 6 for CMS-1500.
- For professional claims, CMS-1500, list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2 International Unit; GR Gram; ML Milliliter; UN Unit), and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.
- Each NDC when billed <u>under the same procedure code on the same date of service</u> is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.
- The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

-Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage



Procedure Code/NDC Detail Attachment Form - DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Diagram 7

Detail #	Sequence #		NDC										Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

I<u>II</u>V. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations, or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include

documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing purchase of drugs and documentation showing what drug (name, strength, and amount) was administered and on what date, to the beneficiary in question.

See Section 292.950 for additional information regarding drug code billing.

292.950 Injections, Therapeutic and/or Diagnostic Agents

2-1-221-1 23

A. Providers billing the Arkansas Medicaid Program for covered injections should bill the appropriate CPT or HCPCS procedure code for the specific injection administered. The procedure codes and their descriptions may be found in the Current Procedure Terminology (CPT) and in the Healthcare Common Procedural Coding System Level II (HCPCS) coding books.

Injection administration code, is payable for beneficiaries of all ages. May be used for billing the administration of subcutaneous and/or intramuscular injections only. This procedure code cannot be billed when the medication is administered "ORALLY." No fee is billable for drugs administered orally.

Cannot be billed separately for Influenza Virus vaccines or Vaccines for Children (VFC) vaccines.

Cannot be billed to administer any medication given for family planning purposes. No other fee is billable when the provider decides not to supply family planning injectable medications.

Cannot be billed when the drug administered is not FDA approved.

See the table below when billing:

<u>View or print the procedure codes for Physician/Independent Lab/CRNA/Radiation</u> Therapy Center services.

Most of the cCovered drugs can be billed electronically or on paper. However, any covered drug marked with an asterisk (*) must be billed on paper with the name of the drug and dosage listed in the "Procedures, Services, or Supplies" column, Field 24D, of the CMS-1500 claim form. View a CMS-1500 sample form. If requested, additional documentation may be required to justify medical necessity. Reimbursement for manually priced drugs is based on a percentage of the average wholesale price.

See Section 292.940 for coverage information of radiopharmaceutical procedure codes.

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs. See Section 292.910 for further information.

Administration of therapeutic agents is payable only if provided in a physician's office, place of service code "11." These procedures are not payable to the physician if performed in any other setting. Therapeutic injections should only be provided by physicians experienced in the provision of these medications and who have the facilities to treat patients who may experience adverse reactions. The capability to treat infusion reactions with appropriate life support techniques should be immediately available. Only one administration fee is allowed per date of service unless "multiple sites" are indicated in the "Procedures, Services, or Supplies" field in the CMS-1500 claim form. Reimbursement for supplies is included in the administration fee. An administration fee is not allowed when drugs are given orally.

Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges for therapeutic and chemotherapy administration procedure codes.

See Section 292.940 for radiopharmaceutical drugs.

- B. For consideration of payable unlisted CPT/HCPCS drug procedure codes:
 - 1. The provider must submit an electronic or paper claim that includes a description of the drug being represented by the unlisted procedure code on the claim form.
 - 2. Documentation that further describes the drug provided must be attached and must include justification for medical necessity.
 - 3. All other billing requirements must be met in order for payment to be approved.

C. Immunizations

Physicians may bill for immunization procedures on the CMS-1500 claim form. <u>View a</u> **CMS-1500 sample form.** See Section 292.950 for covered vaccines and billing protocols.

Coverage criteria for all immunizations and vaccines are listed in Part F of this section the Procedure Code Tables - Arkansas Department of Human Services.

Influenza virus vaccine through the Vaccines for Children (VFC) program is determined by the age of the beneficiary and obviously which vaccine is used.

The administration fee for all vaccines is included in the reimbursement fee for the vaccine CPT procedure code.

D. Vaccines for Children (VFC)

The Vaccines for Children (VFC) Program was established to generate awareness and access for childhood immunizations. Arkansas Medicaid established new procedure codes for billing the administration of VFC immunizations for children under the age of 19 years of age. To enroll in the VFC Program, contact the Arkansas Division of Health. Providers may also obtain the vaccines to administer from the Arkansas Division of Health. View or print Arkansas Division of Health contact information.

Medicaid policy regarding immunizations for adults remains unchanged by the VFC Program.

Vaccines available through the VFC Program are covered for Medicaid-eligible children. Administration fee only is reimbursed. When filing claims for administering VFC vaccines, providers must use the CPT procedure code for the vaccine administered. Electronic and paper claims require modifiers **EP** and **TJ**. ARKids First-B beneficiaries are not eligible for the Vaccines for Children (VFC) Program; however, vaccines can be obtained to administer to ARKids First-B beneficiaries who are under the age of 19 by contacting the Arkansas Department of Health and indicating the need to order ARKids-B SCHIP vaccines. View or print the Department of Health contact information.

When vaccines are administered to beneficiaries of ARKids First-B services, only modifier **SL** must be used for billing. Any additional billing and coverage protocols are listed under the specific procedure code in the tables section of this manual. See Part F of this section.

E. Billing of Multi-Use and Single-Use Vials

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

- Multiple units may be billed when applicable. Take-home drugs are not covered.
 Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.
- 2. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the

dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.

- a. **Single-Use Vials:** If the provider must discard the remainder of a **single-use vial** or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail</u> line with a JW (Drug wastage) modifier.
- b. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
- c. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
- d. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e. for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

See Section 292.910 for additional information regarding National Drug Code (NDC) billing.

F. Tables of Payable Procedure Codes

The tables of payable procedure codes are designed with eight columns of information.

- 1. The **first** column of the list contains the CPT or HCPCS procedure codes.
- 2. The <u>second</u> column indicates any modifiers that must be used in conjunction with the procedure code when billed, either electronically or on paper.
- 3. The <u>third</u> column indicates that the coverage of the procedure code is restricted based on the beneficiary's age in number of years(y) or months (m).
- 4. The **fourth** column indicates specific ICD primary diagnosis restrictions.
- 5. The <u>fifth</u> column contains information about the "diagnosis list" for which a procedure code may be used. See the page header for the diagnosis list 003 detail.
- 6. The <u>sixth</u> column indicates whether a procedure is subject to medical review before payment.
- 7. The <u>seventh</u> column indicates a procedure code requires a prior authorization before the service is provided. (See Section 261.100 for Prior Authorization instructions.)
- GF. Process for Obtaining a Prior Authorization Number from Arkansas Foundation for Medical Care (AFMC)the DHS contracted Prior Authorization vendor.

In collaboration with AFMC, DMS has changed the process for acquiring prior approval for drug procedure codes from a prior approval letter to a Prior Authorization number (PA). Instead of attaching a prior approval letter to a paper claim, providers will now list the Prior Authorization number on the claim. Drug procedure codes requiring Prior Authorization should be billed with the PA number listed on the claim form. These Covered drugs may be billed electronically or on a paper claim.

As part of the transition, AFMC will send a letter to all providers who have approval letters spanning timeframes within the last 365 days at the time of the effective date of this policy. The letter will contain a Prior Authorization number and the total remaining number of the approved units that can be billed. Any providers who have questions regarding Prior Authorization numbers and/or the transition process outlined above can contact AFMC at the following:

Toll Free: 1-877-350-2362, ext. 8741 or (501) 212-8741

A Prior Authorization number (PA) must be requested before treatment is initiated for any drug, therapeutic agent or treatment that indicates a Prior Authorization is required in a provider manual or an official Division of Medical Services correspondence.

The Prior Authorization requests should be completed using the approved AFMC contracted vendor Prior Authorization request form, and must be submitted by mail, fax or https://afmc.org.reviewpoint/ (View or print PA form.)

A decision letter will be returned to the provider by fax or e within five (5) business days.

If approved, the Prior Authorization number must be appended to all applicable claims, within the scope of the approval and may be billed electronically or on a paper claim with additional documentation when necessary.

Denials will be subject to reconsideration if received by AFMC the contracted vendor with additional documentation within fifteen (15) business days of date of denial letter.

A reconsideration decision will be returned within five (5) business days of receipt of the reconsideration request.

HG. Contact Information for Obtaining Prior Authorization

<u>View or print contact information to obtain the DHS or designated vendor step-by-step process for requesting prior authorization.</u>

When obtaining a Prior Authorization from the Arkansas Foundation for Medical Care, please send your request to the following:

When obtaining a Prior Authorization from the Arkansas Foundation for Medical Care, please send your request to the following:

In state and out of state toll free for inpatient reviews, Prior Authorizations for surgical procedures and assistant surgeons only	1 800 426 2234
General telephone contact, local or long distance Fort Smith	(479) 649-8501 1-877-650-2362
Fax for CHMS only	(479) 649 0776
Fax for Molecular Pathology only	(479) 649-9413
Fax General	(479) 649-0799
Fax Physician Drug Reviews Only (PDR)	(501) 212-8663
Web portal	https://afmc.org.reviewpoint/
Mailing address	Arkansas Foundation for Medical Care, Inc. P.O. Box 180001 Fort Smith, AR 72918 0001
Physical site location	5111 Rogers Avenue, Suite 476 Fort Smith, AR 72903
Office hours	8:00 a.m. until 4:30 p.m. (Central Time), Monday through Friday, except holidays

*Procedure code requires paper billing with applicable attachments and must follow NDC protocol. (See Section 292.910 for NDC protocol.)

See Sections 261.000 – 261.220 for prior authorization procedures.

See Section 244.100 for instructions regarding obtaining a Prior Approval Letter.

List <u>003/103603</u> diagnosis codes include: (<u>View ICD Codes</u>.) Diagnosis List <u>003/103603</u> restrictions apply to ages 21y and above unless otherwise indicated in the age restriction column.



TOC not required

242.400 Drug Procedure Codes and National Drug Codes (NDCs)

7-1-201-1 2:

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A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

LABELER CODE	LABELER NAME	EFFECTIVE DATE	TERMINATION DATE
00002	ELI LILLY AND COMPANY	1/1/1991	DAIL
00003	E.R. SQUIBB & SONS, INC	1/1/1991	
00004	HOFFMANN-LA ROCHE	1/1/1991	
00005	LEDERLE LABORATORIES	1/1/1991	
00006	MERCK & CO., INC.	1/1/1991	
00007	GLAXOSMITHKLINE	1/1/1991	
00008	WYETH LABORATORIES	1/1/1991	
00009	PFIZER, INC.	1/1/1991	
00011	BECTON DICKINSON MICROBIOLOGY SYSTEMS	10/1/1991	7/1/1998
00013	PFIZER, INC.	1/1/1991	

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date		
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000		
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000		
00004	GENENTECH, INC.	01/01/1991	01/01/3000		
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000		
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000		
80000	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000		
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000		
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000		
00014	PFIZER, INC	01/01/1991	01/01/3000		
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000		
00023	ALLERGAN INC	01/01/1991	01/01/3000		
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000		
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000		
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000		
00032	ABBVIE INC.	01/01/1991	01/01/3000		

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the NDC termination date. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA-assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

00123	0456	78
LABELER CODE	PRODUCT CODE	PACKAGE CODE
	(4 digits)	(2 digits)
(5 digits)		

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

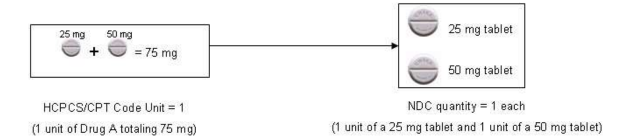
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

C. Claims Filing

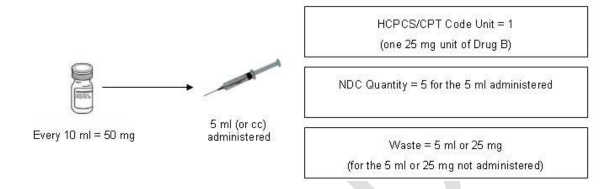
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug, whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets D (Electronic Claims Filing) and E (Paper Claims Filing) below.

Diagram 5



D. Electronic Claims Filing 837I (Outpatient)

Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.

Arkansas Medicaid will require providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

-Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same

NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- E. Paper Claims Filing CMS-1450 (UB-04)

Arkansas Medicaid will require providers billing drug HCPCS/CPT codes, including covered unlisted drug procedure codes, to use the required NDC format.

For institutional outpatient claims on the CMS-1450 (UB-04), use the locator field 43 (Description) to list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2

International Unit; GR - Gram; ML - Milliliter; UN - Unit), and number of units of the actual NDC administered, spaced and arranged exactly as in Diagram 6. Each NDC, when billed under the same procedure code on the same date of service, is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in the detail locator field 47 and total HCPCS/CPT code units in locator field 46. Each subsequent sequence number should show zeros in locator fields 46 and 47. See Detail 1, sequence 2 in Diagram 6. The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

----Diagram 6

	Sequence 1	42 FEX. CO.	43 DESCRIPTION		44 HCPGS / RATE / HPPS 000 E	45 SERV. DATE	46 SERV UNITS	47 TOTAL CHARGES	48 HON-COVERED CHARGES	4	
	sequence i	0636	N4 12345678912 U	N 1.00	Z1234	08/01/07	1	2500	1	П,	
	Sequence 2	0636	N4 01111222233 U	N 1.00	Z1234	08/01/07	0	0.00		,	
Detail 2		0305	Hemogram		85025	08/01/07	1	55 00		1	
		POSSOCIAL DATE:	N4 44444555506 U	N 5.00	Z6789	08/01/07	1	21.00			
	Da-32							1		4	
	Detail 3	1		44 HOPCS / R	VE /HIPPS DODE	46 SERV DATE	40 SERV. UNITS	47 TOTAL O-VARGE	5 42 NON	OOVERED O	DIARGES
REV. CD. 4		78912	UN 1.00	44 HOPOS / R	Towns or			47 TOTAL O-WARGE	2500	OOVERED O	DIARGES
636 N	43 DESCRIPTION	SPASSAGGISSON			KP	46 SERV CATE 01/01/22 01/01/22	1	# 47 TOTAL O-WAGE		COVERED O	DIARGES
636 N	40 DESCRIPTION NA 123456	22233	3 UN 1.00	Z1234	KP KQ	01/01/22	1	& TOTAL CHARGE	2500	COVERED O	DIARGES

F. Procedure Code/NDC Detail Attachment Form-DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. View or print form DMS-664 and instructions for completion.

Diagram 7

Detail #	Sequence #		NDC										Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

GF. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

H. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

IG. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, disputes or review issues, appeal hearings, investigations, or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing the purchase of drugs and documentation showing what drug (name, strength, and amount) was administered and on what date, to the beneficiary in question.

242.410 Billing of Multi-Use and Single-Use Vials

-1-1-151-1 23

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

- A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges 96365 through 96379.
- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description, the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - 1. **Single-Use Vials:** If the provider must discard the remainder of a **single-use vial** or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (Drug wastage) modifier.
 - 2. **Multi-Use Vials:** Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.



ARKids First-B Section II

TOC not required

262.431 Billing of Multi-Use and Single-Use Vials

2-1-221-1-23

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

View or print the procedure codes for ARKids First-B procedures and services.

- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description, the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.
 - 2. **Multi-Use Vials:** Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

TOC not required

272.531 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The Labeler termination date indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the termination date.

LABELER			EFFECTIV	E TERMINATION
CODE	LABELER NAME		DATE	DATE
00002	ELI LILLY AND COMPANY		1/1/1991	
00003	E.R. SQUIBB & SONS, INC		1/1/1991	
00004	HOFFMANN-LA ROCHE		1/1/1991	
00005	LEDERLE LABORATORIES		1/1/1991	
00006	MERCK & CO., INC.		1/1/1991	
00007	GLAXOSMITHKLINE		1/1/1991	
00008	WYETH LABORATORIES		1/1/1991	
00009	PFIZER, INC.		1/1/1991	
00011	BECTON DICKINSON MICROBIOLOG	Y SYSTEMS	10/1/1991	7/1/1998
00013	PFIZER, INC.		1/1/1991	
Labeler ID	Labeler Name	Contract	Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/0	1/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/0	11/1991	01/01/3000
00004	GENENTECH, INC.	01/0	1/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/0	1/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/0	1/1991	01/01/3000
00008	WYETH PHARMACEUTICALS LLC,	01/0	1/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/0	01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/0	1/1991	01/01/3000
00014	PFIZER, INC	01/0	01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/0	1/1991	01/01/3000
00023	ALLERGAN INC	01/0	1/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/0	1/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/0	1/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/0	1/1991	01/01/3000
00032	ABBVIE INC.	01/0	1/1991	01/01/3000

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

00123	0456	78
LABELER	PRODUCT	PACKAGE
CODE	CODE	CODE
(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345-6789-1	123456789 0 1
1111-2222-33	0 1111222233
01111-456-71	01111 0 45671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

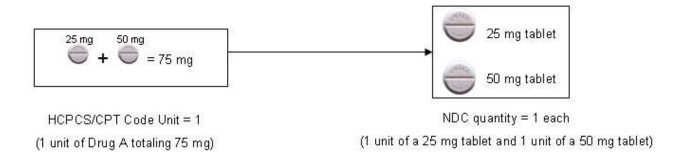
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

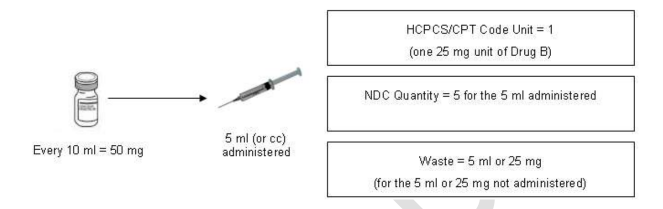
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



- A. Electronic Claims Filing 837P (Professional) and 837I (Outpatient)
 - Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid requires providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500

Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.

See Diagram 6 for CMS-1500.

- For professional claims, CMS-1500, list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2 International Unit; GR Gram; ML Milliliter; UN Unit) and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.
- Each NDC when billed under the same procedure code on the same date of service is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, Sequence 2 in Diagram 6.
- The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = <u>Drug wastage</u>

Sequence 1	MM	DO:	YY	мм	DD	YY	PLACE OF GERVICE		(Explain the CPT/HCPCS	Nutried Cit		DIFER		POINTER	\$ CHARGE	E9.	ONES UNTS	GHICT Family Plan	10. 004L	RENDERING PROVIDER ID. #
1	N4	1234	45678	912	UN	1.00				100	111	med 5	2811	100		av S		00	19900	123456789
Sequence 2	08	01	07	80	01	07	11		Z1234	1				1	25	00	- 1		NFI	
	N4	0111	11222	2233	UN	1.00					,					1111				123456789
	08	01	07	08	01	07	111		Z1234	1	-		1	14	- 0	00	0	1	HE	***********
Detail 2	,		-	Acceptance			-						-			-				123456789
	08	01	07	08	01	07	11		99213	1				1 1	- 55	00	1		NET	
		444	44555	506	ML:	5.00		-								edecomount.				123456789
Sequence 1 4	08	01	07	108	01	07	11		26789	1				1	35	00	-1		NPI	

24. A. MM	From	YY	MM	To	YY	PLACE OF SERVICE	C. EMG			VICES, OR SUPPLIES SUMMERCES)	E. DIAGNOSIS POINTER	\$ CHARGES	DAYS BY	ID.	RENDERING PROVIDER ID. #
V4 :	1234	1567	8912	UN	1.00				Jan H						123456789
)1	01	22	01	01	22	11		Z1234	KP		1	25 00	1	NPI	
14 (0111	122	2223	UN	1.00										123456789
)1	01	22	01	01	22	11		Z1234	KQ		1	25 00	1	NPI	
V4 4	4444	1445	5506	ML	3.0										123456789
	01		The state of the s	1000000	22	11		Z1234	KQ		1	75 00	3	NPL	
to contain	190		5506	200											123456789
)1	01	22	01	01	22	11		Z1234	JW		1	50 00	2	NPI	
							- 1							NPI	
			7		7										
					1	-		Contract of	article.		20000			NPI	No. of Concession,

Procedure Code/NDC Detail Attachment Form - DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the **DMS-664** "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Diagram 7

Detail #	Sequence #						NDC	8					Proc Code /Modifier	Drug Name/Dose/Route	Wasted
9	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

I<u>II</u>V. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations, or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing purchase of drugs and documentation showing what drug (name, strength, and amount) was administered and on what date, to the beneficiary in question.

See Section 272.533 for additional information regarding drug code billing.

272.533 Injections, Therapeutic and/or Diagnostic Agents

2-1-22<u>1-1-</u> 23

A. Providers billing the Arkansas Medicaid Program for covered injections should bill the appropriate CPT or HCPCS procedure code for the specific injection administered. The procedure codes and their descriptions may be found in the Current Procedure Terminology (CPT) and in the Healthcare Common Procedural Coding System Level II (HCPCS) coding books.

Injection administration code is payable for beneficiaries of all ages. May be used for billing the administration of subcutaneous and/or intramuscular injections only. This procedure code cannot be billed when the medication is administered "ORALLY." No fee is billable for drugs administered orally.

Cannot be billed separately for Influenza Virus vaccines or Vaccines for Children (VFC) vaccines.

Cannot be billed to administer any medication given for family planning purposes. No other fee is billable when the provider decides not to supply family planning injectable medications.

Cannot be billed when the drug administered is not FDA approved.

See the table below when billing:

Most of the cC overed drugs can be billed electronically or on paper. However, any covered drug marked with an asterisk (*) must be billed on paper with the name of the drug and desage listed in the "Procedures, Services, or Supplies" column, Field 24D, of the CMS-1500 claim form. View a CMS-1500 sample form. If requested, additional documentation may be required to justify medical necessity. Reimbursement for manually priced drugs is based on a percentage of the average wholesale price.

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs. See Section 272.531 for further information.

Administration of therapeutic agents is payable only if provided in a physician's office, place of service code "11." These procedures are not payable to the certified nurse-midwife if performed in any other setting. Therapeutic injections should only be provided by certified nurse-midwives experienced in the provision of these medications and who have the facilities to treat patients who may experience adverse reactions. The capability to treat infusion reactions with appropriate life support techniques should be immediately available. Only one administration fee is allowed per date of service unless "multiple sites" are indicated in the "Procedures, Services, or Supplies" field in the CMS-1500 claim form. Reimbursement for supplies is included in the administration fee. An administration fee is not allowed when drugs are given orally.

Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges for therapeutic and chemotherapy administration procedure codes.

B. For consideration of payable unlisted CPT/HCPCS drug procedure codes:

 The provider must submit an electronic or paper claim that includes a description of the drug being represented by the unlisted procedure code on the claim form.

- 2. Documentation that further describes the drug provided must be attached and must include justification for medical necessity.
- 3. All other billing requirements must be met in order for payment to be approved.

C. Immunizations

Physicians may bill for immunization procedures on the CMS-1500 claim form. <u>View a CMS-1500 sample form.</u>

Coverage criteria for all immunizations and vaccines are listed in Part F of this sectionthe Procedure Code Tables – Arkansas Department of Human Services.

Influenza virus vaccine through the Vaccines for Children (VFC) program is determined by the age of the beneficiary and which vaccine is used.

The administration fee for all vaccines is included in the reimbursement fee for the vaccine CPT procedure code.

D. Vaccines for Children (VFC)

The Vaccines for Children (VFC) Program was established to generate awareness and access for childhood immunizations. Arkansas Medicaid established new procedure codes for billing the administration of VFC immunizations for children under the age of 19 years of age. To enroll in the VFC Program, contact the Arkansas Department of Health. Providers may also obtain the vaccines to administer from the Arkansas Division of Health. View or print Arkansas Department of Health contact information.

Medicaid policy regarding immunizations for adults remains unchanged by the VFC Program.

Vaccines available through the VFC Program are covered for Medicaid-eligible children. Administration fee only is reimbursed. When filing claims for administering VFC vaccines, providers must use the CPT procedure code for the vaccine administered. Electronic and paper claims require modifiers **EP** and **TJ**. ARKids First-B beneficiaries are not eligible for the VFC Program; however vaccines can be obtained to administer to ARKids First-B beneficiaries who are under the age of 19 by contacting the Arkansas Department of Health and indicating the need to order ARKids First-B SCHIP vaccines. View or print the Department of Health contact information.

When vaccines are administered to beneficiaries of ARKids First-B services, only modifier **SL** must be used for billing. Any additional billing and coverage protocols are listed under the specific procedure code in the tables in this section of this manual. See Part F of this section.

E. Billing of Multi-Use and Single-Use Vials

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

- Multiple units may be billed when applicable. Take-home drugs are not covered.
 Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.
- When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.

a. **Single-Use Vials:** If the provider must discard the remainder of a **single-use vial** or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW (Drug wastage) modifier.</u>

- b. **Multi-Use Vials** are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
- c. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
- d. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

See Section 272.531 for additional information regarding National Drug Code (NDC) billing.

F. Tables of Payable Procedure Codes

The tables of payable procedure codes are designed with eight columns of information.

- The <u>first</u> column of the list contains the CPT or HCPCS procedure codes.
- 2. The <u>second</u> column indicates any modifiers that must be used in conjunction with the procedure code when billed, either electronically or on paper.
- 3. The <u>third</u> column indicates that the coverage of the procedure code is restricted based on the beneficiary's age in number of years(y) or months (m).
- 4. The **fourth** column indicates specific ICD primary diagnosis restrictions.
- 5. The <u>fifth</u> column contains information about the "diagnosis list" for which a procedure code may be used. See the page header for the diagnosis list 003 detail.
- 6. The <u>sixth</u> column indicates whether a procedure is subject to medical review before payment.
- 7. The <u>seventh</u> column indicates a procedure code requires a prior authorization before the service is provided. (See Section 240.000 for prior authorization.)
- GF. Process for Obtaining a Prior Authorization (PA) Number from Arkansas Foundation for Medical Care (AFMC)the DHS contracted Prior Authorization vendor.

In collaboration with AFMC, DMS is changing the process for acquiring prior approval for drug procedure codes from a prior approval letter to a PA number. Instead of attaching a prior approval letter to a paper claim, providers will now list the PA number on the claim. This will mean that effective for claims submitted on and after August 26, 2016, drug procedure codes requiring PA should be billed with the PA number listed on the claim form. TheseCovered drugs may be billed electronically or on a paper claim. Additionally, these procedure codes requiring a PA will no longer require manual review during the processing of the claim.

As part of the transition, AFMC will send a letter to all providers who have approval letters spanning timeframes within the last 365 days at the time of the effective date of this policy.

The letter will contain a PA number and the total remaining number of the approved units that can be billed. Any providers who have questions regarding PA numbers and/or the transition process outlined above can contact AFMC at the following:

Toll Free: 1-877-350-2362, ext. 8741 or (501) 212-8741

A PA must be requested before treatment is initiated for any drug, therapeutic agent or treatment that indicates a PA is required in a provider manual or an official Division of Medical Services correspondence.

The PA requests should be completed using the approved AFMC contracted vendor PA request form and must be submitted by mail, fax or https://afmc.org.reviewpoint/_(View or print PA form.)

A decision letter will be returned to the provider by fax or *e-mail* within five (5) business days.

If approved, the Prior Authorization number must be appended to all applicable claims, within the scope of the approval and may be billed electronically or on a paper claim with additional documentation when necessary.

Denials will be subject to reconsideration if received by AFMC-the contracted vendor with additional documentation within fifteen (15) business days of date of denial letter.

A reconsideration decision will be returned within five (5) business days of receipt of the reconsideration request.

HG. Contact Information for Obtaining Prior Authorization

<u>View or print contact information to obtain the DHS or designated vendor step-by-step process for requesting prior authorization.</u>

When obtaining a Prior Authorization from the Arkansas Foundation for Medical Care, please send your request to the following:

In-state and out-of-state toll free for inpatient reviews, Prior Authorizations for surgical procedures and assistant surgeons only	1-800-426-2234
General telephone contact, local or	(479) 649-8501
long distance Fort Smith	1-877-650-2362
Fax for CHMS only	(479) 649 0776
Fax for Molecular Pathology only	(479) 649-9413
Fax General	(479) 649-0799
Fax Physician Drug Reviews Only (PDR)	(501) 212-8663
Web portal	https://afmc.org.reviewpoint/
Mailing address	Arkansas Foundation for Medical Care, Inc.
	P.O. Box 180001
	Fort Smith, AR 72918-0001
Physical site location	5111 Rogers Avenue, Suite 476 Fort Smith, AR 72903

Office hours	8:00 a.m. until 4:30 p.m. (Central
	Time), Monday through Friday, except
	holidays

IH. All family planning procedures require an FP modifier and a primary family planning diagnosis on the claim.

*Procedure code requires paper billing with applicable attachments and must follow NDC protocol. (See Section 272.531 for NDC protocol.)

See Section 240.000-240.200 for prior authorization procedures.

List 003/103603 -diagnosis codes include: (View ICD Codes.) Diagnosis List 003/103603 restrictions apply to ages twenty-one (21) years and above unless otherwise indicated in the age restriction column.

TOC not required

242.141 Billing of Multi-Use and Single-Use Vials

2-1-221-1-23

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

<u>View or print the procedure codes and modifiers for Child Health Services/Early and</u> Periodic Screening, Diagnosis, and Treatment (EPSDT) services.

- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description, the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.
 - 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

TOC not required

272.102 Drug Procedure Codes and National Drug Codes (NDC)

-1-201-1

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005 for drugs. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on or after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. The effective date is when a manufacturer entered into a rebate agreement with CMS. The Labeler termination date indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the termination date. For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the NDC termination date. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA-assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 1 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

00123	0456	78
LABELER	PRODUCT CODE	PACKAGE CODE
CODE	(4 digits)	(2 digits)

(5 digits)	

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 2 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 2

	Required 11-digit NDC
10-digit FDA NDC on PACKAGE	(5-4-2) Billing Format
12345 6789 1	12345678901
1111-2222-33	01111222233
01111 456 71	01111045671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

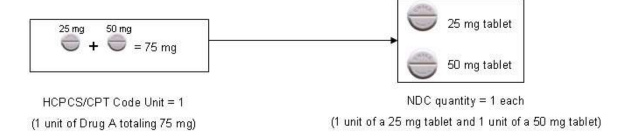
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines.

C. Claims Filing

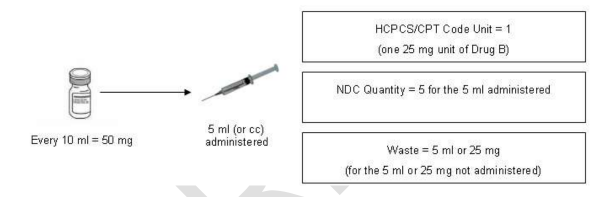
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug, whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example, whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 3.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. For billing wastage, see bullets D (Electronic Claims Filing) and E (Paper Claims Filing) below.

Diagram 4



D. Electronic Claims Filing 837I (Outpatient)

- Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid will require providers using electronic filing through the provider portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- o 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- o 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- o 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- o 4 or more NDCs for same procedure submit via paper claim
- o Wastage of each NDC shall be billed on a separate line with a JW modifier.
- NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

• KP = First drug of a multiple drug unit dose formulation

- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- E. Paper Claims Filing CMS-1450 (UB-04)

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- o 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- o 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- o 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- o Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage

Arkansas Medicaid will require providers billing drug HCPCS/CPT codes, including covered unlisted drug procedure codes, to use the required NDC format.

For institutional outpatient claims on the CMS 1450 (UB 04), use the locator field 43 (Description) to list the qualifier of "N4," the 11-digit NDC, the unit of measure qualifier (F2—International Unit; GR—Gram; ML—Milliliter; UN—Unit) and number of units of the actual NDC administered, spaced and arranged exactly as in Diagram 5. Each NDC when billed under the same procedure code on the same date of service is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail locator field 47 and total HCPCS/CPT code units in locator field 46. Each subsequent sequence number should show zeros in locator fields 46 and 47. See Detail 1, sequence 2 in Diagram 5. The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 5, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

		30	42 REV. CO.	43 DESCRIPTION	44 HCPCS / RATE / HPPS CODE	45 SERV. DATE	46 SERV UNITS	47 TOTAL CHARGES	49 HON-COVERED CHARGES	40		
	Seque	ence I	0636	N4 12345678912 UN 1.00	Z1234	08/01/07	1	2500	1 3	1		
	Seque	ence 2	0636	N4 01111222233 UN 1.00	Z1234	08/01/07	0	0.00		2		
ail 2	-		0305	Hemogram	85025	08/01/07	1	55 00		1		
	Seque	ence 1	0636	N4 44444555506 UN 5.00	Z6789	08/01/07	1	21:00				
r -	\$2.000	7 1							- 35	8		
	Detail 3]								ε		
i		an reserve	NOTION.		AA MIDDING / STATE (MARIDE CODE	de meno o	ATTE LAG	SERVINE LETTE	AL CHARGE	E	versom I	
	42 REV. CD.	43 DESC		-7004D UN 4 00	44 HCPCS / RATE / HPPS CODE	45 EERV. 0	-		N. OWRGES	48 NON-COMERCO O	HVACES	4
ļ	42 REV CD.	N4 1	23450	578912 UN 1.00	Z1234 KP	01/0	1/22	1	2500		HARCES .	4
ļ	42 REV CD.	N4 1	23450	578912 UN 1.00 222233 UN 1.00		01/0	-	1			HVAGES	4
	0636 0636	N4 1 N4 0	23450 1111		Z1234 KP	01/0 01/0	1/22	1 1	2500		OVECES	
2 2	0636 0636 0636	N4 1 N4 0 N4 4	23450 1111: 4444	222233 UN 1.00	Z1234 KP Z1234 KQ	01/0 01/0 01/0	1/22 01/22	1 1 3	2500 2500		-WACES	4

F. Procedure Code/NDC Detail Attachment Form- DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 6 for an example of the completed form. View or print form DMS-664 and instructions for completion.

— Diagram 6

Detail #	Sequence #	NDC											Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

GF. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

H. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

IG. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, disputes or review issues, appeal hearings, investigations, or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing the purchase of drugs and documentation showing what drug (name, strength, and amount) was administered and on what date, to the beneficiary in question.

See Section 272.510 for additional information regarding National Drug Code (NDC) billing.

272.510 Injections, Radiopharmaceuticals and Therapeutic Agents

2-1-221-1-23

Intravenous administration of therapeutic agents is payable only if provided in an outpatient setting. Therapeutic injections should only be provided by facilities that have the capacity to treat patients who may experience adverse reactions. The capability to treat infusion reactions with appropriate life support techniques should be immediately available. Reimbursement for supplies is included in the administration fee.

View or print the procedure codes for Hospital/Critical Access Hospitals/ESRD services.

Use procedure code for IV infusion therapy. For additional hours, sequential and/or concurrent infusions, bill revenue code **0760** (for observation), up to 8 hours maximum per day. For monoclonal antibody intravenous infusion use procedure code.

Multiple units may be billed for drug procedure codes, if appropriate. Take home drugs are not covered. Drugs loaded into an infusion pump are not classified as take home drugs.

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

- A. Multiple units may be billed when applicable. Take home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.
- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the desage included in the HCPCS procedure code description. If the desage given is not a multiple of the number provided in the HCPCS code description, the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - 1. **Single-Use Vials:** If the provider must discard the remainder of a **single-use vial** or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered.
 - 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - <u>31.</u> **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

See Section 272.102 for additional information regarding National Drug Code (NDC) billing.

See Section 272.450 for special billing instructions and coverage of Radiopharmaceuticals.

For coverage information regarding any drug not listed, please contact the Medicaid Reimbursement Unit. View or print Medicaid Reimbursement Unit contact information.

The following is a list of injections with special instructions for coverage and billing:

View or print the procedure codes for Hospital/Critical Access Hospitals/ESRD services.

Tables of Payable Procedure Codes

The tables of payable procedure codes are designed with eight columns of information.

- 1. The first column of the list contains the CPT or HCPCS procedure codes.
- 2. The <u>second</u> column indicates any modifiers that must be used in conjunction with the procedure code when billed, either electronically or on paper.
- 3. The <u>third</u> column indicates that the coverage of the procedure code is restricted based on the beneficiary's age in number of years(y) or months (m).
- 4. The **fourth** column indicates specific ICD-9-CM primary diagnosis restrictions.

- 5. The <u>fifth</u> column contains information about the "diagnosis list" for which a procedure code may be used. See the page header for the diagnosis list 003 detail.
- 6. The <u>sixth</u> column indicates whether a procedure is subject to medical review before payment.
- 7. The <u>seventh</u> column indicates a procedure code requires a prior authorization before the service is provided. (See Section 241.000 for prior authorization.)

*Procedure code requires paper billing with applicable attachments and must follow NDC protocol. (See Section 272.102 for NDC protocol.)

See Section 241.000 for prior authorization procedures.

See Section 272.103 for instructions regarding obtaining a Prior Approval Letter.

List 003/103 diagnosis codes include: (View ICD Codes) Diagnosis List 003/103 restrictions apply to ages 21y and above unless otherwise indicated in the age restriction column.

TOC not required

262.441 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

LABELER CODE	LABELER NAME		EFFECT DATE	Carrier of	TERMINATION DATE
00002	ELI LILLY AND COMPANY		1/1/199		DATE
00003	E.R. SQUIBB & SONS, INC		1/1/199	31	
00004	HOFFMANN-LA ROCHE		1/1/199	91	
00005	LEDERLE LABORATORIES		1/1/199	31	
00006	MERCK & CO., INC.		1/1/199	31	
00007	GLAXOSMITHKLINE		1/1/199	91	
00008	WYETH LABORATORIES		1/1/199	31	
00009	PFIZER, INC.		1/1/199	31	
00011	BECTON DICKINSON MICROBIOLOGY SYS	10/1/19	91	7/1/1998	
00013	PFIZER, INC.		1/1/199	31	
Labeler ID	Labeler Name	Contract B	egin Date	Cor	ntract End Date
00000	FILLIELY AND COMPANY	04/04/	(04/4004		

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000
00004	GENENTECH, INC.	01/01/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000
80000	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00014	PFIZER, INC	01/01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000
00023	ALLERGAN INC	01/01/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000
00032	ABBVIE INC.	01/01/1991	01/01/3000

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

00123	0456	78
LABELER	PRODUCT	PACKAGE
CODE	CODE	CODE
(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

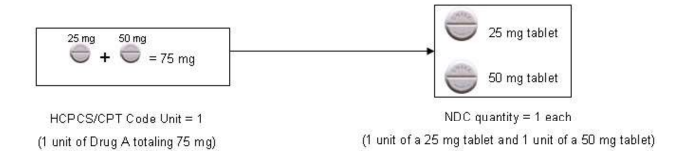
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

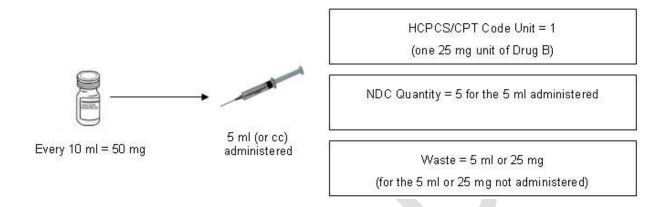
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



- A. Electronic Claims Filing 837P (Professional) and 837I (Outpatient)
 - Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid requires providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500

Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.

See Diagram 6 for CMS-1500.

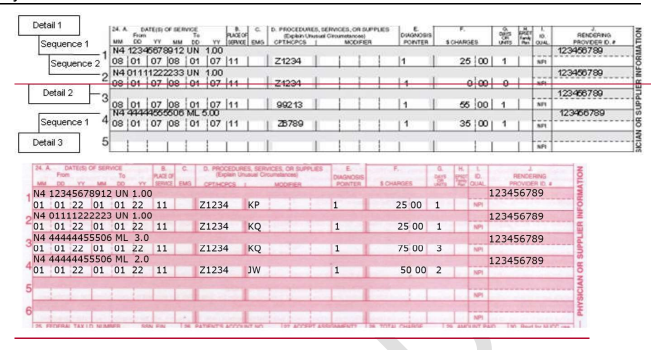
- For professional claims, CMS-1500, list the qualifier of "N4," the 11-digit NDC, the unit of measure qualifier (F2 International Unit; GR Gram; ML Milliliter; UN Unit) and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.
- Each NDC when billed under the same procedure code on the same date of service is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.
- The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences one (1) and two (2). Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence one (1) gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage



Procedure Code/NDC Detail Attachment Form - DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Diagram 7

Detail #	Sequence #						NDC	8					Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

IIIV. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC

invoices showing purchase of drugs and documentation showing what drug (name, strength and amount) was administered and on what date, to the beneficiary in question.

262.442 Billing of Multi-Use and Single-Use Vials

2-1-221-1-23

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

<u>View or print the procedure codes for Federally Qualified Health Center (FQHC)</u> services.

- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description, the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (Drug wastage) modifier.
 - 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

See Section 262.441 for additional information regarding National Drug Code (NDC) billing.

TOC not required

242.143 National Drug Codes (NDCs)

7-1-20

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. The effective date is when a manufacturer entered into a rebate agreement with CMS. The Labeler termination date indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the termination date. For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the NDC termination date. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA-assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 1 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

00123	0456	78

LABELER	PRODUCT CODE	PACKAGE CODE
CODE	(4 digits)	(2 digits)
(5 digits)	, ,	, ,

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 2 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 2

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	12345678901
1111-2222-33	01111222233
01111 456 71	01111045671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

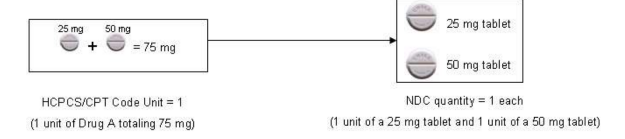
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

C. Claims Filing

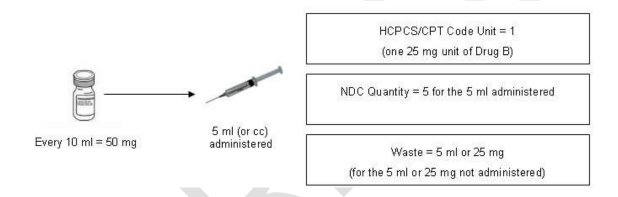
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug, whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example, whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 3.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. For billing wastage, see bullets D (Electronic Claims Filing) and E (Paper Claims Filing) below.

Diagram 4



- D. Electronic Claims Filing 837I (Outpatient)
 - Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid will require providers using electronic filing through the provider portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

• KP = First drug of a multiple drug unit dose formulation

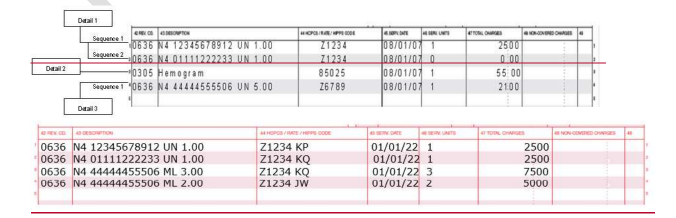
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- E. Paper Claims Filing CMS-1450 (UB-04)
- Arkansas Medicaid will require providers billing drug HCPCS/CPT codes, including covered unlisted drug procedure codes, to use the required NDC format.
- For institutional outpatient claims on the CMS-1450 (UB-04), use the locator field 43 (Description) to list the qualifier of "N4," the 11-digit NDC, the unit of measure qualifier (F2—International Unit; GR—Gram; ML—Milliliter; UN—Unit) and number of units of the actual NDC administered, spaced and arranged exactly as in Diagram 5. Each NDC when billed under the same procedure code on the same date of service is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail locator field 47 and total HCPCS/CPT code units in locator field 46. Each subsequent sequence number should show zeros in locator fields 46 and 47. See Detail 1, sequence 2 in Diagram 5. The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 5, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage



F. Procedure Code/NDC Detail Attachment Form-DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 6 for an example of the completed form. View or print form DMS-664 and instructions for completion.

Diagram 6

Detail #	Sequence #		NDC								Proc Code /Modifier	Drug Name/Dose/Route	Wasted		
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

GF. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

H. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

IG. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, disputes or review issues, appeal hearings, investigations, or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

—At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing the purchase of drugs and documentation showing what drug (name, strength, and amount) was administered and on what date, to the beneficiary in question.

242.144 Billing of Multi-Use and Single-Use Vials

2-1-22

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

View or print the procedure codes for Home Health services.

B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description, the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.

 Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.

- 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
- 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
- 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

See Section 242.143 for additional information regarding National Drug Code (NDC) billing.



TOC not required

242.401 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

LABELER		EFFECTIVE	TERMINATION
CODE	LABELER NAME	DATE	DATE
00002	ELI LILLY AND COMPANY	1/1/1991	
00003	E.R. SQUIBB & SONS, INC	1/1/1991	
00004	HOFFMANN-LA ROCHE	1/1/1991	
00005	LEDERLE LABORATORIES	1/1/1991	
00006	MERCK & CO., INC.	1/1/1991	
00007	GLAXOSMITHKLINE	1/1/1991	
00008	WYETH LABORATORIES	1/1/1991	
00009	PFIZER, INC.	1/1/1991	
00011	BECTON DICKINSON MICROBIOLOGY SYSTEMS	10/1/1991	7/1/1998
00013	PFIZER, INC.	1/1/1991	

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000
00004	GENENTECH, INC.	01/01/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000
80000	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00014	PFIZER, INC	01/01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000
00023	ALLERGAN INC	01/01/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000
00032	ABBVIE INC.	01/01/1991	01/01/3000

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

Diagram 2

00123	0456	78
LABELER	PRODUCT	PACKAGE
CODE	CODE	CODE
(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

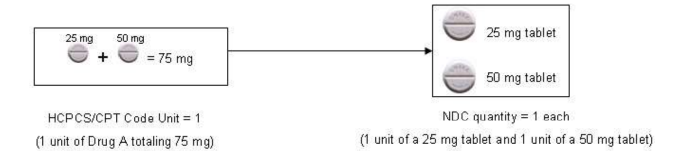
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

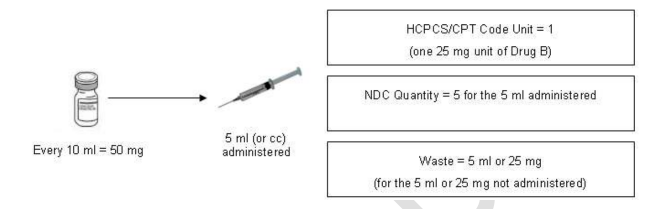
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



- A. Electronic Claims Filing 837P (Professional) and 837I (Outpatient)
- Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid requires providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500
- Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.
- See Diagram 6 for CMS-1500.

For professional claims, CMS-1500, list the qualifier of "N4," the 11-digit NDC, the unit of measure qualifier (F2 — International Unit; GR — Gram; ML — Milliliter; UN — Unit) and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.

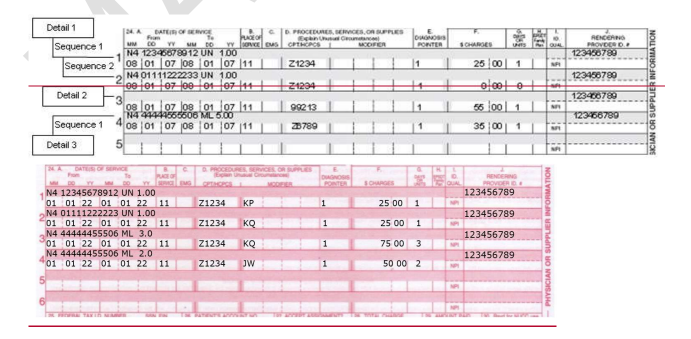
- Each NDC when billed under the same procedure code on the same date of service is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.
- The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage



Procedure Code/NDC Detail Attachment Form- DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Diagram 7

Detail #	Sequence #						NDC	â					Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

IIIV. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing purchase of drugs and documentation showing what drug (name, strength and amount) was administered and on what date, to the beneficiary in question.

242.402 Billing of Multi-Use and Single-Use Vials

2-1-221-1

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

View or print the procedure codes for Hyperalimentation services.

B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.

 Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.

- 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
- 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
- 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

TOC not required

252.438 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000
00004	GENENTECH, INC.	01/01/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000
00008	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00014	PFIZER, INC	01/01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000
00023	ALLERGAN INC	01/01/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000
00032	ABBVIE INC.	01/01/1991	01/01/3000

LABELER		EFFECTIVE	TERMINATION
CODE	LABELER NAME	DATE	DATE
00002	ELI LILLY AND COMPANY	1/1/1991	
00003	E.R. SQUIBB & SONS, INC	1/1/1991	
00004	HOFFMANN-LA ROCHE	1/1/1991	
00005	LEDERLE LABORATORIES	1/1/1991	
00006	MERCK & CO., INC.	1/1/1991	
00007	GLAXOSMITHKLINE	1/1/1991	
00008	WYETH LABORATORIES	1/1/1991	
00009	PFIZER, INC.	1/1/1991	
00011	BECTON DICKINSON MICROBIOLOGY SYSTEMS	10/1/1991	7/1/1998
00013	PFIZER, INC.	1/1/1991	

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

Diagram 2

	00123	0456	78
1	LABELER	PRODUCT	PACKAGE
	CODE	CODE	CODE
L	(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

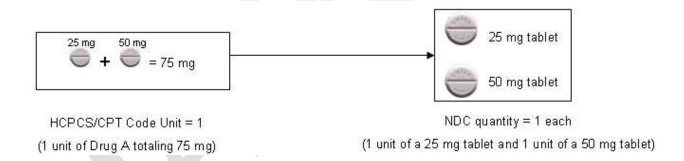
Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

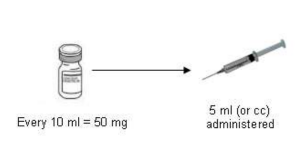
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.

Diagram 4



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.



HCPCS/CPT Code Unit = 1 (one 25 mg unit of Drug B)

NDC Quantity = 5 for the 5 ml administered

Waste = 5 ml or 25 mg (for the 5 ml or 25 mg not administered)

A. Electronic Claims Filing – 837P (Professional) and 837I (Outpatient)

- Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.
- Arkansas Medicaid requires providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500
- Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.
- See Diagram 6 for CMS-1500.
- For professional claims, CMS-1500, list the qualifier of "N4," the 11-digit NDC, the unit of measure qualifier (F2—International Unit; GR—Gram; ML—Milliliter; UN—Unit) and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.
- Each NDC when billed <u>under the same procedure code on the same date of service</u> is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and

total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.

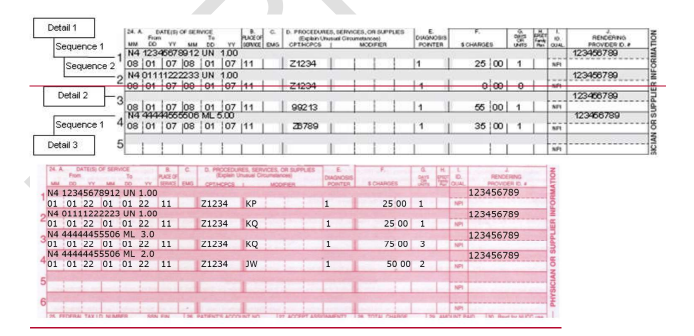
The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences one (1) and two (2). Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence one (1) gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- -----Diagram 6



Procedure Code/NDC Detail Attachment Form - DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Diagram 7

Detail #	Sequence #		v		_		NDC		n	Y			Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

IIIV. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing purchase of drugs and documentation showing what drug (name, strength and amount) was administered and on what date, to the beneficiary in question.

252.439 Billing of Multi-Use and Single-Use Vials

2-1-221-1

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

View or print the procedure codes for Nurse Practitioner services.

- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.
 - Multi-Use Vials: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.

3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.

4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.



TOC not required

242.450 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare & Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000
00004	GENENTECH, INC.	01/01/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000
80000	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00014	PFIZER, INC	01/01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000
00023	ALLERGAN INC	01/01/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000
00032	ABBVIE INC.	01/01/1991	01/01/3000

LABELER CODE	LABELER NAME	EFFECTIVE DATE	TERMINATION DATE
00002	ELI LILLY AND COMPANY	1/1/1991	
00003	E.R. SQUIBB & SONS, INC	1/1/1991	
00004	HOFFMANN-LA ROCHE	1/1/1991	
00005	LEDERLE LABORATORIES	1/1/1991	
00006	MERCK & CO., INC.	1/1/1991	
00007	GLAXOSMITHKLINE	1/1/1991	
00008	WYETH LABORATORIES	1/1/1991	
00009	PFIZER, INC.	1/1/1991	
00011	BECTON DICKINSON MICROBIOLOGY SYSTEMS	10/1/1991	7/1/1998
00013	PFIZER, INC.	1/1/1991	

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

00123	0456	78
LABELER	PRODUCT	PACKAGE
CODE	CODE	CODE
(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

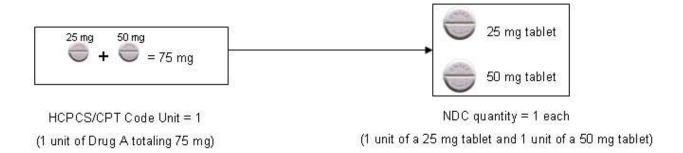
HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPC/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

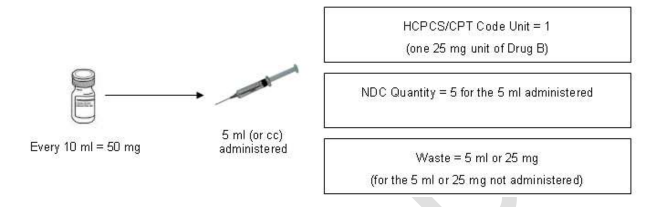
The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



A. Electronic Claims Filing – 837P (Professional) and 837I (Outpatient)

Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.

Arkansas Medicaid requires providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500 and CMS-1450 (UB-04)

Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.

See Diagram 6 for CMS-1500 and Diagram 7 for CMS-1450 (UB-04).

CMS-1500

For professional claims, CMS-1500, list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2 - International Unit; GR - Gram; ML - Milliliter; UN - Unit), and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.

- Each NDC, when billed under the same procedure code on the same date of service is defined as a "sequence". When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in the detail field 24F and total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.
- The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences one (1) and two (2). Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence one (1) gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage

24. A.	From	YY	MM	To	YY	PLACE OF SERVICE	EMG	(Explain U	husual Cir	cumetances)		DIAGNOSIS	\$ CHARGES	DAYS OR UNITS	PRO	ID. QUAL	RENDERING PROVIDER ID. #
V4 1	1234	567	8912	UN	1.00							75100			10		123456789
01	01	22	01	01	22	11		Z1234	KP			1	25 00	1		NP)	
N4 C	0111	122	2223	UN	1.00												123456789
01	01	22	01	01	22	11	100	Z1234	KQ			1	25 00	1		NPI	
14 4			5506	ML	3.0												123456789
O Personal Property of		22	01	01	22	11		Z1234	KQ			1	75 00	3		NP1	
			5506									- 18					123456789
)1	01	22	01	01	22	11		Z1234	JW			1	50 00	2		NPI	
					1000	-										NPI	
					1	-			-		100		market and the same		13	NPL	

Sequence 1	DA. A.	From DO	YY	MM	To DD	w	PLACE OF SERVICE	C. EMG	(Ephin) OPTHOPOS	NES, SER	roumeta	OR BUI NOM) ICIFIER	PPLIES	DIAGNOSIS PONTER	F. S CHARGES	OKYS CR UMB	DECT.	10	RENDERING PROVIDER D. #
Sequence 1	N4	1234	6678	912	UN	1.00	de my	1000	1000 million 1000	700	3711		10011					2026	123456789
Sequence 2	08	01	07	80	01	07	11		Z1234					1	25 00	1		NP1	
	N4 0	3111	1222	233	UN	1.00							72577			- 1			123456789
	00	01	07	00	01	07	111		Z1234	+	-	-	-	11	0 0	0 1	1	NPI	
Detail 2											.0								123456789
		01		08	01	07	111		99213					1 1	55 00	1		MPE	
Casting and State of	N4 2	444	4555	508	ML	5,00	No.			10		0.01	Sim	Total Trans			13 3		123456789
Sequence 1 4	08	01	07	80	01	07	111		26789	1		1		11	35 00	1	T	NPI	
Consessor I	-					1				2.5			1000			1	4	2000	
etail 3 📗 💲)					1	1 1			1	1	1		1 1	1	1	1	11F1	

CMS-1450 (UB-04)

For institutional outpatient claims on the CMS-1450 (UB-04), use the locator field 43 (Description) to list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2 - International Unit; GR - Gram; ML - Milliliter; UN - Unit), and number of units of the actual NDC administered, spaced, and arranged exactly as in Diagram 7.

Each NDC, when billed under the same procedure code on the same date of service is defined as a "sequence". When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in the detail locator field 47 and total HCPCS/CPT code units in locator field 46. Each subsequent sequence number should show zeros in locator fields 46 and 47. See Detail 1, sequence 2 in Diagram 7.

The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 7, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

Providers are instructed to bill as follows:

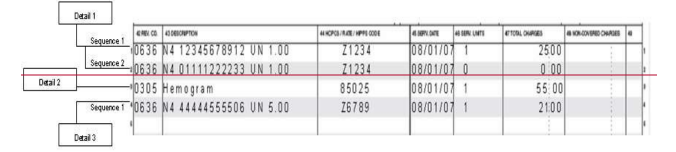
- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage

——Diagram 7

42 REV. CD.	43 DESCRIPTION	44 HOPCE / BATE / HIPPS DODE	40 SERV, DATE	40 SERV. UNITS	47 TOTAL CHARGES	48 NON-COVERED OHARGES	49
0636	N4 12345678912 UN 1.00	Z1234 KP	01/01/22	1	2500	1	
0636	N4 01111222233 UN 1.00	Z1234 KQ	01/01/22	1	2500		
0636	N4 44444455506 ML 3.00	Z1234 KQ	01/01/22	3	7500		
0636	N4 44444455506 ML 2.00	Z1234 JW	01/01/22	2	5000		



Procedure Code/NDC Detail Attachment Form - DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 8 for an example of the completed form. A copy of form DMS-664 is attached and may be copied for claim submission. -View or print form DMS-664 and instructions for completion.

Diagram 8

Detail #	Sequence #		NDC										Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML
	î			ľ									21		Ī

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

IIIV. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, disputes or review issues, appeal hearings, investigations, or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer. At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include NDC invoices showing the purchase of drugs and documentation showing what drug (name, strength, and amount) was administered and on what date, to the beneficiary in question.

TOC not required

242.401 National Drug Codes (NDCs)

7-1-201-1-23

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Health Care Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The *Labeler termination date* indicates that the manufacturer no longer participates in the federal rebate program and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the *termination date*.

Diagram 1

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000
00004	GENENTECH, INC.	01/01/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000
80000	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00014	PFIZER, INC	01/01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000
00023	ALLERGAN INC	01/01/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000
00032	ABBVIE INC.	01/01/1991	01/01/3000

LABELER		EFFECTIVE	TERMINATION			
CODE	LABELER NAME	DATE DATE				
00002	ELI LILLY AND COMPANY	1/1/1991				
00003	E.R. SQUIBB & SONS, INC	1/1/1991				
00004	HOFFMANN-LA ROCHE	1/1/1991				
00005	LEDERLE LABORATORIES	1/1/1991				
00006	MERCK & CO., INC.	1/1/1991				
00007	GLAXOSMITHKLINE	1/1/1991				
00008	WYETH LABORATORIES	1/1/1991				
00009	PFIZER, INC.	1/1/1991				
00011	BECTON DICKINSON MICROBIOLOGY SYSTEMS	10/1/1991	7/1/1998			
00013	PFIZER, INC.	1/1/1991				

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the *NDC termination date*. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code, and a 2-digit package code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

Diagram 2

	00123	0456	78
1	LABELER	PRODUCT	PACKAGE
	CODE	CODE	CODE
	(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid:

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	123456789 0 1
1111-2222-33	0 1111222233
01111 456 71	01111 0 45671

HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

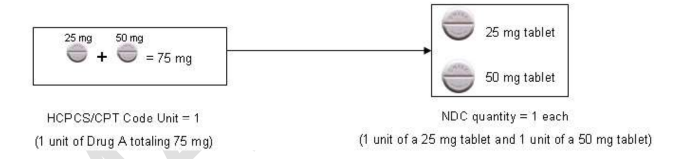
Exception: There is no requirement for an NDC when billing for vaccines, radiopharmaceuticals, and allergen immunotherapy.

II. Claims Filing

The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). —See Diagram 4.

Diagram 4



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. -See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



A. Electronic Claims Filing – 837P (Professional) and 837I (Outpatient)

Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.

Arkansas Medicaid requires providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage
- B. Paper Claims Filing CMS-1500
- Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.
- See Diagram 6 for CMS-1500.
- For professional claims, CMS-1500, list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2 International Unit; GR Gram; ML Milliliter; UN Unit), and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.
- Each NDC when billed <u>under the same procedure code on the same date of service</u> is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and

total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.

The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.

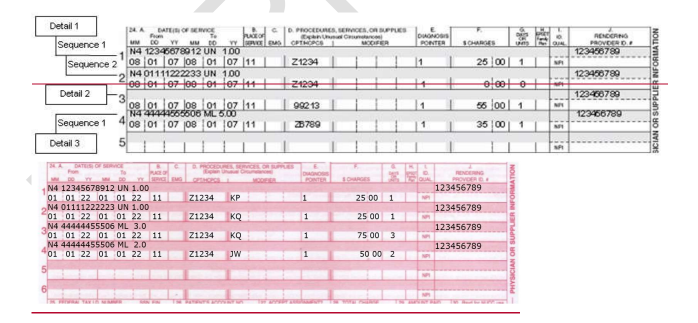
Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- JW = Drug wastage

Diagram 6



Procedure Code/NDC Detail Attachment Form—DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Diagram 7

Detail #	Sequence #		NDC										Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

IIIV. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing purchase of drugs and documentation showing what drug (name, strength and amount) was administered and on what date, to the beneficiary in question.

242.402 Billing of Multi-Use and Single-Use Vials

2-1-221-1

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

 Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

<u>View or print the procedure codes and modifiers for Durable Medical Equipment</u> (<u>DME</u>), oxygen equipment and supplies, orthotic appliances, prosthetic devices and medical supplies, procedures and services.

- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (Drug wastage) modifier.

2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.

- 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
- 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.



Rural Health Clinic Section II

TOC not required

252.103 Billing of Multi-Use and Single-Use Vials

2-1-221-1-23

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.

View or print the procedure codes for Rural Health Clinic (RHC) services.

- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.
 - 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

TOC not required

252.100 Ambulance Procedure Codes

2-1-221-1

The covered ambulance procedure codes are listed below.

View or print the procedure codes for Transportation (Ambulance) services.

Drug procedure codes require National Drug Codes (NDC) billing protocol. See Section 252.110 below.

*Procedure code can be billed <u>only</u> in conjunction with procedure code (please keep all documentation supporting the medical necessity of all codes billed for retrospective review of claims).

Arkansas Medicaid follows the billing protocol per the Federal Deficit Reduction Act of 2005 for drugs.

- A. Multiple units may be billed when applicable. Take-home drugs are not covered. Drugs loaded into an infusion pump are not classified as "take-home drugs." Refer to payable CPT code ranges.
- B. When submitting Arkansas Medicaid drug claims, drug units should be reported in multiples of the dosage included in the HCPCS procedure code description. If the dosage given is not a multiple of the number provided in the HCPCS code description the provider shall round up to the nearest whole number in order to express the HCPCS description number as a multiple.
 - Single-Use Vials: If the provider must discard the remainder of a single-use vial or other package after administering the prescribed dosage of any given drug, Arkansas Medicaid will cover the amount of the drug discarded along with the amount administered. <u>Discarded drugs shall be billed on a separate detail line with a JW</u> (<u>Drug wastage</u>) modifier.
 - 2. **Multi-Use Vials**: Multi-use vials are not subject to payment for any discarded amounts of the drug. The units billed must correspond with the units administered to the beneficiary.
 - 3. **Documentation:** The provider must clearly document in the patient's medical record the actual dose administered in addition to the exact amount wasted and the total amount the vial is labeled to contain.
 - 4. Paper Billing: For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing.

Remember to verify the milligrams given to the patient and then convert to the proper units for billing.

Follow the Centers for Disease Control (CDC) requirements for safe practices regarding expiration and sterility of multi-use vials.

252.110 National Drug Codes (NDC) Billing Protocol

7-1-201-1-

Effective for claims with dates of service on or after January 1, 2008, Arkansas Medicaid implemented billing protocol per the Federal Deficit Reduction Act of 2005. This explains policy and billing protocol for providers that submit claims for drug HCPCS/CPT codes with dates of service on and after January 1, 2008.

The Federal Deficit Reduction Act of 2005 mandates that Arkansas Medicaid require the submission of National Drug Codes (NDCs) on claims submitted with Healthcare Common Procedure Coding System, Level II/Current Procedural Terminology, 4th edition (HCPCS/CPT) codes for **drugs** administered. The purpose of this requirement is to assure that the State Medicaid Agencies obtain a rebate from those manufacturers who have signed a rebate agreement with the Centers for Medicare and Medicaid Services (CMS).

A. Covered Labelers

Arkansas Medicaid, by statute, will only pay for a drug procedure billed with an NDC when the pharmaceutical labeler of that drug is a covered labeler with Centers for Medicare and Medicaid Services (CMS). A "covered labeler" is a pharmaceutical manufacturer that has entered into a federal rebate agreement with CMS to provide each state a rebate for products reimbursed by Medicaid Programs. A covered labeler is identified by the first five (5) digits of the NDC. To assure a product is payable for administration to a Medicaid beneficiary, compare the labeler code (the first five (5) digits of the NDC) to the list of covered labelers which is maintained on the DHS contracted Pharmacy vendor website.

A complete listing of "Covered Labelers" is located on the website. See Diagram 1 for an example of this screen. The effective date is when a manufacturer entered into a rebate agreement with CMS. The Labeler termination date indicates that the manufacturer no longer participates in the federal rebate program, and therefore the products cannot be reimbursed by Arkansas Medicaid on or after the termination date.

Diagram 1

Labeler ID	Labeler Name	Contract Begin Date	Contract End Date
00002	ELI LILLY AND COMPANY	01/01/1991	01/01/3000
00003	E.R. SQUIBB & SONS, LLC.	01/01/1991	01/01/3000
00004	GENENTECH, INC.	01/01/1991	01/01/3000
00006	MERCK SHARP & DOHME CORP.	01/01/1991	01/01/3000
00007	GLAXOSMITHKLINE LLC	01/01/1991	01/01/3000
80000	WYETH PHARMACEUTICALS LLC,	01/01/1991	01/01/3000
00009	PHARMACIA AND UPJOHN COMPANY LLC	01/01/1991	01/01/3000
00013	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00014	PFIZER, INC	01/01/1991	01/01/3000
00015	MEAD JOHNSON AND COMPANY	01/01/1991	01/01/3000
00023	ALLERGAN INC	01/01/1991	01/01/3000
00024	SANOFI-AVENTIS, US LLC	01/01/1991	01/01/3000
00025	PFIZER LABORATORIES DIV PFIZER INC	01/01/1991	01/01/3000
00026	BAYER HEALTHCARE LLC	01/01/1991	01/01/3000
00032	ABBVIE INC.	01/01/1991	01/01/3000

For a claim with drug HCPCS/CPT codes to be eligible for payment, the detail date of service must be prior to the NDC termination date. The NDC termination date represents the shelf-life expiration date of the last batch produced, as supplied on the Centers for Medicare and Medicaid Services (CMS) quarterly update. The date is supplied to CMS by the drug manufacturer/distributor.

Arkansas Medicaid will deny claim details with drug HCPCS/CPT codes with a detail date of service equal to or greater than the NDC termination date.

When completing a Medicaid claim for administering a drug, indicate the HIPAA standard 11-digit NDC with no dashes or spaces. The 11-digit NDC is comprised of three (3) segments or codes: a 5-digit labeler code, a 4-digit product code and a 2-digit package

code. The 10-digit NDC assigned by the FDA printed on the drug package must be changed to the 11-digit format by inserting a leading zero (0) in one (1) of the three (3) segments. Below are examples of the FDA assigned NDC on a package changed to the appropriate 11-digit HIPAA standard format. Diagram 2 displays the labeler code as five (5) digits with leading zeros; the product code as four (4) digits with leading zeros; the package code as two (2) digits without leading zeros, using the "5-4-2" format.

Diagram 2

00123	0456	78
LABELER CODE	PRODUCT CODE	PACKAGE CODE
(5 digits)	(4 digits)	(2 digits)

NDCs submitted in any configuration other than the 11-digit format will be rejected/denied. NDCs billed to Medicaid for payment must use the 11-digit format without dashes or spaces between the numbers.

See Diagram 3 for sample NDCs as they might appear on drug packaging and the corresponding format which should be used for billing Arkansas Medicaid.

Diagram 3

10-digit FDA NDC on PACKAGE	Required 11-digit NDC (5-4-2) Billing Format
12345 6789 1	12345678901
1111-2222-33	01111222233
01111 456 71	01111045671

B. Drug Procedure Code (HCPCS/CPT) to NDC Relationship and Billing Principles

HCPCS/CPT codes and any modifiers will continue to be billed per the policy for each procedure code. However, the NDC and NDC quantity of the administered drug is now also required for correct billing of drug HCPCS/CPT codes. To maintain the integrity of the drug rebate program, it is important that the specific NDC from the package used at the time of the procedure be recorded for billing. HCPCS/CPT codes submitted using invalid NDCs or NDCs that were unavailable on the date of service will be rejected/denied. We encourage you to enlist the cooperation of all staff members involved in drug administration to assure collection or notation of the NDC from the actual package used. It is not recommended that billing of NDCs be based on a reference list, as NDCs vary from one (1) labeler to another, from one (1) package size to another, and from one (1) time period to another.

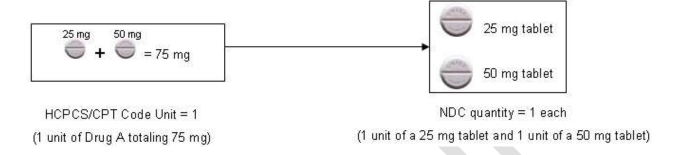
Exception: There is no requirement for an NDC when billing for vaccines.

II. Claims Filing

The HCPCS/CPT codes billing units and the NDC quantity do not always have a one-to-one relationship.

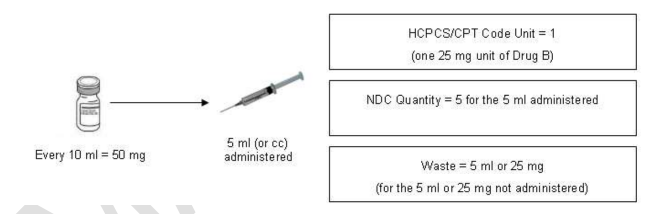
Example 1: The HCPCS/CPT code may specify up to 75 mg of the drug whereas the NDC quantity is typically billed in units, milliliters or grams. If the patient is provided 2 oral tablets, one at 25 mg and one at 50 mg, the HCPCS/CPT code unit would be 1 (1 total of 75 mg) in the example whereas the NDC quantity would be 1 each (1 unit of the 25 mg tablet and 1 unit of the 50 mg tablet). See Diagram 4.

Diagram 4



Example 2: If the drug in the example is an injection of 5 ml (or cc) of a product that was 50 mg per 10 ml of a 10 ml single-use vial, the HCPCS/CPT code unit would be 1 (1 unit of 25 mg) whereas the NDC quantity would be 5 (5 ml). In this example, 5 ml or 25 mg would be documented as wasted. See Diagram 5. For billing wastage, see bullets A (Electronic Claims Filing) and B (Paper Claims Filing) below.

Diagram 5



A. Electronic Claims Filing – 837P (Professional)

Procedure codes that do not require paper billing may be billed electronically. Any procedure codes that have required modifiers in the past will continue to require modifiers.

Arkansas Medicaid will require providers using electronic filing through the Provider Portal to use the required NDC format when billing HCPCS/CPT codes for administered drugs.

Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier

- 4 or more NDCs for same procedure submit via paper claim
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: The NDCs listed above are not the same (unless with a JW modifier). Same NDCs shall be billed on a single line with appropriate units.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- <u>JW = Drug wastage</u>
- B. Paper Claims Filing CMS-1500

Arkansas Medicaid will require providers billing drug HCPCS/CPT codes including covered unlisted drug procedure codes to use the required NDC format.

See Diagram 6 for CMS-1500.

CMS-1500

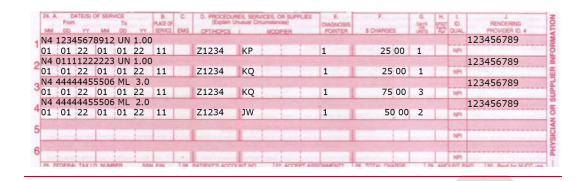
For professional claims, CMS-1500, list the qualifier of "N4", the 11-digit NDC, the unit of measure qualifier (F2 – International Unit; GR – Gram; ML - Milliliter; UN – Unit), and the number of units of the actual NDC administered in the shaded area above detail field 24A, spaced and arranged exactly as in Diagram 6.

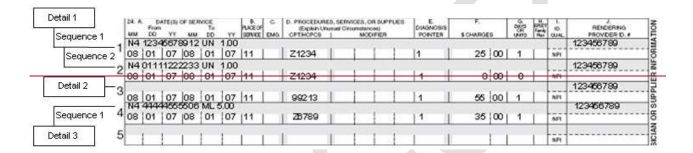
Providers are instructed to bill as follows:

- 1 NDC for a procedure 1st/only detail shall be billed with no modifier
- 2 NDCs for same procedure 1st detail shall be billed with a KP and 2nd gets billed with a KQ modifier
- 3 NDCs for same procedure 1st detail shall be billed with a KP and 2nd & 3rd detail get billed with a KQ modifier
- 4 or more NDCs for same procedure 1st detail shall be billed with a KP and 2nd and subsequent details shall be billed with a KQ modifier
- Wastage of each NDC shall be billed on a separate line with a JW modifier.

NOTE: CMS definitions of modifiers:

- KP = First drug of a multiple drug unit dose formulation
- KQ = Second or subsequent drug of a multiple drug unit dose formulation
- <u>JW = Drug wastage</u>
- Each NDC, when billed under the same procedure code on the same date of service is defined as a "sequence." When billing a single HCPCS/CPT code with multiple NDCs as detail sequences, the first sequence should reflect the total charges in detail field 24F and total HCPCS/CPT code units in detail field 24G. Each subsequent sequence number should show zeros in detail fields 24F and 24G. See Detail 1, sequence 2 in Diagram 6.
- The quantity of the NDC will be the total number of units billed for each specific NDC. See Diagram 6, first detail, sequences 1 and 2. Detail 2 is a Procedure Code that does not require an NDC. Detail 3, sequence 1 gives an example where only one (1) NDC is associated with the HCPCS/CPT code.





Procedure Code/NDC Detail Attachment Form- DMS-664

For drug HCPCS/CPT codes requiring paper billing (i.e., for manual review), complete every field of the DMS-664 "Procedure Code/NDC Detail Attachment Form." Attach this form and any other required documents to your claim when submitting it for processing. See Diagram 7 for an example of the completed form. Section V of the provider manual includes this form.

Complete instructions for accurate completion of form DMS-664 (including indication of required attachments) accompany the form. All forms are listed and accessible in Section V of each Provider Manual.

Diagram 7

Detail #	Sequence #		NDC										Proc Code /Modifier	Drug Name/Dose/Route	Wasted
1	1	1	2	3	4	5	6	7	8	9	1	2	Z1234	ABC drug/25 MG/Oral	0
1	2	0	1	1	1	1	2	2	2	2	3	3	Z1234	XYZ drug/50 MG/Oral	0
3	1	4	4	4	4	4	5	5	5	5	0	6	Z6789	PRQ drug/5 ML/IV	5 ML

III. Adjustments

Paper adjustments for paid claims filed with NDC numbers will not be accepted. Any original claim will have to be voided and a replacement claim will need to be filed. Providers have the option of adjusting a paper or electronic claim electronically.

IIIV. Remittance Advices

Only the first sequence in a detail will be displayed on the remittance advice reflecting either the total amount paid or the denial EOB(s) for the detail.

V. Record Retention

Each provider must retain all records for five (5) years from the date of service or until all audit questions, dispute or review issues, appeal hearings, investigations or administrative/judicial litigation to which the records may relate are concluded, whichever period is longer.

At times, a manufacturer may question the invoiced amount, which results in a drug rebate dispute. If this occurs, you may be contacted requesting a copy of your office records to include documentation pertaining to the billed HCPCS/CPT code. Requested records may include NDC invoices showing purchase of drugs and documentation showing what drug (name, strength and amount) was administered and on what date, to the beneficiary in question.

See Section 252.100 for additional information regarding drug code billing.

