

MEDICARE B Update!

A NEWSLETTER FOR FLORIDA MEDICARE PART B PROVIDERS

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The *Medicare B Update!* should be shared with all health care practitioners and managerial members of the provider/supplier staff. Publications issued beginning in 1997 are available at no cost from our provider education Web sites which may be accessed at: <http://www.feso.com>.

Routing Suggestions:

- Physician/Provider
- Office manager
- Billing/Vendor
- Nursing Staff
- Other _____



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Medicare B Update!

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PUBLICATIONS STAFF

TERRI DRURY
MILLIE C. PÉREZ
MARY BARNES
BETTY ALIX

The *Medicare B Update!* is published monthly by First Coast Service Options, Inc. (FCSO) Provider Outreach and Education Division, to provide timely and useful information to Medicare Part B providers in Connecticut and Florida.

Questions concerning this publication or its contents may be faxed to 1-904- 361-0723.

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THE FCSO MEDICARE B UPDATE!

ABOUT THE CONNECTICUT AND FLORIDA MEDICARE B UPDATE!

The *Medicare B Update!* is a comprehensive publication developed by First Coast Service Options, Inc. (FCSO) for Part B providers in Connecticut and Florida.

The Provider Outreach & Education Publications team distributes the *Medicare B Update!* on a monthly basis.

Important notifications that require communication in between publications will be posted to the FCSO Medicare provider education Web site, <http://www.fcsso.com>. In some cases, additional unscheduled special issues may be posted.

WHO RECEIVES THE UPDATE?

Anyone may view, print, or download the *Update!* from our provider education Web site(s). Providers who cannot obtain the *Update!* from the Internet are required to register with us to receive a complimentary hardcopy or CD-ROM.

Distribution of the *Update!* in hardcopy or CD-ROM format is limited to individual providers and professional association (PA) groups who have billed at least one Part B claim to either Connecticut or Florida Medicare for processing during the twelve months prior to the release of each issue. Providers meeting these criteria are eligible to receive a complimentary copy of that issue, *if a technical barrier exists that prevents them from obtaining it from the Internet and they have returned a completed registration form to us*. Registration forms must be submitted annually or when you experience a change in circumstances that impacts your electronic access.

For additional copies, providers may purchase a separate annual subscription in hardcopy or CD-ROM format (see order form in the back of this issue). All issues published since 1997 may be downloaded from the Internet, free of charge.

We use the same mailing address for *all* correspondence, and cannot designate that the *Update!* be sent to a specific person/department within a provider's office. To ensure continued receipt of all Medicare correspondence, providers must keep their addresses current with the Medicare Provider Enrollment department. Please remember that address changes must be done using the appropriate CMS-855.

CLEAR IDENTIFICATION OF STATE-SPECIFIC CONTENT

Articles common to both states appear at the beginning of the publication. Within common articles, references to phone numbers, addresses, reimbursement amounts, past publications, etc., are state-specific as appropriate. Content specific to Connecticut is next, followed by content specific to Florida. Connecticut and Florida local coverage determination (LCD) summaries are combined into one section. Articles in this section applies to both Connecticut and Florida unless otherwise noted.

PUBLICATION FORMAT

The *Update!* is arranged into distinct sections.

Following the table of contents, a letter from the carrier medical director (as needed), and an administrative information section, the *Update!* provides content applicable to both states, as noted previously. Within this section, information is categorized as follows.

- The **claims** section provides claim submission requirements and tips, plus correspondence (appeals and hearings) information.
- The **coverage/reimbursement** section discusses specific CPT and HCPCS procedure codes. It is arranged by specialty *categories* (not specialties). For example, "Mental Health" would present coverage information of interest to psychiatrists, clinical psychologists and clinical social workers, rather than listing articles separately under individual provider specialties. Also presented in this section are changes to the Medicare physician fee schedule, and other pricing issues.
- The section pertaining to **electronic data interchange** (EDI) submission also includes information pertaining to the Health Insurance Portability and Accountability Act (HIPAA).
- The **general information** section includes fraud and abuse, and Medicare Secondary Payer topics, plus additional topics not included elsewhere.

Educational resources. Important **addresses, phone numbers, and Web sites** will *always* be in state-specific sections.

QUARTERLY PROVIDER UPDATE

The Centers for Medicare & Medicaid Services (CMS) publishes the Quarterly Provider Update (QPU) at the beginning of each quarter to inform the public about:

- Regulations and major policies currently under development during this quarter.
- Regulations and major policies completed or canceled.
- New/revised manual instructions.

CMS regulations establish or modify the way CMS administers the Medicare program. These regulations impact providers and suppliers providing services to Medicare beneficiaries.

Providers may access the Quarterly Provider Update by going to the CMS Web site at <http://www.cms.hhs.gov/QuarterlyProviderUpdates/>.

Providers may join the CMS-QPU listserv to ensure timely notification of all additions to the QPU.

ADVANCE BENEFICIARY NOTICES

Medicare Part B allows coverage for services and items deemed medically reasonable and necessary for treatment and diagnosis of the patient. For some services, to ensure that payment is made only for medically necessary services or items, coverage may be limited based on one or more of the following factors (this list is not inclusive):

- Coverage for a service or item may be allowed only for specific diagnoses/conditions. Always code to the highest level of specificity.
- Coverage for a service or item may be allowed only when documentation supports the medical need for the service or item.
- Coverage for a service or item may be allowed only when its frequency is within the accepted standards of medical practice (i.e., a specified number of services in a specified timeframe for which the service may be covered).

If the provider believes that the service or item may not be covered as medically reasonable and necessary, the patient must be given an acceptable advance notice of Medicare's possible denial of payment if the provider does not want to accept financial responsibility for the service or item. Advance beneficiary notices (ABNs) advise beneficiaries, before items or services actually are furnished, when Medicare is likely to deny payment.

PATIENT LIABILITY NOTICE

The Centers for Medicare & Medicaid Services' (CMS) has developed the CMS-R131form as part of the Beneficiary Notices Initiative (BNI) The ABNs are designed to be beneficiary-friendly, readable and understandable, with patient options clearly defined.

There are two ABN forms - the General Use form (CMS-R-131G) and the Laboratory Tests form (CMS-R-131L). Both are standard forms that *may not be modified*; however, both contain customizable boxes for the individual requirements of users. Reproducible copies of Form CMS-R-131 ABNs (in English and Spanish) and other BNI information may be found on CMS's BNI Web site at

http://www.cms.hhs.gov/BNI/01_overview.asp#TopOfPage.

ABN MODIFIERS

When a patient is notified in advance that a service or item may be denied as not medically necessary, the provider must annotate this information on the claim (for both paper and electronic claims) by reporting modifier **GA** (waiver of liability statement on file) or **GZ** (item or service expected to be denied as not reasonable and necessary) with the service or item.

Failure to report modifier **GA** in cases where an appropriate advance notice was given to the patient may result in the provider having to assume financial responsibility for the denied service or item.

Modifier **GZ** may be used in cases where a signed ABN is *not* obtained from the patient; however, when modifier **GZ** is billed, the provider assumes financial responsibility if the service or item is denied.

“GA” MODIFIER AND APPEALS

When a patient is notified in advance that a service or item may be denied as not medically necessary, the provider must annotate this information on the claim (for both paper and electronic claims) by reporting the modifier **GA** (waiver of liability statement on file).

Failure to report modifier **GA** in cases where an appropriate advance notice was given to the patient may result in the provider having to assume financial responsibility for the denied service or item.

Nonassigned claims containing the modifier GA in which the patient has been found liable **must** have the patient's **written consent** for an appeal. Written appeals requests should be sent to:

Connecticut

Medicare Part B Redeterminations Appeals
PO Box 45010
Jacksonville, FL 32232-5010

OR

Florida

Medicare Part B Redeterminations Appeals
PO Box 2360
Jacksonville, FL 32231-0018

SIGN UP TO OUR *eNEWS* ELECTRONIC MAILING LIST

Join our **eNews** mailing list and receive urgent and other critical information issued by First Coast Service Options, Inc. (FCSO), your Medicare carrier. By signing up, you will receive automatic e-mail notification when new or updated information is posted to the provider education Web site. It's very easy to do. Simply go to our Web site <http://www.fcso.com>, select Medicare Providers, Connecticut or Florida, click on the "**eNews**" link located on the upper-right-hand corner of the page and follow the prompts.

ANNUAL *MEDICARE B UPDATE!* HARDCOPY/CD-ROM REGISTRATION FORM

To receive the *Medicare B Update!* in hardcopy, CD-ROM, or e-mail format, you must complete this registration form. To receive a hardcopy, CD-ROM or e-mail of future issues of the *Medicare B Update!* your form must be faxed to the number below on or before **June 30, 2008**. Providers currently receiving hardcopy publications that do not return this form by June 30, 2008, will not receive hardcopy versions after that date.

Please note that you are not obligated to complete this form to access information published in the *Medicare B Update!* Issues published beginning in 1997 are available **free** on our provider education Web site <http://www.fcso.com>, select *Medicare Providers, Florida Part A or B*, then select *Medicare Part B*, and then the *News & Bulletins* page on the left menu bar.

Provider/Facility Name:

National Provider Identifier (NPI):

Address:

City, State, ZIP Code:

Contact Person/Title:

Telephone Number:

Fax Number:

Email Address:

Rationale for needing a hardcopy:

Does your office have Internet access? YES NO

Do you have a PC with a CD-ROM drive? YES NO

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Fax your completed form to: **Medicare Publications at 904-361-0723**

Please share your questions and/or concerns regarding this initiative with us.

Additional questions or concerns may be submitted via the Medicare provider education Web site at <http://www.fcso.com>, select *Medicare Providers, Florida Part A or B or Connecticut Part B*, then *Contact* in the upper right corner of the page. You may also fax your questions or comments to 904-361-0723. **Our Provider Contact Center will not be able to respond to inquiries about this form.**

SIGN UP TO OUR *eNEWS* ELECTRONIC MAILING LIST

Join our **eNews** mailing list and receive urgent and other critical information issued by First Coast Service Options, Inc. (FCSO), your Medicare carrier. By signing up, you will receive automatic e-mail notification when new or updated information is posted to the provider education Web site. It's very easy to do. Simply go to our Web site <http://www.fcso.com>, select *Medicare Providers, Connecticut or Florida*, click on the "**eNews**" link located on the upper-right-hand corner of the page and follow the prompts.

CLAIMS

MEDICARE SHARED SYSTEMS MODIFICATIONS NECESSARY TO ACCEPT AND CROSSOVER TO MEDICAID NATIONAL DRUG CODES AND CORRESPONDING QUANTITIES SUBMITTED ON CMS-1500 PAPER CLAIMS

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

All physicians, providers, and suppliers who submit paper claims using the CMS-1500 to Medicare contractors (carriers, Medicare administrative contractors [A/B MACs], and durable medical equipment Medicare administrative contractors [DME/MACs]) for certain physician administered drugs provided to Medicare beneficiaries.

PROVIDER ACTION NEEDED

STOP – Impact to You

The Centers for Medicare & Medicaid Services (CMS) issued change request (CR) 5835 that notifies physicians and suppliers who use the CMS-1500 (those providers who qualify for a waiver from the Administrative Simplification Compliance Act [ASCA]) that changes are being made to Medicare systems to conform with instructions for submitting national drug codes (NDC) and quantity information on the CMS-1500.

CAUTION – What You Need to Know

This article only applies to those providers eligible to submit paper claims and who do so for patients who are dually eligible for Medicaid and Medicare. Such claims need to include NDCs and corresponding quantity amounts for physician-administered drugs. The *Key Points* section of this CR outlines the changes required in the CMS-1500.

GO – What You Need to Do

Make certain your office staffs are aware of these changes in the content requirements of your paper claims.

BACKGROUND

The Deficit Reduction Act (DRA) of 2005 required state Medicaid agencies to provide for the collection of NDC on all claims for certain physician-administered drugs for the purpose of billing manufacturers for Medicaid drug rebates. Prior to the DRA, physicians' offices, outpatient hospital departments and clinics generally used Healthcare Common Procedure Coding System (HCPCS) codes to bill Medicaid for drugs dispensed to Medicaid patients. However, because state Medicaid agencies are required to invoice manufacturers for rebates using NDCs for drugs for which the states have made payments, often states were not able to fulfill the rebate requirements for physician-administered drugs. The requirements for the collection of NDCs became effective beginning January 1, 2007. In addition, beginning January 1, 2008, in order for federal financial participation (FFP) to be available for these drugs, state Medicaid agencies must be in compliance with the requirements. These requirements were implemented in a final rule published on July 17, 2007.

Also, the quantity field of the CMS-1500 paper claim should be captured on all crossover claims for Medicaid billing, as provided for by the National Uniform Claims Committee (NUCC). Information regarding the quantities of physician-administered drugs billed to Medicaid is also necessary for states to bill manufacturers for Medicaid drug rebates.

Key Points

When required to submit NDC and quantity information for Medicaid rebates on the CMS-1500 paper claim be aware of the following:

- Submit the NDC in the red shaded portion of the detail line item in positions 01 through position 13.
- The NDC is to be preceded with the qualifier N4 and followed immediately by the 11-digit NDC (e.g. N49999999999).
- Report the NDC quantity in positions 17 through 24 of the same red shaded portion. The quantity is to be preceded by the appropriate qualifier: UN (units), F2 (international units), GR (gram) or ML (milliliter). There are six positions available for quantity. If the quantity is less than six positions, the entry should be left justified with spaces filling the remaining positions.

ADDITIONAL INFORMATION

To see the official instruction (CR 5835) issued to your Medicare carrier, DME/MAC, or A/B MAC refer to <http://www.cms.hhs.gov/Transmittals/downloads/R1401CP.pdf> on the CMS Web site.

If you have questions, please contact your Medicare carrier, DME/MAC, or A/B MAC at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5835

Related Change Request (CR) #: 5835

Related CR Release Date: December 21, 2007

Effective Date: April 7, 2008

Related CR Transmittal #: R1401CP

Implementation Date: April 7, 2008

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AMBULATORY SURGICAL CENTER

APRIL 2008 UPDATE TO THE AMBULATORY SURGICAL CENTER PAYMENT SYSTEM; SUMMARY OF PAYMENT POLICY CHANGES

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

This article is based on change request (CR) 5994 which describes changes to, and billing instructions for, payment policies implemented in the April 2008 ambulatory surgical center (ASC) update. This update provides updated payment rates for selected separately payable drugs and biologicals and provides rates and descriptors for newly created Level II Healthcare Common Procedure Coding System (HCPCS) codes for drugs and biologicals.

KEY POINTS

Billing for Drugs and Biologicals

- ASCs are strongly encouraged to report charges for all separately payable drugs and biologicals, using the correct HCPCS codes for the items used. ASCs billing for these products must make certain that the reported units of service of the reported HCPCS code are consistent with the quantity of the drug or biological that was used in the care of the patient. ASCs should not report HCPCS codes and separate charges for drugs and biologicals that receive packaged payment through the payment for the associated covered surgical procedure.
- If commercially available drug and biological products are being mixed together to facilitate their concurrent administration, the ASC should report the quantity of each product (reported by HCPCS code) that is separately payable in the ASC used in the care of the patient. Alternatively, if the ASC is compounding drugs that are not a mixture of commercially available products, but are a different product that has no applicable HCPCS code, the payment is packaged and no HCPCS coding is required. In these situations, ASCs should not report HCPCS code C9399. HCPCS code C9399, Unclassified drug or biological, is for new drugs and biologicals that are approved by the Food and Drug Administration (FDA) on or after January 1, 2004, for which a HCPCS code has not been assigned.

Drugs and Biologicals with Payment Based on Average Sales Price (ASP) Effective April 1, 2008

- Payments for separately payable drugs and biologicals based on the ASP will be updated on a quarterly basis as later quarter ASP submissions become available. In cases where adjustments to payment rates for previous quarters (January 2008) are necessary based on the most recent ASP submissions, the Centers for Medicare & Medicaid Services (CMS) will incorporate changes to the payment rates in the April 2008 release of the ASC drug file.
- Your Medicare contractors will make available to the ASCs the list of any newly added codes and previous quarter payment rate changes as identified in CR 5994.
- Providers take note that if your claims were processed prior to the installation of the revised January 2008 ASC Drug file, your Medicare AB/MAC or carrier will adjust, as appropriate, claims you bring to their attention that have dates of service on or after January 1, 2008 but prior to April 1, 2008.

New HCPCS Drug Codes Separately Payable under the ASC Payment System as of April 1, 2008

Four new HCPCS codes have been created effective April 1, 2008. These new HCPCS codes and their descriptors are listed in Table 1 below.

Table 1
New Drugs Separately Payable under the ASC Payment System as of April 1, 2008

HCPCS Code	Long Descriptor
C9241	Injection, doripenem, 10 mg
Q4096	Injection, von Willebrand Factor Complex, human, ristocetin cofactor (Not otherwise specified), per I.U. VWF:RCO,
Q4097	Injection, immune globulin (Privigen), intravenous, non-lyophilized (e.g., liquid), 500 mg
Q4098	Injection, iron dextran, 50 mg

The payment rates for the drugs in Table 1 can be found in the April 2008 update of the ASC Addendum BB which will be posted on the CMS Web site at the end of March.

HCPCS Drug Codes No Longer Payable under the ASC Payment System Effective April 1, 2008

The following drug codes have been deleted and are no longer payable by Medicare, effective April 1, 2008.

April 2008 Update to the ASC Payment System; Summary of Payment Policy Changes, continued

Table 2
Drugs HCPCS codes no longer eligible for payment under Medicare as of April 1, 2008

HCPCS Code	Long Descriptor	ASC Payment Status
J1751	Injection, iron dextran 165, 50 mg	Not payable by Medicare
J1752	Injection, iron dextran 267, 50 mg	Not payable by Medicare

Correct Reporting of Units for Drugs

ASCs are reminded to ensure that units of drugs administered to patients are accurately reported in terms of the dosage specified in the full HCPCS code descriptor. That is, units should be reported in multiples of the units included in the HCPCS descriptor.

- For example, if the drug's HCPCS code descriptor specifies 6 mg, and 6 mg of the drug were administered to the patient, the units billed should be 1.
- As another example, if the drug's HCPCS descriptor specifies 50 mg and 200 mg of the drug were administered to the patient, the units billed should be 4.
- ASCs should not bill the units based on how the drug is packaged, stored, or stocked. That is, if the HCPCS descriptor for the drug code specifies 1 mg and a 10 mg vial of the drug was administered to the patient, 10 units should be reported on the bill, even though only 1 vial was administered.
- HCPCS short descriptors are limited to 28 characters, including spaces; so short descriptors do not always capture the complete description of the drug. Therefore, before submitting Medicare claims for drugs and biologicals, it is extremely important to review the complete long descriptors for the applicable HCPCS codes.

ADDITIONAL INFORMATION

To see the official instruction (CR 5994) issued to your Medicare carrier or A/B MAC refer to <http://www.cms.hhs.gov/Transmittals/downloads/R1488CP.pdf> on the CMS Web site.

If you have questions, please contact your Medicare A/B MAC or carrier at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5994
Related Change Request (CR) #: 5994
Related CR Release Date: April 9, 2008
Effective Date: April 1, 2008
Related CR Transmittal #: R1488CP
Implementation Date: April 7, 2008

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DRUGS AND BIOLOGICALS

APRIL 2008 QUARTERLY AVERAGE SALES PRICE MEDICARE PART B DRUG PRICING FILES AND REVISIONS TO PRIOR QUARTERLY PRICING FILES

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

All physicians, providers and suppliers who submit claims to Medicare contractors (Medicare administrative contractors [A/B MACs], fiscal intermediaries [FIs], carriers, durable medical equipment Medicare administrative contractors [DME MACs] or regional home health intermediaries [RHHIs]) for services provided to Medicare beneficiaries.

WHAT YOU NEED TO KNOW

Change request (CR) 5982, from which this article is taken, instructs Medicare contractors to download and implement the April 2008 average sales price (ASP) drug pricing file for Medicare Part B drugs; and if released by CMS, also the revised January 2008, January 2007, April 2007, July 2007, October 2007, and October 2006 files.

BACKGROUND

Section 303(c) of the Medicare Modernization Act of 2003 revised the payment methodology for Part B covered drugs and biologicals that are not paid on a cost or prospective payment basis. Beginning January 1, 2005, the vast majority of drugs and biologicals not paid on a cost or prospective payment basis are paid based on the ASP

methodology, and pricing for compounded drugs has been performed by the local contractor.

Additionally, beginning in 2006, all end-stage renal disease (ESRD) drugs (that both independent and hospital-based ESRD facilities furnish), as well as specified covered outpatient drugs, and drugs and biologicals with pass-through status under the outpatient prospective payment system (OPPS), are paid based on the ASP methodology.

The ASP methodology is based on quarterly data that drug manufacturers submit to the Centers for Medicare & Medicaid Services (CMS), which CMS then provides (quarterly) to Medicare contractors (carriers, DME MACs, FIs, A/B MACs, and/or RHHIs) through the ASP drug pricing files for Medicare Part B drugs.

As announced in late 2006, CMS has been working further to ensure that accurate and separate payment is made for single source drugs and biologicals as required by section 1847A of the Social Security Act. As part of the effort to ensure compliance with this requirement, CMS has also reviewed how the terms "single source drug," "multiple source drug," and "biological product" have been operationalized in the context of payment under section 1847A.

April 2008 Quarterly ASP Medicare Part B Drug Pricing Files and Revisions to Prior Files, continued

For the purpose of identifying “single source drugs” and “biological products” subject to payment under section 1847A, CMS (and its contractors) will generally utilize a multi-step process that will consider:

- The FDA approval
- Therapeutic equivalents as determined by the FDA
- The date of first sale in the United States.

The payment limit for the following will be based on the pricing information for products marketed or sold under the applicable FDA approval:

- A biological product (as evidenced by a new FDA Biologic License Application or other relevant FDA approval), first sold in the United States after October 1, 2003; or
- A single source drug (a drug for which there are not two or more drug products that are rated as therapeutically equivalent in the most recent FDA Orange Book), first sold in the United States after October 1, 2003.

As appropriate, a unique HCPCS code will be assigned to facilitate separate payment. Separate payment may be operationalized through use of “not otherwise classified, (NOC)” HCPCS codes.

ASP Methodology

Beginning January 1, 2005, the payment allowance limits for Medicare Part B drugs and biologicals that are not paid on a cost or prospective payment basis are 106 percent of the ASP. Further, beginning January 1, 2006, payment allowance limits are paid based on 106 percent of the ASP for the following:

- ESRD drugs (when separately billed by freestanding and hospital-based ESRD facilities)
- Specified covered outpatient drugs and drugs and biologicals with pass-through status under the OPPS.

Beginning January 1, 2008, under the OPPS, payment allowance limits for specified covered outpatient drugs are paid based on 105 percent of the ASP. Drugs and biologicals with pass-through status under the OPPS continue to have a payment allowance limit of 106 percent of the ASP. CMS will update the payment allowance limits quarterly.

Exceptions are summarized as follows:

- The payment allowance limits for blood and blood products (other than blood clotting factors) that are not paid on a prospective payment basis are 95 percent of the average wholesale price (AWP) as reflected in the published compendia. The payment allowance limits are updated on a quarterly basis. Blood and blood products furnished in the hospital outpatient department are paid under OPPS at the amount specified for the ambulatory payment class (APC) to which the product is assigned.
- Payment allowance limits for infusion drugs furnished through a covered item of durable medical equipment on or after January 1, 2005, will continue to be 95 percent of the AWP reflected in the published compendia as of October 1, 2003, unless the drug is compounded or the drug is furnished incident to a professional service. The payment allowance limits will not be updated in 2008. The payment allowance limits for infusion drugs furnished through a covered item of durable medical equipment that were not listed

in the published compendia as of October 1, 2003, (i.e., new drugs) are 95 percent of the first published AWP unless the drug is compounded or the drug is furnished incident to a professional service.

- The payment allowance limits for influenza, pneumococcal and hepatitis B vaccines are 95 percent of the AWP as reflected in the published compendia except where the vaccine is furnished in a hospital outpatient department. Where the vaccine is administered in the hospital outpatient department, the vaccine is paid at reasonable cost.
 - The payment allowance limits for drugs and biologicals that are not included in the ASP Medicare Part B drug pricing file or NOC pricing file, other than new drugs and biologicals that are produced or distributed under a new drug application (or other application) approved by the FDA, are based on the published wholesale acquisition cost (WAC) or invoice pricing, except under OPPS where the payment allowance limit is 95 percent of the published AWP. In determining the payment limit based on WAC, the contractors follow the methodology specified in Pub. 100-04, chapter 17, Drugs and Biologicals, for calculating the AWP but substitute WAC for AWP. The payment limit is 100 percent of the lesser of the lowest-priced brand or median generic WAC. For 2006, the blood clotting furnishing factor of \$0.146 per I.U. is added to the payment amount for the blood-clotting factor when the blood-clotting factor is not included on the ASP file. For 2007, the blood clotting furnishing factor of \$0.152 per I.U. is added to the payment amount for the blood-clotting factor when the blood-clotting factor is not included on the ASP file. For 2008, the blood clotting furnishing factor of \$0.158 per I.U. is added to the payment amount for the blood-clotting factor when the blood-clotting factor is not included on the ASP file.
 - The payment allowance limits for new drugs and biologicals that are produced or distributed under a new drug application (or other new application) approved by the FDA and that are not included in the ASP Medicare Part B drug pricing file or NOC pricing file are based on 106 percent of the WAC, or invoice pricing if the WAC is not published, except under OPPS where the payment allowance limit is 95 percent of the published AWP. This policy applies only to new drugs and biologicals that were first sold on or after January 1, 2005.
 - The payment allowance limits for radiopharmaceuticals are not subject to the ASP payment methodology. Medicare contractors determine payment limits for radiopharmaceuticals based on the methodology in place as of November 2003 in the case of radiopharmaceuticals furnished in other than the hospital outpatient department. Radiopharmaceuticals furnished in the hospital outpatient department are paid charges reduced to cost by the hospital's overall cost to charge ratio.
- On or after March 18, 2008, the April 2008 ASP file will be available for download along with revisions to prior ASP payment files, if CMS determines that revisions to these prior files are necessary. On or after March 18, 2008, the April 2008 ASP NOC files will be available for retrieval from the CMS ASP Web page along with revisions to prior ASP NOC files, if CMS determines that revisions to these prior files are necessary. The payment limits included in revised

April 2008 Quarterly ASP Medicare Part B Drug Pricing Files and Revisions to Prior Files, continued

ASP and NOC payment files supersede the payment limits for these codes in any publication published prior to this document.

The payment files will be applied to claims processed or reprocessed on or after the implementation date of CR 5982 for the dates of service noted in the following table:

Payment Allowance Limit Revision Date	Applicable Dates of Service
April 2008 ASP and ASP NOC files	April 1, 2008, through June 30, 2008
January 2008 ASP and ASP NOC files	January 1, 2008, through March 31, 2008
October 2007 ASP and ASP NOC files	October 1, 2007, through December 31, 2007
July 2007 ASP and ASP NOC files	July 1, 2007, through September 30, 2007
April 2007 ASP and ASP NOC files	April 1, 2007, through June 30, 2007
January 2007 ASP and ASP NOC files	January 1, 2007, through March 31, 2007
October 2006 ASP and ASP NOC files	October 1, 2006, through December 31, 2006

Note: The absence or presence of a HCPCS code and its associated payment limit does not indicate Medicare coverage of the drug or biological. Similarly, the inclusion of a payment limit within a specific column does not indicate Medicare coverage of the drug in that specific category. The local Medicare contractor processing the claim makes these determinations.

Drugs Furnished During Filling or Refilling an Implantable Pump or Reservoir

Physicians may be paid for filling or refilling an implantable pump or reservoir when it is medically necessary for the physician (or other practitioner) to perform the service. Medicare contractors must find the use of the implantable pump or reservoir medically reasonable and necessary in order to allow payment for the professional service to fill or refill the implantable pump or reservoir and to allow payment for drugs furnished incident to the professional service.

If a physician (or other practitioner) is prescribing medication for a patient with an implantable pump, a nurse may refill the pump if the medication administered is accepted as a safe and effective treatment of the patient's illness or injury; there is a medical reason that the medication cannot be taken orally; and the skills of the nurse are needed to infuse the medication safely and effectively. Payment for drugs furnished incident to the filling or refilling of an implantable pump or reservoir is determined under the ASP methodology as described above. Note that your local Medicare contractor does pricing for compounded drugs.

ADDITIONAL INFORMATION

To see the official instruction (CR 5982) issued to your Medicare contractor visit <http://www.cms.hhs.gov/Transmittals/downloads/R1484CP.pdf> on the CMS Web site.

If you have questions, please contact your Medicare carrier at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5982

Related Change Request (CR) #: 5982

Related CR Release Date: March 26, 2008

Effective Date: April 1, 2008

Related CR Transmittal #: R1484CP

Implementation Date: April 7, 2008

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MODIFIER JW NOT REQUIRED

Change request (CR) 5923 allows contractors to decide on the use of the modifier JW when identifying unused drugs or biologicals from single-use vials or single-use packages that are appropriately discarded when processing all drugs except for those provided under the Competitive Acquisition Program (CAP).

First Coast Service Options Inc. (FCSO) has made a decision not to require the use of modifier JW. However, JW will remain an active modifier in the FCSO system in order to reimburse for wastage when billed on a single line item along with administered drug dosage.

For example, if a claim is received where the administered portion of the drug is billed on one line and the wastage amount is billed on a second line with modifier JW, then one line will be denied as a duplicate. The drug wastage portion should be included in the same line item billed for the actual administered portion. In addition, FCSO will deny claims for drugs provided under the CAP when billed with modifier JW.

Here is the link to the *MLN Matters* article MM5923 <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM5923.pdf>.

Here is the link to the official instruction issued to your Medicare carrier, DME/MAC, FI and/or A/B MAC for CR 5923 <http://www.cms.hhs.gov/Transmittals/downloads/R1478CP.pdf>.

Source: CMS Publication 100-04, Chapter 17, Section 40

NEW HCPCS CODES FOR THE APRIL 2008 UPDATE

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Physicians, providers, and suppliers submitting claims to Medicare contractors (carriers, DME Medicare administrative contractors [DME MACs], fiscal intermediaries [FIs], Part A/B Medicare administrative contractors [A/B MACs], and/or regional home health intermediaries [RHHIs]) for services provided to Medicare beneficiaries.

PROVIDER ACTION NEEDED

This article is based on change request (CR) 5981, which instructs Medicare contractors to implement Healthcare Common Procedure Coding System (HCPCS) code changes effective April 1, 2008. Make sure that your billing staffs are aware of these changes.

BACKGROUND

The Centers for Medicare & Medicaid Services (CMS) updates the HCPCS code set on a quarterly basis.

Effective for claims with dates of service on or after April 1, 2008, the following HCPCS codes will no longer be payable for Medicare:

HCPCS Code	Short Description	Long Description
J7602	Albuterol inh non-comp con	Albuterol, all formulations including separated isomers, inhalation solution, FDA-approved final product, noncompounded, administered through DME, concentrated form, per 1 mg (albuterol) or per 0.5 mg (levalbuterol)
J7603	Albuterol inh non-comp u d	Albuterol, all formulations including separated isomers, inhalation solution, FDA-approved final product, non-compounded, administered through DME, unit dose, per 1 mg (albuterol) or per 0.5 mg (levalbuterol)
J1751	Iron dextran 165 injection	Injection, iron dextran 165, 50 mg
J1752	Iron dextran 267 injection	Injection, iron dextran 267, 50 mg

Effective for claims with dates of service on or after April 1, 2008, the following HCPCS codes will be payable for Medicare:

HCPCS Code	Short Description	Long Description
J7611	Albuterol non-comp con	Albuterol, inhalation solution, FDA-approved final product, noncompounded, administered through DME, concentrated form, 1mg
J7612	Levalbuterol non-comp con	Levalbuterol, inhalation solution, FDA-approved final product, noncompounded, administered through DME, concentrated form, 0.5 mg
J7613	Albuterol non-comp unit	Albuterol, inhalation solution, FDA-approved final product, non-compounded, administered through DME, unit dose, 1mg
J7614	Levalbuterol non-comp unit	Levalbuterol, inhalation solution, FDA-approved final product, noncompounded, administered through DME, unit dose, 0.5 mg
Q4096	VWF complex, NOS	Injection, von Willebrand factor complex, human, ristocetin cofactor (not otherwise specified), per I.U. vWF:RCO
Q4097	Inj IVIG Privigen 500 mg	Injection, immune globulin (Privigen™), intravenous, nonlyophilized (e.g., liquid), 500 mg
Q4098	Inj iron dextran	Injection, iron dextran, 50mg
Q4099	Formoterol fumarate, inh	Formoretol fumarate, inhalation solution, FDA-approved final product, noncompounded, administered through DME, unit dose form, 20 micrograms

Currently, Alphanate® is the only product that should be billed using HCPCS code Q4096. HCPCS code J7190 should continue to be billed when Alphanate® is furnished for purposes of administering factor VIII. The blood clotting furnishing fee is payable when payment is allowed for HCPCS code Q4096. When a payment allowance limit for HCPCS code Q4096 is included on the quarterly Part B drug pricing files, the payment allowance limit will include payment for the blood clotting furnishing fee.

Effective for dates of service on or after April 1, 2008, the requirements under CR 5713 (See the *MLN Matters* article for CR 5713, which is on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM5713.pdf>) are being updated by CR 5981 to apply to claims that bill Intravenous Immune globulins (IVIG) using Q4097 as follows:

- Effective for dates of service on or after April 1, 2008, Medicare contractors will:
 - Only pay a claim for preadministration-related services (G0332) associated with IVIG administration if HCPCS code G0332, the drug (IVIG, HCPCS codes: J1566, J1568, J1569, J1561, J1572 and/or Q4097), and the drug administration service are all billed on the same claim for the same date of service.
 - Return institutional claims for G0332 to the provider if J1566, J1568, J1569, J1561, J1572 and/or Q4097 and a drug administration service are not also billed for the same date of service on the same claim.

New HCPCS Codes for the April 2008 Update, continued

- Reject professional claims as unprocessable for G0332 if J1566, J1568, J1569, J1561, J1572 and/or Q4097 and a drug administration service are not billed for the same date of service on the same claim.
- Use the appropriate reason/remark messages such as: M67 "Missing other procedure codes" and/or 16 "Claim/service lacks information" which are needed for adjudication when claims are returned/rejected.

ADDITIONAL INFORMATION

The official instruction, CR 5981, issued to your carrier, FI, RHHI, A/B MAC, and DME MAC regarding these changes may be viewed on the CMS Web site at <http://www.cms.hhs.gov/transmittals/downloads/R1492CP.pdf>.

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END-STAGE RENAL DISEASE

MANUALIZATION OF PAYMENT FOR OUTPATIENT ESRD RELATED SERVICES

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Physicians and other practitioners who bill Medicare contractors (carriers or Medicare administrative contractors [A/B MAC]) for providing outpatient end-stage renal disease (ESRD) services to Medicare beneficiaries.

WHAT YOU NEED TO KNOW

CR 5931, from which this article is taken, updates the *Medicare Claims Processing Manual*, chapter 8 (Outpatient ESRD Hospital, Independent Facility, and Physician/Supplier Claims), section 140 (Monthly Capitation Payment Method for Physicians' Services Furnished to Patients on Maintenance Dialysis) to reflect changes in the payment methodology for ESRD services (as discussed in the CY 2004 and CY 2005 Medicare physician fee schedule final rules).

ESRD related services (per full month), as described by *Current Procedural Terminology (CPT)* codes 90918-90921, and those (less-than-full-month), as described by *CPT* codes 90922-90925, are no longer valid for Medicare. They have been replaced by Healthcare Common Procedure Coding System (HCPCS) codes G0308 through G0327 (HCPCS codes G0308-G0319 are used for center based patients on dialysis, HCPCS codes G0320 – G0323 are used for home dialysis patients and HCPCS codes G0324 through G0327 are used for less-than-full-month services).

These policies have been discussed in prior communications to providers and CMS is now incorporating these changes into their manuals. Thus, while this article is informational in nature, be sure that your billing staffs are aware of these coding policies for ESRD services.

BACKGROUND

In the *Federal Register* published November 7, 2003, (68 FR 63216), CMS established new G codes for managing dialysis patients; with varying monthly capitation payments (MCP) based on 1) the number of visits provided within each month, and 2) the beneficiary's age.

Under this payment methodology, physicians bill separate codes for providing one end stage renal disease (ESRD) related visit per month, two to three visits per month, or four or more visits per month; and in turn, receive the lowest payment amount when providing one visit per

If you have questions, please contact your Medicare carrier at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5981

Related Change Request (CR) Number: 5981

Related CR Release Date: April 18, 2008

Related CR Transmittal Number: R1492CP

Effective Date: April 1, 2008

Implementation Date: April 7, 2008

month, a higher payment when providing two to three visits per month, and the highest payment amount when providing at least four visits per month.

On September 17, 2004, CR 3414, "Payment for Outpatient ESRD-Related Services," provided interim billing instructions for specific less-than-full-month ESRD-related scenarios (e.g. transient patients) and for visits furnished to patients in hospital observation status. In these two instances, physicians and practitioners were instructed to use the unlisted dialysis procedure code (*CPT* code 90999). To view the related *MLN Matters* article on "Payment for Outpatient ESRD-Related Services", please visit <http://www.cms.hhs.gov/MLNMattersArticles/Downloads/MM3414.pdf> on the CMS Web site.

Subsequently, in the *Federal Register* published November 15, 2004, (69 FR 66357), CMS: 1) changed the descriptor of the G codes for ESRD-related home dialysis services, less-than-full-month (G0324 through G0327) to allow other partial month scenarios (in addition to patients dialyzing at home); and 2) established policy that permits visits furnished to beneficiaries in hospital observation status to be counted for purposes of billing the MCP service. These policy changes superseded the interim billing instructions contained in CR 3414.

Finally, CR 3595, "Emergency Update to the CY 2005 Physician Fee Schedule Data Base", which is discussed in *MLN Matters* article MM3595 at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM3595.pdf> on the CMS Web site, published December 23, 2004, included descriptors of G0324-G0327 to allow these codes to be used for other scenarios in addition to home dialysis less than full month, (e.g. transient patients, partial month due to hospitalization, transplant or when the patient expired, and when a permanent change in MCP physician occurs during the month).

Now, CR 5931, from which this article is taken, updates the *Medicare Claims Processing Manual*, chapter 8 (Outpatient ESRD Hospital, Independent Facility, and Physician/Supplier Claims), section 140 (Monthly Capitation Payment Method for Physicians' Services Furnished to Patients on Maintenance Dialysis) to reflect these requirement changes.

Manualization of Payment for Outpatient ESRD Related Services, continued

Additionally, it notifies you that this section has been reorganized. Some of the information previously contained in sections 140.1 and 140.2 has been moved to section 140; and the title of section 140.1 has been changed to "Payment for ESRD-Related Services Under the Monthly Capitation Payment (Center Based Patients)" and the title of section 140.2 has been changed to "Payment for ESRD-Related Services (Per Diem)". Further, Sections 140.5, (Determining MCP Amount for Physician's Service to Maintenance Dialysis Patients) and 140.51 (Temporary Absence Under MCP) were deleted from chapter 8 as these sections are superseded by the new instructions.

Note: ESRD-related services as described by Healthcare Common Procedure Coding System (HCPCS) codes G0308 - G0327 are already included as part of the HCPCS payment file, and Medicare contractors are currently making payment for these service.

ADDITIONAL INFORMATION

You may see these policies for the payment for outpatient ESRD-related services by going to CR 5931, located at <http://www.cms.hhs.gov/Transmittals/downloads/R1456CP.pdf> on the CMS Web site. You will find the updated *Medicare Claims Processing Manual*, Chapter 8 (Outpatient ESRD Hospital, Independent Facility, and Physician/Supplier Claims), Sections 140 (Monthly Capitation Payment Method for Physicians' Services Furnished to Patients on Maintenance Dialysis), 140.1 (Payment for ESRD-Related Services Under the Monthly Capitation Payment [Center Based Patients]), 140.1.1

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(Payment for Managing Patients on Home Dialysis), 140.1.2 (Patients That Switch Modalities [Center to Home and Vice Versa]), 140.2 (Payment for ESRD-Related Services [Per Diem]), 140.2.1 (Guidelines for Physician or Practitioner Billing [Per Diem]), 140.3 (Data Elements Required on Claim for Monthly Capitation Payment), and 140.4 (Controlling Claims Paid Under the Monthly Capitation Payment Method) as an attachment to that CR.

You might also want to look at *MLN Matters* articles Payment for Outpatient ESRD-Related Services (September 17, 2004) and Emergency Update to the 2005 Medicare Physician Fee Schedule Database (MPFSDB) (December 23, 2004) which you may find at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM3414.pdf> and <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM3595.pdf> on the CMS Web site.

If you have any questions, please contact your carrier or /B MAC at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5931
Related Change Request (CR) #: 5931
Related CR Release Date: February 22, 2008
Effective Date: March 24, 2008
Related CR Transmittal #: R1456CP
Implementation Date: March 24, 2008

SERVICE INCLUDED IN THE MONTHLY CAPITATION PAYMENT

As a reminder, interpretations of the following tests are included in monthly capitation payment (MCP):

- Bone mineral density studies (CPT codes 76070, 76075, 78350, and 78351)
- Noninvasive vascular diagnostic studies of hemodialysis access (CPT codes 93925, 93926, 93930, 93931, and 93990)
- Nerve conduction studies (CPT codes 95900, 95903, 95904, 95925, 95926, 95927, 95934, 95935, and 95936)
- Electromyography studies (CPT codes 95860, 95861, 95863, 95864, 95867, 95867, 95869, and 95872).

Note: Separate payment may be made for medically necessary services that are included or bundled into the MCP (e.g., test interpretations) when furnished by physicians other than the monthly capitation payment physician.

Source: Publication 100-04, Chapter 8, Section 140A

EVALUATION AND MANAGEMENT SERVICES**NURSING FACILITY SERVICES (CODES 99304 - 99318)**

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Physicians and qualified nonphysician practitioners who bill Medicare contractors (carriers or Medicare administrative contractors [A/B MAC]) for services provided to Medicare beneficiaries in skilled nursing facilities (SNF) or nursing facilities (NF).

WHAT YOU NEED TO KNOW

Change request (CR) 5986, from which this article is taken, updates the *Medicare Claims Processing Manual*, chapter 12, (Physicians/Non-physician Practitioners), section 30.6.13 (Nursing Facility Services [codes 99304 -

99318]) subsection F (Use of the Prolonged Services Codes and Other Time-Related Services) by noting that the typical/average time units for evaluation and management (E/M) visit codes in the NF services code family are reestablished and applicable, as of January 1, 2008.

Effective for services on or after July 1, 2008, you may bill Medicare for medically necessary prolonged services for E/M visits (codes 99356 and 99357) in a SNF or NF with NF services codes (99304 - 99306, 99307 - 99310 and 99318). Additionally, you may use these prolonged services codes (99356 and 99357) with NF services in the code range (99304 - 99306, 99307 - 99310, and 99318) to bill

Nursing Facility Services (Codes 99304 - 99318), continued

for counseling and/or coordination of care services that are based on time.

Make sure that your billing staffs are aware of these new billing changes.

BACKGROUND

Effective January 1, 2006, the American Medical Association (AMA) *Current Procedural Terminology (CPT)* panel removed the typical/average time units for evaluation and management (E/M) services in the NF code family. Until these typical/average times were reestablished, this action precluded the billing of: 1) prolonged services for E/M visits in an SNF or NF; and 2) time-based counseling and/or coordination of care for NF Services.

CR 5986, from which this article is taken, updates the *Medicare Claims Processing Manual*, chapter 12, (Physicians/Non-physician Practitioners), section 30.6.13 (Nursing Facility Services [Codes 99304 – 99318]), subsection F (Use of the Prolonged Services Codes and Other Time-Related Services) by announcing that the AMA CPT panel has reestablished these typical/average time units beginning January 1, 2008.

Further, CR 5968 announces that, effective July 1, 2008, you may bill for medically necessary prolonged services for SNF or NF E/M visits (CPT codes 99356 and 99357) with NF services (code range 99304 – 99306, 99307 – 99310 and 99318); and you may also use the medically necessary prolonged services CPT codes (99356 and 99357) to bill for medically necessary E/M visits for time-based counseling and/or coordination of care for NF services in the code range

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99304 – 99306, 99307 – 99310, and 99318.

ADDITIONAL INFORMATION

You may find more information about using the prolonged services codes CPT codes (99356 and 99357) billing for medically necessary prolonged services for E/M visits in an SNF or NF, and for time-based counseling and/or coordination of care for NF services by going to CR 5968, located at <http://www.cms.hhs.gov/Transmittals/downloads/R1489CP.pdf> on the Centers for Medicare & Medicaid Services (CMS) Web site. You will find the updated *Medicare Claims Processing Manual*, chapter 12, (Physicians/Non-physician Practitioners), section 30.6.13 (Nursing Facility Services (Codes 99304 - 99318) subsection F (Use of the Prolonged Services Codes and Other Time-Related Services) as an attachment to that CR.

If you have any questions, please contact your carrier or A/B MAC at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5968

Related Change Request (CR) #: 5968

Related CR Release Date: April 11, 2008

Effective Date: July 1, 2008

Related CR Transmittal #: R1489CP

Implementation Date: July 7, 2008

MEDICARE PHYSICIAN FEE SCHEDULE DATABASE

APRIL UPDATE TO THE 2008 MEDICARE PHYSICIAN FEE SCHEDULE DATABASE

PROVIDER TYPES AFFECTED

Physicians and providers submitting claims to Medicare contractors (carriers, fiscal intermediaries (FIs), and/or Part A/B Medicare administrative contractors (A/B MACs) for professional services provided to Medicare beneficiaries that are paid under the Medicare physician fee schedule (MPFS).

PROVIDER ACTION NEEDED

This article is based on change request (CR) 5980 which amends payment files previously issued to Medicare contractors based upon the 2008 MPFS final rule. CR 5980 also includes new/revised codes for the Physician Quality Reporting Initiative (PQRI).

BACKGROUND

Attachment 1 of CR 5980 contains changes included in the April Update to the 2008 MPFSDB, and CR 5980 may be reviewed at <http://www.cms.hhs.gov/Transmittals/downloads/R1482CP.pdf> on the Centers for Medicare & Medicaid Services (CMS) Web site. Specific changes are detailed in Attachment 1 of CR 5980 and are summarized as follows:

CPT/HCPCS Code Revisions

A number of CPT/HCPCS codes have been modified to reflect revised bilateral indicators, relative value unit (RVU) revisions, or procedure status changes retroactive to January 1, 2008.

Reinstated "J" Codes

A number of "J" codes (J7611 through J7614) are reinstated with a status indicator of "E" and the reinstated codes are effective for dates of service on or after April 1, 2008. Descriptors and payment indicators for the reinstated codes are in attachment 1 of CR 5980.

New "Q" Codes

There are several new "Q" codes (Q4096 through Q4098) with a status indicator of "E" and which are effective for dates of service on or after April 1, 2008. The codes with their descriptors are in the following table:

April Update to the 2008 MPFSDB, continued

Benefits for Which Medicare Makes Payment	Services for Which Medicare Does NOT Pay – No Benefit Category
A conventional intraocular lens (IOL) implanted following cataract surgery.	The astigmatism-correcting functionality of an IOL implanted following cataract surgery.
Facility or physician services and supplies required to insert a conventional IOL following cataract surgery.	Facility or physician services and resources required to insert and adjust an AC-IOL following cataract surgery that exceeds the services and resources furnished for insertion of a conventional IOL.
One pair of eyeglasses or contact lenses as a prosthetic device furnished after each cataract surgery with insertion of an IOL.	The surgical correction of cylindrical lenses of eyeglasses or contact lenses that may be required to compensate for imperfect curvature of the cornea (astigmatism). Eye examinations performed to determine the refractive state of the eyes specifically associated with insertion of an AC-IOL (including subsequent monitoring services), that exceed the one-time eye examination following cataract surgery with insertion of a conventional IOL.

New Category II Codes for PQRI

There are new category II codes for the PQRI for dates of service on or after April 1, 2008. These new codes and their descriptors are in the following table:

Code	Long Descriptor	Short Descriptor
0525F	Initial visit for episode	Initial visit for episode
0526F	Subsequent visit for episode	Subs visit for episode
1130F	Back pain and function assessed, including all of the following: Pain assessment AND functional status AND patient history, including notation of presence or absence of "red flags" (warning signs) AND assessment of prior treatment and response, AND employment status	Bk pain + fxn assessed
1134F	Episode of back pain lasting six weeks or less	Epsd bk pain for =< 6 wks
1135F	Episode of back pain lasting longer than six weeks	Epsd bk pain for > 6 wks
1136F	Episode of back pain lasting 12 weeks or less	Epsd bk pain for <= 12 wks
1137F	Episode of back pain lasting longer than 12 weeks	Epsd bk pain for > 12 wks
2040F	Physical examination on the date of the initial visit for low back pain performed, in accordance with specifications	Bk pn xm on init visit date
2044F	Documentation of mental health assessment prior to intervention (back surgery or epidural steroid injection) or for back pain episode lasting longer than six weeks	Doc mntl tst b/4 bk trxmnt
3330F	Imaging study ordered	Imaging study ordered (bkp)
3331F	Imaging study not ordered	Bk imaging tst not ordered
3340F	Mammogram assessment category of "incomplete: need additional imaging evaluation", documented	Mammo assess inc xray docd
3341F	Mammogram assessment category of "negative", documented	Mammo assess negative docd
3342F	Mammogram assessment category of "benign", documented	Mammo assess bengn docd
3343F	Mammogram assessment category of "probably benign", documented	Mammo probably bengn docd
3344F	Mammogram assessment category of "suspicious", documented	Mammo assess susp docd
3345F	Mammogram assessment category of "highly suggestive of malignancy", documented	Mammo assess hghlymalig doc
3350F	Mammogram assessment category of "known biopsy proven malignancy", documented	Mammo bx proven malig docd
4240F	Instruction in therapeutic exercise with follow-up by the physician provided to patients during episode of back pain lasting longer than 12 weeks	Instr xrcz 4bk pn >12 weeks
4242F	Counseling for supervised exercise program provided to patients during episode of back pain lasting longer than 12 weeks	Sprvsd xrcz bk pn >12 weeks
4245F	Patient counseled during the initial visit to maintain or resume normal activities	Pt instr nrml lifest

April Update to the 2008 MPFSDB, continued

Code	Long Descriptor	Short Descriptor
4248F	Patient counseled during the initial visit for an episode of back pain against bed rest lasting 4 days or longer	Pt instr--no bd rest>= 4 days
4250F	Active warming used intraoperatively for the purpose of maintaining normothermia, OR at least one body temperature equal to or greater than 36 degrees Centigrade (or 96.8 degrees Fahrenheit) recorded within the 30 minutes immediately before or the 30 minutes immediately after anesthesia end time	Wrmng 4 surg - normothermia
5060F	Findings from diagnostic mammogram communicated to practice managing patient's on-going care within 3 business days of exam interpretation	Fndngs mammo 2pt w/in 3 days
5062F	Findings from diagnostic mammogram communicated to the patient within 5 days of exam interpretation	Doc f2fmammo fndng in 3 days
6040F	Use of appropriate radiation dose reduction devices OR manual techniques for appropriate moderation of exposure, documented	Appro rad ds dvcs techs docd
6045F	Radiation exposure or exposure time in final report for procedure using fluoroscopy, documented	Radxps in end rpt4fluro pxd
7020F	Mammogram assessment category [eg, Mammography Quality Standards Act (MQSA), Breast Imaging Reporting and Data System (BI-RADS®), or FDA approved equivalent categories] entered into an internal database to allow for analysis of abnormal interpretation (recall) rate	Mammo assess cat in dbase
7025F	Patient information entered into a reminder system with a target due date for the next mammogram	Pt infosys alarm 4 nxt mammo

Revised Descriptors for PQRI Codes

Attachment 1 of CR 5980 also contains a list of editorial changes to the short and/or long descriptors for a number of PQRI codes.

Connecticut Fee Revisions

CPT/Mod	Par	Non-Par	Limiting Charge
93501	979.28	930.32	1069.86
93501 TC	805.58	765.30	880.10
93508	1,156.12	1098.31	1263.06
93508 TC	908.64	863.21	992.69
93510	1,757.26	1669.40	1919.81
93510 TC	1,496.67	1421.84	1635.11
93526	2,271.31	2157.74	2481.41
93526 TC	1,909.71	1814.22	2086.36
93642	566.57	538.24	618.98

Florida Fee Revisions

CPT/Mod	Participating			Nonparticipating			Limiting Charge		
	Loc 01/02	Loc 03	Loc 04	Loc 01/02	Loc 03	Loc 04	Loc 01/02	Loc 03	Loc 04
93501	832.13	902.26	976.62	790.52	857.15	927.79	909.10	985.72	1066.96
93501 TC	670.37	733.22	798.84	636.85	696.56	758.90	732.38	801.04	872.73
93508	969.27	1041.86	1113.75	920.81	989.77	1058.06	1058.93	1138.23	1216.77
93508 TC	740.62	802.73	862.31	703.59	762.59	819.19	809.13	876.98	942.07
93510	1496.13	1629.50	1773.49	1421.32	1548.02	1684.82	1634.52	1780.23	1937.54
93510 TC	1255.16	1377.42	1508.34	1192.40	1308.55	1432.92	1371.26	1504.83	1647.86
93526	1938.64	2111.61	2299.36	1841.71	2006.03	2184.39	2117.96	2306.93	2512.05
93526 TC	1604.19	1761.68	1931.17	1523.98	1673.60	1834.61	1752.58	1924.64	2109.80
93642	495.16	526.88	560.46	470.40	500.54	532.44	540.96	575.62	612.30

ADDITIONAL INFORMATION

The official instruction, CR 5980, issued to your carrier, FI, and A/B MAC regarding this change may be viewed at <http://www.cms.hhs.gov/Transmittals/downloads/R1482CP.pdf> on the CMS Web site.

If you have any questions, please contact your carrier, FI, or A/B MAC at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

April Update to the 2008 MPFSDB, continued

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SURGERY**ASTIGMATISM-CORRECTING INTRAOCULAR LENS—IMPLEMENTATION OF CMS-1536 RULING**

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals. This information was previously published in the June 2007 Medicare B Update! pages 19-22.

Note: This article was revised on April 10, 2008, to add a reference to *MLN Matters* article MM5853 (<http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM5853.pdf>), which provides instructions regarding the use of HCPCS code V2787 when billing for intraocular lens procedures and services involving recognized A-C IOLs that take place in ambulatory surgery centers (ASCs), physician offices, or hospital outpatient departments (HOPDs).

PROVIDER TYPES AFFECTED

Physicians, providers, and suppliers submitting claims to Medicare contractors (carriers, fiscal intermediaries [FIs], or Part A/B Medicare administrative contractors [A/B MACs] for services provided to Medicare beneficiaries

PROVIDER ACTION NEEDED

This article is based on change request (CR) 5527 which discusses a recent administrator ruling from the Centers for Medicare & Medicaid Services (CMS) regarding astigmatism-correcting intraocular lenses (A-C IOLs) following cataract surgery (CMS-1536-R). The new policy is effective for dates of service on and after January 22, 2007. Physicians and providers need to be aware that effective January 22, 2007:

- Medicare will pay the same amount for cataract extraction with A-C IOL insertion that it pays for cataract extraction with conventional IOL insertion.
- The beneficiary is responsible for payment of that portion of the hospital or ambulatory surgery center (ASC) charge for the procedure that exceeds the facility's usual charge for cataract extraction and insertion of a conventional IOL following cataract surgery, as well as any fees that exceed the physician's usual charge to perform a cataract extraction with insertion of a conventional IOL.

In addition, CMS reminds physicians that they can be reimbursed for the conventional or A-C IOL (V2632) only when the service is performed in a physician's office. Also, when physicians perform cataract surgery in an ASC or hospital outpatient setting, the physician may only bill for the professional service because payment for the lens is bundled into the facility payment for the cataract extraction.

BACKGROUND

The Centers for Medicare & Medicaid Services (CMS) Administrator rulings serve as 1) precedent final opinions and orders and 2) statements of policy and interpretation. The Administrator rulings provide clarification and interpretation of complex or ambiguous provisions of the law or regulations relating to Medicare, Medicaid, utilization and peer review by quality improvement organizations, private health insurance, and related matters. These rulings also promote consistency in interpretation of policy and adjudication of disputes, and they are binding on all CMS components, Medicare contractors, the Provider Reimbursement Review Board, the Medicare Geographic Classification Review Board, and administrative law judges who hear Medicare appeals.

CR 5527 discusses a recent CMS administrator ruling concerning requirements for determining payment for insertion of intraocular lenses (IOLs) that replace beneficiaries' natural lenses and correct pre-existing astigmatism following cataract surgery under the Social Security Act:

Note: CR 5527 basically restates CMS policy provided in CR 3927 (*MLN Matters* article MM3927), except that CR 3927 focused on presbyopia-correcting IOLs and this article focuses on A-C IOLs.

Coverage Policy

In general, an item or service covered by Medicare must satisfy the following three basic requirements:

- Fall within a statutorily-defined benefit category
- Be reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body part
- Not be excluded from coverage.

Astigmatism-Correcting IOL—Implementation of CMS-1536 Ruling, continued

The Social Security Act specifically excludes eyeglasses and contact lenses from coverage, with an exception for one pair of eyeglasses or contact lenses covered as a prosthetic device furnished after each cataract surgery with insertion of an IOL. In addition, there is no Medicare benefit category to allow payment for the surgical correction or cylindrical lenses of eyeglasses or contact lenses that may be required to compensate for the imperfect curvature of the cornea (astigmatism).

An A-C IOL is intended to provide what is otherwise achieved by two separate items:

- An implantable conventional IOL (one that is not astigmatism - correcting) that is covered by Medicare
- The surgical correction, eyeglasses, or contact lenses that are not covered by Medicare.

Although A-C IOLs may serve the same function as eyeglasses or contact lenses furnished following removal of a cataract, A-C IOLs are neither eyeglasses nor contact lenses. The following table is a summary of benefits for which Medicare makes payment, and services for which Medicare does not pay (no benefit category):

Benefits for Which Medicare Makes Payment	Services for Which Medicare Does NOT Pay – No Benefit Category
A conventional intraocular lens (IOL) implanted following cataract surgery.	The astigmatism-correcting functionality of an IOL implanted following cataract surgery.
Facility or physician services and supplies required to insert a conventional IOL following cataract surgery.	Facility or physician services and resources required to insert and adjust an AC-IOL following cataract surgery that exceeds the services and resources furnished for insertion of a conventional IOL.
One pair of eyeglasses or contact lenses as a prosthetic device furnished after each cataract surgery with insertion of an IOL.	The surgical correction of cylindrical lenses of eyeglasses or contact lenses that may be required to compensate for imperfect curvature of the cornea (astigmatism). Eye examinations performed to determine the refractive state of the eyes specifically associated with insertion of an AC-IOL (including subsequent monitoring services), that exceed the one-time eye examination following cataract surgery with insertion of a conventional IOL.

Currently, there is one NTIOL class approved for special payment when furnished by an ASC, and this currently active NTIOL category for "Reduced Spherical Aberration" was established on February 27, 2006, and expires on February 26, 2011.

Effective for services furnished on or after January 22, 2007, CMS now recognizes the following as A-C IOLs:

- Acrysof® Toric IOL (models: SN60T3, SN60T4, and SN60T5), manufactured by Alcon Laboratories, Inc
- Silicon 1P Toric IOL (models: AA4203TF and AA4203TL), manufactured by STAAR Surgical.

Payment Policy for Facility Services and Supplies

The following applies to an IOL inserted following removal of a cataract in a hospital (on either an outpatient or inpatient basis) that is paid under 1) the hospital outpatient prospective payment system (OPPS) or 2) the inpatient prospective payment system (IPPS), respectively (or in a Medicare-approved ASC that is paid under the ASC fee schedule):

- Medicare does not make separate payment to the hospital or the ASC for an IOL inserted subsequent to extraction of a cataract. Payment for the IOL is packaged into the payment for the surgical cataract extraction/lens replacement procedure.
- Any person or ASC, who presents or causes to be presented a bill or request for payment for an IOL inserted during or subsequent to cataract surgery for which payment is made under the ASC fee schedule, is subject to a civil money penalty.

For an A-C IOL inserted subsequent to removal of a cataract in a hospital (on either an outpatient or inpatient basis) that is paid under the OPPS or the IPPS, respectively (or in a Medicare-approved ASC that is paid under the ASC fee schedule):

- The facility should bill for removal of a cataract with insertion of a conventional IOL, regardless of whether a

conventional or A-C IOL is inserted. When a beneficiary receives an A-C IOL following removal of a cataract, hospitals and ASCs should report the same CPT code that is used to report removal of a cataract with insertion of a conventional IOL (see "Coding" below).

- There is no Medicare benefit category that allows payment of facility charges for services and supplies required to insert and adjust an A-C IOL following removal of a cataract that exceed the facility charges for services and supplies required for the insertion and adjustment of a conventional IOL.
- There is no Medicare benefit category that allows payment of facility charges for subsequent treatments, services and supplies required to examine and monitor the beneficiary who receives an AC-IOL following removal of a cataract that exceed the facility charges for subsequent treatments, services, and supplies required to examine and monitor a beneficiary after cataract surgery followed by insertion of a conventional IOL.

Payment Policy for Physician Services and Supplies

For an IOL inserted following removal of a cataract in a physician's office Medicare makes separate payment, based on reasonable charges, for an IOL inserted subsequent to extraction of a cataract that is performed at a physician's office.

For an A-C IOL inserted following removal of a cataract in a physician's office:

- A physician should bill for a conventional IOL, regardless of whether a conventional or A-C IOL is inserted (see "Coding," below).
- There is no Medicare benefit category that allows payment of physician charges for services and supplies required to insert and adjust an A-C IOL following removal of a cataract that exceed the physician charges for services and supplies for the insertion and adjustment of a conventional IOL.

Astigmatism-Correcting IOL—Implementation of CMS-1536 Ruling, continued

- There is no Medicare benefit category that allows payment of physician charges for subsequent treatments, services, and supplies required to examine and monitor a beneficiary following removal of a cataract with insertion of an AC-IOL that exceed the physician charges for services and supplies to examine and monitor a beneficiary following removal of a cataract with insertion of a conventional IOL.

For an A-C IOL inserted following removal of a cataract in a hospital or ASC:

- A physician may not bill Medicare for the A-C IOL inserted during a cataract procedure performed in those settings because payment for the lens is included in the payment made to the facility for the entire procedure.
- There is no Medicare benefit category that allows payment of physician charges for services and supplies required to insert and adjust an A-C IOL following removal of a cataract that exceed physician charges for services and supplies required for the insertion of a conventional IOL.
- There is no Medicare benefit category that allows payment of physician charges for subsequent treatments, services, and supplies required to examine and monitor a beneficiary following removal of a cataract with insertion of an A-C IOL that exceed the physician charges for services and supplies required to examine and monitor a beneficiary following cataract surgery with insertion of a conventional IOL.

Coding

No new codes are being established at this time to identify an A-C IOL or procedures and services related to an A-C IOL, and hospitals, ASCs, and physicians should report one of the following CPT codes to bill Medicare for removal of a cataract with IOL insertion:

66982 - *Extracapsular cataract removal with insertion of intraocular lens prosthesis (one stage procedure), manual or mechanical technique (e.g., irrigation and aspiration or phacoemulsification), complex, requiring devices or techniques not generally used in routine cataract surgery (e.g., iris expansion device, suture support for intraocular lens, or primary posterior capsulorrhexis) or performed on patients in the amblyogenic developmental stage*

66983 - *Intracapsular cataract extraction with insertion of intraocular lens prosthesis (one stage procedure)*

66984 - *Extracapsular cataract removal with insertion of intraocular lens prosthesis (one stage procedure), manual or mechanical technique (e.g., irrigation and aspiration or phacoemulsification).*

Physicians inserting an IOL or an A-C IOL in an office setting may bill code V2632 (posterior chamber intraocular lens) for the IOL or the A-C IOL, which is paid on a reasonable charge basis.

If appropriate, hospitals and physicians may use the proper CPT code(s) to bill Medicare for evaluation and management services usually associated with services following cataract extraction surgery, if appropriate.

Beneficiary Liability

When a beneficiary requests insertion of an A-C IOL instead of a conventional IOL following removal of a cataract and that procedure is performed, the beneficiary is responsible for payment of facility charges for services and supplies attributable to the astigmatism-correcting functionality of the A-C IOL:

- In determining the beneficiary's liability, the facility and physician may take into account any additional work and resources required for insertion, fitting, vision acuity testing, and monitoring of the AC-IOL that exceeds the work and resources attributable to insertion of a conventional IOL.
- The physician and the facility may not charge for cataract extraction with insertion of an A-C IOL unless the beneficiary requests this service.
- The physician and the facility may not require the beneficiary to request an A-C IOL as a condition of performing a cataract extraction with IOL insertion.

Provider Notification Requirements

When a beneficiary requests insertion of an A-C IOL instead of a conventional IOL following removal of a cataract:

- Prior to the procedure to remove a cataractous lens and insert an A-C IOL, the facility and the physician must inform the beneficiary that Medicare will not make payment for services that are specific to the insertion, adjustment, or other subsequent treatments related to the astigmatism-correcting functionality of the IOL.
- The correcting functionality of an A-C IOL does not fall into a Medicare benefit category and, therefore, is not covered. Therefore, the facility and physician are not required to provide an advanced beneficiary notice to beneficiaries who request an A-C IOL.
- Although not required, CMS strongly encourages facilities and physicians to issue a notice of exclusion from Medicare benefits to beneficiaries in order to identify clearly the non-payable aspects of an A-C IOL insertion. This notice may be found on the CMS Web site at:

- English version: <http://www.cms.hhs.gov/BNI/downloads/CMS20007English.pdf>
- Spanish version: <http://www.cms.hhs.gov/BNI/downloads/CMS20007Spanish.pdf>

ADDITIONAL INFORMATION

The official instruction, CR 5527, issued to your Medicare carrier, intermediary, and A/B MAC regarding this change may be viewed at <http://www.cms.hhs.gov/Transmittals/downloads/R1228CP.pdf> on the CMS Web site.

If you have any questions, please contact your Medicare carrier, intermediary, or A/B MAC at their toll-free number, which may be found on the CMS Web site at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

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GENERAL COVERAGE

ERYTHROPOIESIS STIMULATING AGENTS IN CANCER AND RELATED NEOPLASTIC CONDITIONS

CMS has issued the following MLN Matters article. *Information for Medicare Fee-for-Service Health Care Professionals.*

Note: This article was April 25, 2008, to correct the third bullet under *Reasonable and Necessary ESA Use* regarding the "Maintenance of ESA therapy" (See bullet in **bold**). It should have stated that the "starting dose if the hemoglobin level remains below 10 g/dL (or hematocrit is < 30 percent) four weeks after initiation of therapy and the rise in hemoglobin is \geq 1g/dL (hematocrit \geq 3 percent)." All other information remains the same.

PROVIDER TYPES AFFECTED

Providers and suppliers who bill Medicare contractors (carriers, fiscal intermediaries [FI], regional home health intermediaries [RHHI], Medicare administrative contractors [A/B MAC] and durable medical equipment Medicare administrative contractors [DME MAC]) for administering or supplying erythropoiesis stimulating agents (ESAs) for cancer and related neoplastic conditions to Medicare beneficiaries.

WHAT YOU NEED TO KNOW

Following a national coverage analysis (NCA) to evaluate the uses of ESAs in non-renal disease applications, the Centers for Medicare & Medicaid Services (CMS), on July 30, 2007, issued a decision memorandum (DM) that addressed ESA use in non-renal disease applications (specifically in cancer and other neoplastic conditions).

Change request 5818 communicates the NCA findings and the coverage policy in the national coverage determination (NCD). Specifically, CMS determines that ESA treatment is reasonable and necessary for anemia secondary to myelosuppressive anticancer chemotherapy in solid tumors, multiple myeloma, lymphoma, and lymphocytic leukemia under specified conditions; and not reasonable and necessary for beneficiaries with certain other clinical conditions, as listed below.

The HCPCS codes specific to non-end-stage renal disease (ESRD) ESA use are J0881 and J0885. Claims processed with dates of service July 30, 2007, through December 31, 2007, do not have to include the ESA modifiers, as the modifiers are not effective until January 1, 2008. However, providers are to begin using the modifiers as of January 1, 2008, even though full implementation of related system edits are not effective until April 7, 2008.

Make sure that your billing staffs are aware of this guidance regarding ESA use.

BACKGROUND

Emerging safety concerns (thrombosis, cardiovascular events, tumor progression, and reduced survival) derived from clinical trials in several cancer and non-cancer populations prompted CMS to review its coverage of ESAs. In so doing, on March 14, 2007, CMS opened an NCA to evaluate the uses of ESAs in non-renal disease applications, and on July 30, 2007, issued a DM specifically narrowed to the use of ESAs in cancer and other neoplastic conditions.

Reasonable and Necessary ESA Use

CMS has determined that ESA treatment for the anemia secondary to a regimen of myelosuppressive anticancer chemotherapy in solid tumors, multiple myeloma, lymphoma, and lymphocytic leukemia is reasonable and necessary only under the following specified conditions:

- The hemoglobin level immediately prior to the first administration is < 10 g/dL (or the hematocrit is < 30 percent) and the hemoglobin level prior to any maintenance administration is < 10g/dL (or the hematocrit is < 30 percent).
 - The starting dose for ESA treatment is up to either of the recommended Food and Drug Administration (FDA) approved label starting doses for cancer patients receiving chemotherapy, which includes the, 150 U/kg/3 times weekly or the 40,000 U weekly doses for epoetin alfa and the 2.25 mcg/kg/weekly or the 500 mcg once every three week dose for darbepoetin alpha.
 - **Maintenance of ESA therapy is the starting dose if the hemoglobin level remains below 10 g/dL (or hematocrit is < 30 percent) 4 weeks after initiation of therapy and the rise in hemoglobin is \geq 1g/dL (hematocrit \geq 3 percent).**
 - For patients whose hemoglobin rises < 1 g/dl (hematocrit rise < 3 percent) compared to pretreatment baseline over 4 weeks of treatment and whose hemoglobin level remains < 10 g/dL after 4 weeks of treatment (or the hematocrit is < 30 percent), the recommended FDA label starting dose may be increased once by 25 percent. Continued use of the drug is not reasonable and necessary if the hemoglobin rises < 1 g/dl (hematocrit rise < 3 percent) compared to pretreatment baseline by 8 weeks of treatment.
 - Continued administration of the drug is not reasonable and necessary if there is a rapid rise in hemoglobin > 1 g/dl (hematocrit > 3 percent) over any two week period of treatment unless the hemoglobin remains below or subsequently falls to < 10 g/dL (or the hematocrit is < 30 percent). Continuation and reinstitution of ESA therapy must include a dose reduction of 25 percent from the previously administered dose.
 - ESA treatment duration for each course of chemotherapy includes the eight weeks following the final dose of myelosuppressive chemotherapy in a chemotherapy regimen.
- Not Reasonable and Necessary ESA Use**
- Either because of a deleterious effect of ESAs on the underlying disease, or because the underlying disease increases the risk of adverse effects related to ESA use, CMS has also determined that ESA treatment is not reasonable and necessary for beneficiaries with the following clinical conditions:
- Any anemia in cancer or cancer treatment patients due to folate deficiency (diagnosis code 281.2), B-12 deficiency (281.1 or 281.3), iron deficiency (280.0-280.9), hemolysis (282.0, 282.2, 282.9, 283.0, 283.2, 283.9, 283.10, 283.19), bleeding (280.0 or 285.1), or

ESAs in Cancer and Related Neoplastic Conditions, continued

- bone marrow fibrosis.
- Anemia associated with the treatment of acute and chronic myelogenous leukemias (CML, AML) (205.00-205.21, 205.80-205.91), or erythroid cancers (207.00-207.81)
- Anemia of cancer not related to cancer treatment
- Any anemia associated only with radiotherapy
- Prophylactic use to prevent chemotherapy-induced anemia
- Prophylactic use to reduce tumor hypoxia
- Erythropoietin-type resistance due to neutralizing antibodies
- Anemia due to cancer treatment if patients have uncontrolled hypertension.

Claims Processing

Effective for claims with dates of service on or after January 1, 2008, Medicare will deny non-ESRD ESA services for J0881 or J0885 when:

- Billed with modifier EC (ESA, anemia, non-chemo/radio) when a diagnosis on the claim is present for any anemia in cancer or cancer treatment patients due to folate deficiency (diagnosis code 281.2), B-12 deficiency (281.1 or 281.3), iron deficiency (280.0-280.9), hemolysis (282.0, 282.2, 282.9, 283.0, 283.2, 283.9, 283.10, 283.19), bleeding (280.0 or 285.1), anemia associated with the treatment of acute and chronic myelogenous leukemias (CML, AML) (205.00-205.21, 205.80-205.91), or erythroid cancers (207.00-207.81).
- Billed with modifier EC for any anemia in cancer or cancer treatment patients due to bone marrow fibrosis, anemia of cancer not related to cancer treatment, prophylactic use to prevent cancer-induced anemia, prophylactic use to reduce tumor hypoxia, erythropoietin-type resistance due to neutralizing antibodies, and anemia due to cancer treatment if patients have uncontrolled hypertension.
- Billed with modifier EA (ESA, anemia, chemo-induced) for anemia secondary to myelosuppressive anticancer chemotherapy in solid tumors, multiple myeloma, lymphoma, and lymphocytic leukemia when a hemoglobin 10.0g/dL or greater or hematocrit 30.0 percent or greater is reported.
- Billed with modifier EB (ESA, anemia, radio-induced).

Note: Denial of claims for non-ESRD ESAs for cancer and related neoplastic indications as outlined in NCD 110.21 are based on reasonable and necessary determinations. A provider may have the beneficiary sign an advance beneficiary notice (ABN), making the beneficiary liable for services not covered by Medicare. When denying ESA claims, contractors will use Medicare Summary Notice 15.20. The following policies [NCD 110.21] were used when we made this decision, and remittance reason code 50. These are noncovered services because this is not deemed a 'medical necessity' by the payer. However, standard systems shall assign liability for the denied charges to the provider unless documentation of the ABN is present on the claim. Denials are subject to appeal and standard systems shall allow for medical review override of denials. Contractors may reverse the denial if the review results in a determination of clinical necessity.

Medicare contractors have discretion to establish local coverage policies for those indications not included in NCD 110.21.

Medicare contractors will not search files to retract payment for claims paid prior to April 7, 2008. However, contractors shall adjust claims brought to their attention.

ADDITIONAL INFORMATION

If you have any questions, please contact your Medicare contractor at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

This addition/revision of section 110.21 of Pub.100-03 is an NCD. NCDs are binding on all carriers, FIs, quality improvement organizations, qualified independent contractors, the Medicare Appeals Council, and administrative law judges (ALJs) (see 42 CFR section 405.1060[a][4] [2005]). An NCD that expands coverage is also binding on a Medicare advantage organization. In addition, an ALJ may not review an NCD. (See section 1869[f][1][A][I] of the Social Security Act.)

The official instruction, CR 5818, was issued to your contractor in two transmittals. The first is the NCD transmittal and that is available at <http://www.cms.hhs.gov/Transmittals/downloads/R80NCD.pdf> on the CMS Web site. The second transmittal revises the *Medicare Claims Processing Manual* and it is at <http://www.cms.hhs.gov/Transmittals/downloads/R1413CP.pdf> on the same site.

MLN Matters Number: MM5818 *Revised*

Related Change Request (CR) #: 5818

Related CR Release Date: January 14, 2008

Effective Date: July 30, 2007

Related CR Transmittal #: R80NCD and R1413CP

Implementation Date: April 7, 2008

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NATIONAL PROVIDER IDENTIFIER

MAY 23RD IS ONLY TWO WEEKS AWAY—ARE YOU PREPARED?

NPI IS HERE. NPI IS NOW. ARE YOU USING IT? INFORMATION FOR ALL HEALTH CARE PROVIDERS MEDICARE AND NON-MEDICARE

The Centers for Medicare & Medicaid Services (CMS) encourages all health care providers to contact other health plans with which you interact in order to ensure you fully understand their expectations for May 23, 2008.

IMPORTANT INFORMATION FOR MEDICARE FEE-FOR-SERVICE PROVIDERS

Clarification of April 3, 2008, Statement “Institutional Providers Submitting Taxonomy Codes to Identify Subparts – What Medicare Is Using to Obtain NPI/OSCAR Match”

Providers who submit Medicare claims may continue to send their Medicare provider taxonomy codes. However, Medicare fee-for-service claim processing systems will not use this data to adjudicate claims. The taxonomy codes will be crossed over to the secondary payers as CMS understands that some payers may use this information to adjudicate claims.

When To Update NPPES if an Update to Medicare Enrollment Information Is Also Needed

The NPI final rule requires covered providers to update their required NPPES data within 30 days of the change. If a Medicare provider needs to update information in NPPES, it will also need to update the corresponding information in its Medicare enrollment record via the CMS-855. Providers should not make updates to NPPES data until after their CMS-855s are processed and those updates are effective in the Medicare enrollment system (PECOS, or the NSC for Medicare DMEPOS suppliers). After the update is effective in PECOS or the NSC (whichever is appropriate), providers have up to 30 days to make the corresponding updates in NPPES. In a change of ownership (CHOW) situation, for example, the new owner would not make changes in the NPPES record of the provider that is being sold until after the CMS-855 is processed and its changes are effective in the Medicare enrollment system. If a new NPI is to be obtained as part of the CHOW and an existing NPI is to be deactivated (those decisions are up to the buyer and the seller), the NPI should not be deactivated until after all claims using that NPI reach final settlement (this could involve health plans in addition to Medicare).

May 23rd Is Only Two Weeks Away, ARE YOU PREPARED?

Urgent: CMS continues to be concerned about the low percentage of claims being submitted with an NPI alone in the primary provider identifier fields. See below for specific steps to begin using the NPI alone in the primary provider identifier fields.

Don't Be Surprised on May 23...TRY NPI-ONLY NOW

Now that the NPI is required on all Medicare claims in the primary provider fields, if your claims are being successfully processed with NPI/legacy pairs (and most are) now is the time to begin sending a small batch of claims with NPI alone. If the Medicare NPI crosswalk cannot match your NPI to your Medicare legacy number, the claim with an NPI-only will reject. You can and should try sending NPI-only now! If the claim is processed and you are paid, continue to increase the volume of claims sent

with only your NPI. If the claims reject, go into your NPPES record and validate that the information you are sending on the claim is consistent with the information in NPPES. If it is different, make the updates in NPPES and resend a small batch of claims three-four days later. If your claims are still rejecting, you may need to update your Medicare enrollment information to correct this problem. Call the Customer Service Representative at your Medicare carrier, FI, or A/B MAC or at your DME MAC to discuss your situation and, if necessary, have it investigated. Have a copy of your NPPES record or your NPI registry record available. The contractor telephone numbers are likely to be quite busy, so don't wait.

If you bill Medicare using a billing service or clearinghouse, you should work with them to establish a way to try sending NPI-only claims. It may be difficult for some of these third party vendors to send small batches of your NPI-only claims and continue sending NPI and legacy claims as well, so contact them and develop an alternative solution so you can try NPI-only.

Sending a sample of NPI-only claims will allow time for any needed corrections prior to May 23, 2008, the date when only the NPI will be accepted in all provider fields.

NPIs in Secondary Provider Fields

May 23, 2008, is also the deadline for using the NPI-only in the secondary provider identifier fields on a claim transaction. This includes the prescriber field in a Medicare fee-for-service retail pharmacy drug claim submitted in an NCPDP 5.1 transaction. CMS will be providing guidance with respect to the reporting of NPIs in the Service Facility Location loop in the X12 N 837 claim transactions.

NPIs on ALL HIPAA Standard Transactions

May 23, 2008, is also the deadline for the use of NPI on ALL HIPAA standard transactions (e.g., 837I, 837P, NCPDP, DDE, 276/277, 270/271 and 835).

NEED MORE INFORMATION?

Still not sure what an NPI is and how you can get it, share it and use it? As always, more information and education on the NPI may be found at the CMS NPI Web page <http://www.cms.hhs.gov/NationalProviderStand>.

Providers can apply for an NPI online at <https://nppes.cms.hhs.gov> or can call the NPI enumerator to request a paper application at 1-800-465-3203.

Having trouble viewing any of the URLs in this message? If so, try to cut and paste any URL in this message into your Web browser to view the intended information.

Note: All current and past CMS NPI communications are available by clicking “CMS Communications” in the left column of the CMS Web page <http://www.cms.hhs.gov/NationalProvIdentStand>.

Source: CMS Provider Education Resource 200804-13

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STEPS TO FACILITATE A SMOOTH TRANSITION TO NPI-ONLY MEDICARE BILLING AND MORE NPI IS HERE. NPI IS NOW. ARE YOU USING IT?

INFORMATION FOR ALL HEALTH CARE PROVIDERS MEDICARE AND NON-MEDICARE

The Centers for Medicare & Medicaid Services (CMS) encourages all health care providers to ensure they understand the readiness of other health plans with which they interact, especially if those health plans may be primary or secondary to Medicare. Medicare will only accept/send NPI-only transactions beginning May 23, 2008, and providers need to understand from these other plans what will happen if they are unable to send/receive **NPI-only** transactions.

IMPORTANT INFORMATION FOR MEDICARE FEE-FOR-SERVICE PROVIDERS

CMS is pleased to announce that Medicare is receiving more than 98 percent of claims with an NPI. The next milestone – May 23rd – requires providers to take the next step so they do not risk disruption in cash flow. Begin billing with **NPI-only** now to test how May 23rd will impact you.

CMS is concerned that the percentage of Medicare claims with **NPI-only** is not growing fast enough.

STEPS TO FACILITATE A SMOOTH TRANSITION TO NPI-ONLY

- 1) Bill with Medicare legacy ID & NPI
 - Once claims are successfully processed, move to Step 2.
- 1) Bill with NPI-only
 - Start with a small batch of claims. If, or when, the results are positive, begin sending a greater volume and move to Step 3.
 - Billing with NPI-only also tests the ability to receive the NPI on 835 transactions.
- 1) Test NPI-only on other HIPAA transactions
 - CMS will require use of the NPI on the 270/271, 276/277 and NCPDP transactions. Providers should begin testing the use of the NPI on these transactions, in small quantities, prior to May 23rd to ensure a smooth transition. Also, be prepared to accept the NPI-only on the 835-remittance advice transaction.

INSTITUTIONAL PROVIDERS SUBMITTING TAXONOMY CODES TO IDENTIFY SUBPARTS

UPDATE: MEDICARE IS USING ALTERNATIVE DATA TO OBTAIN NPI/OSCAR MATCH

On January 1, 2007, Medicare implemented change request (CR) 5243, which required the submission of taxonomy codes all claims submitted by institutional Medicare providers who submit claims for their primary facility and its subparts (such as psychiatric unit, rehabilitation unit, etc.).

The intent of CR 5243 was to enable Medicare to appropriately crosswalk a provider NPI to each of the provider's subparts through the reporting of taxonomy codes in the claims.

Medicare has found that using taxonomy codes has been unsuccessful in obtaining a one-to-one match on the crosswalk for those providers having one NPI tied to multiple OSCAR/certification numbers. As a result, the taxonomy code is no longer used as part of the crosswalk criteria that are used to attempt to match an NPI with an OSCAR/certification number. Currently, the fiscal

intermediary shared system (FISS) uses these matching criteria to obtain a one-to-one match between an institutional Medicare provider's NPI and its OSCAR/certification number:

- First level of match: **Type of bill (TOB) to OSCAR/certification number.** If the system is unable to identify a valid match, the search will continue with the next level of match.
- Second level of match: **Revenue code to OSCAR/certification number.** If the system is unable to identify a valid match, the search will continue with the next level of match.
- Third and final level of match: **Facility ZIP code on the claim.** This final level prompts the systems logic to limit the list of appropriate OSCAR numbers by matching the facility ZIP code on the claim against the ZIP code of the master address in the FISS provider address file.

Note: If the system is unable to make a valid match, the claim will suspend with reason code 32105, and the provider will receive an additional development letter (ADR) requesting the OSCAR number.

ACTION REQUIRED BY INSTITUTIONAL PROVIDERS WITH SUBPARTS

Providers are strongly encouraged to enumerate their subparts. The following documents may assist providers in answering additional questions on this subject.

- CMS Medicare Subpart Expectations paper may be accessed on the CMS Web site at <http://www.cms.hhs.gov/NationalProvidentStand/Downloads/Medsubparts012>
- Read NPI Fact Sheet titled "For Health Care Providers Who are Organizations" at http://www.cms.hhs.gov/NationalProvidentStand/Downloads/NPI_FactSheet_03-07.pdf
- Review the article titled "Information Regarding National Plan and Provider Enumeration System Errors, Using the NPI on Medicare Claims and 835 Remittance Advice Changes" available in the August 2007 *Medicare A Bulletin* at http://www.floridamedicare.com/Part_A/Medicare_A_Bulletins/Archive/11004
- Review special edition article <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0608.pdf>
- National Plan and Provider Enumeration System (NPPES) <https://nppes.cms.hhs.gov/NPPES>.

ENCOURAGE CLEARINGHOUSES TO ALLOW TESTING OF NPI-ONLY

It has come to CMS' attention that some clearinghouses may not allow important NPI-only testing prior to May 23rd. CMS encourages Medicare providers to work with their clearinghouses to allow use of the NPI-only to facilitate this testing. If you do not test, you will not be aware, in advance, of any problems that could prohibit Medicare from processing and paying claims.

Registration Available for NPI Roundtable CMS subject matter experts will be available to address questions from the provider community on April 17, 2008 from 2-3:30PM ET. Participants are able to submit questions using the online registration system for this call. To register, visit http://www.cms.hhs.gov/NationalProvidentStand/Downloads/listserv_wording_4-10-08_call.pdf on the CMS Web site.

Steps to Facilitate a Smooth Transition to NPI-Only Medicare Billing and More, continued**NEED MORE INFORMATION?**

Not sure what an NPI is and how you can get it, share it and use it? As always, more information and education on the NPI may be found at the CMS NPI Web page <http://www.cms.hhs.gov/NationalProvidentStand>.

Providers can apply for an NPI online at <https://nppes.cms.hhs.gov> or can call the NPI enumerator to request a paper application at 1-800-465-3203.

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Note: All current and past CMS NPI communications are available by clicking "CMS Communications" in the left column of the CMS Web page <http://www.cms.hhs.gov/NationalProvidentStand>.

Source: CMS Provider Education Resource 200804-02

Visit The Medicare Learning Network – It's Free!

SELECTING A PREFERRED PROVIDER IDENTIFICATION NUMBER

The purpose of this article is to assist providers who are receiving claim development letters because they have multiple provider identification numbers (PINs) tied to one national provider identifier (NPI). Providers now have an option to select a "preferred PIN."

While the preferred PIN process was established to eliminate development to providers as much as possible, it will not solve all one NPI to multiple PINs situation, nor will it work for all providers. If the provider is an organization, it must separately enumerate its subparts that bill Medicare electronically. Individuals in the one-to-many situation that do not qualify for the preferred PIN will ultimately have no choice but to deactivate the PINs they no longer need to avoid receiving development letters.

DO YOU QUALIFY?

First, determine if you qualify to use the preferred PIN process.

PROVIDERS WHO DON'T QUALIFY

- Providers/group practices with multiple PIN(s), each having their own unique specialty
- Providers billing for services that are paid off reasonable charges rather than fee schedules and

have multiple PIN(s) established because their practice locations cross multiple localities.

Providers Who Might Not Qualify

- Providers rendering services in a health professional shortage area (HPSA) location as they may be negatively impacted if a preferred PIN is used.
- Independent diagnostic testing facility (IDTF) providers, as they are independently credentialed based on their location. Each IDTF location has a separate table of HCPCS/CPT procedure codes and corresponding equipment they are allowed to bill. If the procedure code submitted on the claim is not on the table, the claim would be denied. If a single IDTF had multiple PIN(s) and the preferred PIN(s) had separate/specific procedures credentialed specific to that location, there could be inappropriate denials or payments.

FOR MORE INFORMATION

If you have received a development letter requesting your PIN, annotate on your response that you would like to select a preferred pin or contact the provider customer service area at 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

VISIT THE NPI MLN MATTERS WEB PAGE AND QUARTERLY JOURNAL ADVERTISEMENT

As Medicare May 23rd national provider identifier (NPI) implementation approaches, the Centers for Medicare & Medicaid Services (CMS) reminds providers to visit the NPI *MLN Matters* national provider education articles, courtesy of the **Medicare Learning Network**.

The national provider identifier (NPI) is a Health Insurance Portability and Accountability Act (HIPAA) Administrative Simplification standard. The NPI is a unique identification number for covered health care providers. Covered health care providers and all health plans and health care clearinghouses will use the NPIs in the administrative and financial transactions adopted under HIPAA.

The NPI is a 10-position, intelligence-free numeric identifier (10-digit number). This means that the numbers do not carry other information about health care providers, such as the state in which they live or their medical specialty.

The **Medicare Learning Network** has created many **MLN Matters** articles on the various aspects of Medicare's NPI implementation. A comprehensive list of the NPI articles is available on the CMS Web site at http://www.cms.hhs.gov/NationalProvidentStand/Downloads/MMArticles_NPI.pdf.

New MLN Quarterly Journal Ad—NPI *MLN Matters* Articles – This quarterly journal ad features the *MLN Matters* articles available regarding Medicare's implementation of the national provider identifier (NPI).

Each calendar quarter, the Medicare Learning Network creates a journal advertisement based on an initiative or new product of particular importance during that time frame. National, state and local associations are encouraged to use this journal ad in their publications and/or newsletters and Web sites, as appropriate.

The files for this quarter's ad, as well as future ads, may be found on the CMS Web site at <http://www.cms.hhs.gov/MLNGenInfo/>.

Source: CMS Provider Education Resource 200804-14

GENERAL INFORMATION

NEW MLN MATTERS ARTICLES NOW AVAILABLE ON DMEPOS COMPETITIVE BIDDING

Note: The Centers for Medicare & Medicaid Services (CMS) has issued *MLN Matters* special edition articles SE0806 and SE0807 since this provider education resource message was originated. The three in a series of articles, which are being published in the following pages, are intended to educate providers on the July 1, 2008, implementation of the Medicare for the durable medical equipment, prosthetics, orthotics, and supplies (DMEPOS) competitive bidding program.

Now Available! The *Medicare Learning Network (MLN Matters)* special edition article SE0805 titled – “Overview of New Medicare Competitive Bidding Program for Durable Medical Equipment, Prosthetics, Orthotics, and Supplies (DMEPOS) – The first in a series of articles on the implementation of this program.” is now posted at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0805.pdf> on the CMS Web site.

This is the first in a series of educational articles that will assist you in understanding this new DMEPOS program and will help you interact with your patients. The new program begins July 1, 2008, and additional educational materials will be made available to you as this date approaches.

This series of articles will be of particular interest to any provider that that may order, refer, or supply durable medical equipment to a Medicare beneficiary affected by the new Medicare DMEPOS competitive bidding program.

The Centers for Medicare & Medicaid Services (CMS) has developed a fact sheet that explains the program for Medicare beneficiaries. This fact sheet, entitled, “What You Should Know if You Need Medicare-covered Equipment or Supplies” is available at, <http://www.medicare.gov/Publications/Pubs/pdf/11307.pdf>.

You may want to provide this fact sheet to your Medicare patients.

Source: CMS Provider Education Resource 200804-01

OVERVIEW OF NEW MEDICARE COMPETITIVE BIDDING PROGRAM FOR DURABLE MEDICAL EQUIPMENT, PROSTHETICS, ORTHOTICS, AND SUPPLIES

The first in a series of articles on the implementation of this program.

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Any Medicare fee-for-service (FFS) provider that may be in a position of ordering, referring, or supplying durable medical equipment, prosthetics, orthotics, and supplies (DMEPOS) to a Medicare beneficiary may be affected by this program. This includes DMEPOS suppliers, physicians (including podiatric physicians), other treating practitioners (nurse practitioners, physician assistants, and clinical nurse specialists), physical and occupational therapists, and institutional providers (especially skilled nursing facilities and their social workers or care coordinators, hospitals and their discharge planners, home health agencies and pharmacists).

Note that those who refer or order DMEPOS for Medicare beneficiaries are being described as “referral agents” throughout this series.

PROVIDER ACTION NEEDED

STOP – Impact to You

Effective July 1, 2008, Medicare will begin implementation of a new program for purchasing DMEPOS for Medicare patients. For Medicare beneficiaries whose permanent residence is in one of the 10 metropolitan statistical areas (MSAs) affected by the first phase of this program, only contract suppliers, in most instances, will be eligible to provide competitive bid items and receive payment from Medicare. While new payment rules may not impact referral agents directly, they may impact your patients. Therefore, the Centers for Medicare & Medicaid Services (CMS) is providing this information to make you aware of the program so you can discuss it with your patients when necessary.

CAUTION – What You Need to Know

This program, initially, will affect patients obtaining DMEPOS in 10 competitive bidding areas (CBAs) that align with the 10 MSAs affected by the first phase of this program

and will include 10 product categories of DMEPOS. These areas and product categories will be identified later in this article. In general, if your patients reside in one of the CBAs, they must use a Medicare contract supplier for competitive bid items, unless they are willing to be responsible for full payment of these items. This means that some of your patients may have to change from a noncontract supplier to a contract supplier. Also, certain suppliers that rent DMEPOS that were not awarded contracts may be “grandfathered” under this program and may be able to continue to supply certain DMEPOS items/services should the beneficiary choose to continue to receive these items from a grandfathered supplier.

GO – What You Need to Do

It is important that all affected providers know this information. This program determines how much Medicare will pay for competitive bidding items and which suppliers are eligible to receive Medicare payments for these items. Be aware that the new program impacts payment amounts for certain DMEPOS items received by beneficiaries residing in one of the CBAs no matter where in the country they obtain their DMEPOS.

Be prepared for this program if you treat Medicare patients in one of the 10 areas affected by the first phase of this program, which are listed later in this article. Note that the program will expand to 70 additional MSAs in 2009.

BACKGROUND

Currently, Medicare payment for most DMEPOS is based on fee schedules. Recent amendments to the Social Security Act (the Act), however, will alter the process for determining payment amounts for certain DMEPOS items. Specifically, section 1847 of the Act mandates that competitive bidding payment amounts replace the current DMEPOS fee schedule payment amounts for selected items in selected areas. The intent is to improve the

Overview of New Medicare Competitive Bidding Program for DMEPOS, continued

effectiveness of the Medicare methodology for setting DMEPOS payment amounts, which will reduce beneficiary out-of-pocket expenses and save the Medicare program money while ensuring beneficiary access to quality items and services. The new method brings the payment amount for these items in line with that of a competitive market and reduces your patients' out-of-pocket expenses. The program also ensures the availability of a sufficient number of accredited suppliers for access to quality items and services. For more information on accreditation of DME suppliers, visit the CMS Web site http://www.cms.hhs.gov/CompetitiveAcqforDMEPOS/04_New_Quality_Standards.asp.

The law also provides for phasing in competitive bidding beginning in 10 of the largest MSAs. The program will be expanded into 70 additional MSAs in 2009 and the program will be expanded into additional areas after 2009. Areas that may be exempt from competitive acquisition of DMEPOS include rural areas and areas with low population density that are not competitive, unless there is a significant national market through mail order for a particular item or service. An area is chosen for the competitive bidding program based on several variables, including the size of its Medicare population and the amount of money spent on medical equipment and supplies in those areas.

Definitions

The following definitions are provided to explain several terms and their usage in this series of articles:

- **Contract Supplier** – An entity that is awarded a contract by CMS to furnish items under a competitive bidding program.
- **Noncontract Supplier** – A supplier that is not awarded a contract by CMS to furnish items included in a competitive bidding program.
- **Referral Agents** – This term applies to the range of physicians, practitioners or providers who prescribe DMEPOS (in essence, “order” or “refer”) for their patients.
- **Grandfathered Supplier** – A noncontract supplier that chooses to continue to furnish grandfathered items to a beneficiary in a CBA.
- **Grandfathered Item** – Any one of the items (as described in CFR section 414.220, 222, 226, and 229) for which payment is made on a rental basis prior to the implementation of a competitive bidding program and for which payment is made after implementation of a competitive bidding program to a grandfathered supplier that continues to furnish the items in accordance with section 414.408(j).
- **Single payment amount** – It means the allowed payment for an item furnished under a competitive bidding program.

The following table details how DMEPOS supplies may be acquired, given different scenarios:

If a beneficiary permanently lives in...	And travels to...	Type of supplier a beneficiary may go to...
A competitive bidding area	A competitive bidding area	A beneficiary must get competitively bid items from a contract supplier located in the competitive bidding area to which he/she traveled.
A competitive bidding area	An area not covered by the competitive bidding program	A beneficiary may get items from any Medicare-enrolled DME supplier, and Medicare will pay the supplier as if it were in the beneficiary's competitive bidding area.

For more information on single payment amounts, visit on the Internet <http://dmecompetitivebid.com/SPA>.

Initial Competitive Bidding Areas

Effective July 1, 2008, the competitive bidding program will be implemented in the following CBAs within these 10 MSAs:

- Charlotte-Gastonia-Concord, North Carolina and South Carolina
- Cincinnati-Middletown, Ohio, Kentucky, and Indiana
- Cleveland-Elyria-Mentor, Ohio
- Dallas-Fort Worth-Arlington, Texas
- Kansas City, Missouri and Kansas
- Miami-Fort Lauderdale-Miami Beach, Florida
- Orlando-Kissimmee, Florida
- Pittsburgh, Pennsylvania
- Riverside-San Bernardino-Ontario, California
- San Juan-Caguas-Guaynabo, Puerto Rico.

Product Categories

Effective July 1, 2008, the competitive bidding program will be implemented for the following product categories:

- Oxygen supplies and equipment
- Standard power wheelchairs, scooters, and related accessories
- Complex rehabilitative power wheelchairs and related accessories
- Mail-order diabetic supplies
- Enteral nutrients, equipment, and supplies
- Continuous positive airway pressure (CPAP), respiratory assist devices (RADs), and related supplies and accessories
- Hospital beds and related accessories
- Negative pressure wound therapy (NPWT) pumps and related supplies and accessories
- Walkers and related accessories
- Support surfaces (group 2 mattresses and overlays [**Miami MSAs only**]).

Traveling Beneficiaries

As previously mentioned, any beneficiary obtaining competitive bidding items in one of the CBAs is affected by the rules of the Medicare DMEPOS competitive bidding program. Beneficiaries who reside in a CBA and travels outside their CBAs may obtain competitive bid items and the supplier will be paid the single payment amount under the program.

In addition, beneficiaries who do not reside in CBAs and who travel to CBAs are also affected. If they require competitive bid items, they must obtain competitive bid items from a contract supplier for that CBA. In such instances, Medicare will pay that contract supplier the DMEPOS fee schedule amount.

Overview of New Medicare Competitive Bidding Program for DMEPOS, continued

If a beneficiary permanently lives in...	And travels to...	Type of supplier a beneficiary may go to...
An area not covered by the competitive bidding program	A competitive bidding area	A beneficiary must get the competitively bid item from a contract supplier in the competitive bidding area. If the beneficiary does not use a contract supplier, the noncontract supplier must ask him/her to sign an advance beneficiary notice. Medicare will not pay for competitively bid items furnished by noncontract suppliers.
An area not covered by the competitive bidding program	An area not covered by the competitive bidding program	A beneficiary may get items from any Medicare-enrolled DMEPOS supplier.

CMS is conducting extensive outreach to Medicare beneficiaries who reside in the CBAs and will be offering to help them identify contract suppliers.

If DMEPOS suppliers or referral agents are unsure whether a beneficiary resides in a CBA and is affected by this program effective July 1, they can make that determination by comparing the ZIP code of the patient's residence to the list of ZIP codes for the CBAs, which is available on the Internet at <http://dmecompetitivebid.com/Palmetto/Cbic.nsf/docsCat/DMEPOS%20Competitive%20Bidding%20Areas%20Zip%20Codes?opendocument>.

Payment

Payment for contract DMEPOS items will be the single payment amounts that were announced by CMS on March 20, 2008 (versus the current fee schedule determination of payment) for:

- Contract Suppliers
- Noncontract Suppliers that provide item to traveling beneficiaries.

ADDITIONAL INFORMATION

DMEPOS suppliers should note that previous articles have explained the program in more detail as it relates to

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DMEPOS suppliers. *MLN Matters* article SE0714, "Pre-Bidding Activities for the Medicare Durable Medical Equipment, Prosthetics, Orthotics, and Supplies (DMEPOS) Competitive Bidding Program," is available on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0714.pdf>.

Also, *MLN Matters* article MM5574, "Program Instructions Designating the Competitive Bidding Areas and Product Categories Included in the CY 2007 DMEPOS Competitive Bid Program," is available on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM5574.pdf>.

In addition, all providers may find more detailed information on the Internet at <http://www.dmecompetitivebid.com> and on the CMS Web site at <http://www.cms.hhs.gov/CompetitiveAcqforDMEPOS/>.

As this is the first in a series of *MLN Matters* articles on this issue, further articles will be released in the very near future.

MLN Matters Number: SE0805
 Related Change Request (CR) Number: N/A
 Related CR Release Date: N/A
 Related CR Transmittal Number: N/A
 Effective Date: N/A
 Implementation Date: N/A

GRANDFATHERING, REPAIR AND REPLACEMENT, MAIL ORDER DIABETIC SUPPLIES AND ADVANCED BENEFICIARY NOTICES

The second in a series of articles on the new DMEPOS competitive bidding program.

CMS has issued the following *MLN Matters* article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Any Medicare fee-for-service (FFS) provider supplying durable medical equipment, prosthetics, orthotics, and supplies (DMEPOS) to a Medicare beneficiary. This article also contains information of interest to those who order DMEPOS and to referral agents as defined in *MLN Matters* article SE0805.

PROVIDER ACTION NEEDED

The first article (SE0805) in this series on the DMEPOS competitive bidding program being instituted by the Centers for Medicare & Medicaid Services (CMS) presented an overview of how the program may affect your patients. There are also some key provisions of the program about which your patients may raise questions. While the competitive bidding program only affects ten areas of the

country as of July 1, 2008, it will expand to 70 additional geographic areas in 2009. Thus, it is important for you to be familiar with this program.

BACKGROUND

MLN Matters article SE0805, entitled "Overview of New Medicare Competitive Bidding Program for Durable Medical Equipment, Prosthetics, Orthotics, and Supplies (DMEPOS)," which is available on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0805.pdf>, summarizes information on competitive bidding that may impact your patients. Article SE0805 contains the list of competitive bidding areas for the first phase of competitive bidding as well as a list of the DMEPOS product categories that are included in the program's initial implementation.

Grandfathering, Repair and Replacement, Mail Order Diabetic Supplies and Advanced Beneficiary Notices, continued

In using this series of DMEPOS articles, it is important to remember that in most instances, beneficiaries maintaining a permanent residence in one of the competitive bidding areas (CBAs) must obtain competitive bidding items from a contract supplier. There are also program requirements that apply to beneficiaries who reside in CBAs but travel outside of those CBAs and to beneficiaries who do not live in CBAs but travel to them.

Grandfathered Suppliers

The Medicare DMEPOS competitive bidding program requires Medicare beneficiaries to obtain competitive bidding items from a contract supplier, unless an exception applies. Therefore, in some instances, your patient may be required to change from a non-contract supplier to a contract supplier. However, the program does allow for certain suppliers to be "grandfathered." Grandfathered suppliers are allowed to continue to provide certain rented DME items and services even though they are not contract suppliers.

Grandfathering only applies when the patient is renting DME or oxygen equipment at the time the competitive bidding program becomes effective and the rental period for the item began before the start of the competitive bidding program.

Beneficiaries who are receiving oxygen, oxygen equipment or rented DME at the time the competitive bidding program becomes effective may elect to continue to receive these items from a non-contract supplier, if the supplier is willing to continue furnishing these items. If a noncontract supplier chooses not to be "grandfathered" or if a beneficiary wants to change to a contract supplier, the noncontract supplier must pick up the rental equipment and oxygen equipment. Unless a beneficiary relocates outside of the CBA and the supplier service area, the supplier cannot discontinue services by picking up a medically necessary item prior to the end of a rental month for which the supplier was eligible to receive a rental payment, even if the last day of a rental month is after the start date of the program. If the date of the beginning of a monthly rental period is prior to the start of the competitive bidding program, the supplier must submit a claim for that month. Note that the grandfathering provision also applies to Medicare beneficiaries who transition from a Medicare Advantage plan to the fee-for-service program.

If the beneficiary stays with a "grandfathered" supplier, he or she may elect to change to a contract supplier at any time, and the contract supplier would be required to accept the beneficiary as a customer. For more details on the grandfathering provision, visit the CMS Web site <http://www.dmecompetitivebid.com>.

**Repair and Replacement of Beneficiary-Owned Items
Repair ONLY**

A beneficiary who owns a competitively bid item that needs to be repaired may have the repairs performed by either a contract supplier or by a non-contract supplier. In these cases, Medicare pays for reasonable and necessary labor not otherwise covered under a manufacturer's or supplier's warranty.

Repair and Replacement

If a part needs to be replaced in order to make the beneficiary-owned equipment serviceable, and the replacement part is also a competitively bid item for the CBA in which the beneficiary maintains a permanent residence, the part may be obtained from either a contract supplier or a non-contract supplier. In either case, Medicare pays the single payment amount provided under the competitive bidding program for the replacement part.

Replacement ONLY

Beneficiaries maintaining permanent residences in a CBA are required to obtain replacement of all items subject to competitive bidding from a contract supplier. This includes replacement of base equipment and replacement of parts or accessories for base equipment that are being replaced for reasons other than servicing of the base equipment.

Beneficiaries who are not permanent residents of a CBA but require a replacement of a competitively bid item while visiting a CBA, must obtain the replacement item from a contract supplier. The supplier will be paid the fee schedule amount for the state where the beneficiary is a permanent resident.

Mail Order Diabetic Supplies under the Program

Medicare beneficiaries who permanently reside in a CBA may purchase their diabetic testing supplies from:

- A mail order contract supplier for the area in which the beneficiary maintains a permanent residence; or
- A noncontract supplier in cases where the supplies are not furnished on a mail order basis.

The mail order contract period covers diabetic testing supplies furnished from **July 1, 2008 through March 31, 2010**. The term "mail order" refers to items ordered remotely (i.e., by phone, email, internet, or mail) and delivered to the beneficiary's residence by common carriers (e.g., U.S. Postal Service, Federal Express, United Parcel Service) and does not include items obtained by beneficiaries from local supplier storefronts.

Mail order contract suppliers will be reimbursed at the single payment amount for the CBA where the beneficiary maintains a permanent residence.

For diabetic supplies that are not furnished through mail order, suppliers will be paid the fee schedule amount.

Medicare payment will not be made to noncontract suppliers that furnish mail order diabetic testing supplies to Medicare beneficiaries residing in a CBA. A special modifier, **KL**, will be used on each claim to indicate that the item was furnished on a mail order basis.

Note: Suppliers that furnish diabetic testing supplies on a mail order basis and do not attach the mail order modifier could be subject to significant penalties under the False Claims Act.

Both the Medicare program and beneficiaries will save money each time a mail order contract supplier is used; **however, it is solely up to the beneficiaries to decide whether or not they wish to obtain their diabetic testing supplies on a mail order basis.**

All mail order contract suppliers are required to report the manufacturer or make and model number of products they furnish and must update this list on a quarterly basis. This information will be made available to the public once the contract suppliers have been announced and will be updated on a routine basis. Contract suppliers will be required to make available the same range of products to Medicare beneficiaries that they make available to non-Medicare customers.

Advance Beneficiary Notice Information

In general, if a noncontract supplier in a CBA furnishes a competitively bid item to any Medicare beneficiary regardless of whether that beneficiary maintains a permanent residence in the CBA or another area, and no applicable exceptions apply, Medicare will not make payment. In addition, the beneficiary is not liable for payment unless the noncontract supplier in a CBA obtains an advance beneficiary notice (ABN) signed by the beneficiary.

Grandfathering, Repair and Replacement, Mail Order Diabetic Supplies and Advanced Beneficiary Notices, continued

A signed ABN indicates that the beneficiary was informed in writing prior to receiving the item that there would be no Medicare coverage due to the supplier's contract status, and that the beneficiary understands that he/she will be liable for all costs that the noncontract supplier may charge the beneficiary for the item.

If a noncontract supplier furnishes a competitively bid item to a beneficiary and the beneficiary signs an ABN, the supplier must use modifier **GA** on their claim. If modifier the **GA** is not present on the claim, the supplier may not hold the beneficiary liable for the cost of the item.

ADDITIONAL INFORMATION

CMS contracted with the competitive bidding implementation contractor (CBIC) to administer the DMEPOS competitive bidding program. Downloadable **Patient Education Fact Sheets** may be found at: <http://www.dmecompetitivebid.com/palmetto/CBIC.nsf/docsCat/CBIC-Referral%20Providers-Patient%20Education%20Fact%20Sheets?open&cat=CBIC-Referral%20Providers-Patient%20Education%20Fact%20Sheets>.

If you have concerns, questions, or complaints about the quality of an item or the service that a patient received from a contract supplier, please call the competitive bidding program helpline at 1-877-577-5331.

For more information about the competitive bidding program, call 1-877-577-5331. TTY users call 1-877-486-2048. Stay tuned for additional articles in this series. You may also visit the CMS Web site <http://www.cms.hhs.gov/CompetitiveAcqforDMEPOS/> or <http://www.dmecompetitivebid.com/> on the Internet for more details.

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IMPORTANT EXCEPTIONS AND SPECIAL CIRCUMSTANCES UNDER THE DURABLE MEDICAL EQUIPMENT, PROSTHETICS, ORTHOTICS, AND SUPPLIES COMPETITIVE BIDDING PROGRAM

The third in a series of articles on the new DMEPOS competitive bidding program.

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

The following providers may be affected by this program:

- Physicians and other treating practitioners who are Medicare enrolled DMEPOS suppliers.
- Physicians and others who order or refer DMEPOS items or services for their patients.
- Skilled nursing facilities (SNFs) and nursing facilities (NFs).
- Physical therapists and occupational therapists in private practice who are Medicare enrolled DMEPOS suppliers.

Many Medicare fee-for-service (FFS) providers may be in a position of ordering, referring, or supplying DMEPOS to a Medicare beneficiary. This includes physicians (including podiatric physicians), other treating practitioners (nurse practitioners, physician assistants, and clinical nurse specialists), physical and occupational therapists, and institutional providers (especially skilled nursing facilities and their social workers or care coordinators, hospitals and their discharge planners, home health agencies and pharmacists).

PROVIDER ACTION NEEDED

Understand these special program rules that may affect you. This article is especially important if you are a Medicare enrolled DMEPOS supplier of items governed by the new program, even if you are not located in a competitive bidding area (CBA). It is important to understand that the program affects any beneficiaries who permanently reside in or travel to CBAs. Some program requirements apply to beneficiaries who reside in CBAs even if these beneficiaries travel outside their CBAs. Thus, it is important for you to be familiar with this program.

While the first phase of the competitive bidding program only affects ten CBAs in the country as of July 1, 2008, the second phase will expand to 70 additional

geographic areas in 2009. See *MLN Matters* article SE0805 for information about CBAs and items governed by this new program and for information about how the program applies to traveling beneficiaries.

BACKGROUND

MLN Matters article SE0805 that is entitled, "Overview of New Medicare Competitive Bidding Program for Durable Medical Equipment, Prosthetics, Orthotics, and Supplies (DMEPOS)," which is available on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0805.pdf>, summarizes information on competitive bidding that may impact your patients. *MLN Matters* article SE0805 contains the list of competitive bidding areas for the first phase of competitive bidding as well as a list of the DMEPOS product categories that are included in the program's initial implementation.

MLN Matters article SE0806 that is entitled, "Durable Medical Equipment, Prosthetics, Orthotics, and Supplies (DMEPOS) Competitive Bidding Program: Grandfathering, Repair and Replacement, Mail Order Diabetic Supplies and Advanced Beneficiary Notices (ABNs)," which is available on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0806.pdf>, provides an overview of the rules regarding grandfathered suppliers, repair and replacement of beneficiary-owned equipment, mail order diabetic supplies under the program, and ABNs.

In this, the third in a series of articles on the new DMEPOS competitive bidding program, we provide information on some special circumstances and exceptions of particular interest to physicians and other treating practitioners, SNFs and NFs, and physical and occupational therapists in independent practice.

Note: It is important to note that the Competitive Bidding Program does not affect your patients' choice of physician or treating practitioner.

Important Exceptions and Special Circumstances under the DMEPOS Competitive Bidding Program, continued

In using this series of DMEPOS articles, remember that in most instances, beneficiaries maintaining a permanent residence in one of the competitive bidding areas (CBAs) must obtain competitive bidding items from a contract supplier. There are also program requirements that apply to beneficiaries who reside in CBAs but travel outside of those CBAs and to beneficiaries who do not live in CBAs but travel to them.

Physicians and Other Treating Practitioners Who are Enrolled Medicare DMEPOS Suppliers

Medicare physicians and treating practitioners who have also enrolled as Medicare DMEPOS suppliers via the 855S enrollment form have the option to furnish certain types of competitively bid items to their own patients without submitting a bid or being awarded a competitive bid contract, provided the following requirements are met:

- For the first phase of the program being implemented July 1 2008, the item furnished must be a walker. In the future, the items will be limited to crutches, canes, walkers, folding manual wheelchairs, blood glucose monitors, and infusion pumps that are DME.
- The physician or treating practitioner DMEPOS supplier must furnish the items **to his or her own patients as part of his or her professional service.**
- The items must be billed to a DME MAC using the DMEPOS billing number that is assigned to the physician, the treating practitioner (if possible), or a group practice to which the physician or treating practitioner has reassigned the right to receive Medicare payment.

Where the furnished item is a bid item and the beneficiary resides in a CBA, the physician or treating practitioner will be paid the single payment amount established by this program for the item. This exception does not affect the applicability of the physician self-referral (Stark law) provisions in section 1877 of the Act. All provisions of the physician self-referral law remain fully in effect.

Physicians and Other Treating Practitioners Who Prescribe Specific Brand or Mode of Delivery to Avoid an Adverse Medical Outcome

A physician (including a podiatric physician) or treating practitioner may prescribe, in writing, a particular brand of DMEPOS bid item or mode of delivery for an item if he or she determines that the particular brand or mode of delivery would avoid an adverse medical outcome for the beneficiary. The physician or treating practitioner must document in the beneficiary's medical record the reason why the specific brand or mode of delivery is necessary to avoid an adverse medical outcome.

In these cases, the contract supplier under the competitive bidding program must:

- Furnish the particular brand or mode of delivery as prescribed by the physician or treating practitioner;
- Consult with the physician or treating practitioner to find another appropriate brand of item or mode of delivery for the beneficiary and obtain a revised written prescription from the physician or treating practitioner; or
- Assist the beneficiary in locating a contract supplier that can furnish the particular brand of item or mode of delivery prescribed by the physician or treating practitioner.

Any change in the prescription requires a revised written prescription. A contract supplier is prohibited from submitting a claim to Medicare if it furnishes an item different from that specified in the written prescription received from the beneficiary's physician or treating practitioner.

Skilled Nursing Facilities and Nursing Facilities Specialty Suppliers

The DMEPOS competitive bidding program applies to skilled nursing facilities (SNFs) and nursing facilities (NFs) to the extent that their residents receive competitively bid items under Medicare Part B. Unlike most suppliers, SNFs and NFs have the option to bid for, and be awarded, contracts to be "specialty suppliers" that **only furnish competitively bid items to their own residents.** SNFs and NFs that become specialty suppliers may not furnish competitively bid items and services to Medicare beneficiaries outside their facilities for purposes of Medicare payment. SNFs and NFs can also become regular contract suppliers that furnish competitively bid items to beneficiaries throughout a CBA.

If a SNF or NF is not a contract supplier (either a specialty contract supplier or a regular contract supplier), it must use a contract supplier for its CBA to furnish competitively bid items to its residents.

Physical Therapists and Occupational Therapists in Private Practice Who are Enrolled Medicare DMEPOS Suppliers

Physical therapists and occupational therapists in private practice who are enrolled DMEPOS suppliers may eventually have the option to furnish certain types of competitively bid items to their own patients and be paid the single payment amount for such items without being contract suppliers, provided the following requirements are met:

- The items are limited to off-the-shelf (OTS) orthotics; and
- The items must be furnished only to their own patients as part of the physical or occupational therapy service.

Note: OTS orthotics are not included in the first phase of competitive bidding, this exception is not relevant in the first phase of the DMEPOS competitive bidding program beginning July 1, 2008.

ADDITIONAL INFORMATION

If you have concerns, questions, or complaints about the quality of an item or the service that a patient received from a contract supplier please call the competitive bidding program helpline at 1-877-577-5331.

For more information about the competitive bidding program, call 1-877-577-5331. TTY users call 1-877-486-2048. Stay tuned for additional articles in this series. You may also visit <http://dmecompetitivebid.com/palmetto/cbic.nsf/DocsCat/Home> on the Internet and at <http://www.cms.hhs.gov/CompetitiveAcqforDMEPOS/> on the CMS Web site for more details.

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PRE-BIDDING ACTIVITIES FOR THE MEDICARE DURABLE MEDICAL EQUIPMENT, PROSTHETICS, ORTHOTICS, AND SUPPLIES COMPETITIVE BIDDING PROGRAM

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Suppliers of durable medical equipment (DME) that wish to participate in the upcoming Medicare durable medical equipment, prosthetics, orthotics, and supplies (DMEPOS) competitive bidding program.

PROVIDER ACTION NEEDED

In order to participate in the second round of the DMEPOS competitive bidding program, suppliers will be required to register in the Centers for Medicare & Medicaid Services (CMS) security system known as the individuals authorized access to CMS computer services (IACS). This includes suppliers that bid in the first round of competition last year and are interested in competing in the second round. Although the bidding window for the second round of competition may not be announced before the issue date of this article, CMS urges suppliers planning to bid in the 2008 bidding cycle to make sure their provider enrollment record is current. Specifically, suppliers should verify their supplier number(s) and authorized official(s) information associated with that supplier number(s) on file with the national supplier clearinghouse (NSC). The accuracy of this data is critical for successful bid registration.

BACKGROUND

In this year's bid cycle, suppliers who wish to bid will need to first register in IACS, before the bidding window opens. There will be three user roles available, which are described as follows:

Authorized Official (AO) – Each supplier's organization will be allowed one AO. The AO role can approve all other users associated with their organization who are requesting access to the bidding system. The AO will be able to input bid data, approve Form A and certify Form B in the bidding system.

Backup-Authorized Official (BAO) – Each supplier organization will be allowed to designate one or more BAOs. In this role, the BAO can approve the supplier's end user registration for access to the bidding system. Like the AO, the BAO can also input bid data, approve Form A and certify Form B in the bidding system.

End User – Each supplier organization will be allowed one or more end user(s). The end user can input bid data, but cannot approve Form A or certify Form B.

Save Time and Delay by Verifying NSC Information Prior to Registering to Bid

Only those AOs listed on the CMS-855S (Medicare Enrollment Application) as an AO may register in IACS to approve and certify as described above. As part of the CMS-855S, a supplier designates one or more AO(s). The AO is an appointed official to whom the organization has granted the legal authority to enroll it in the Medicare program and to commit the organization to fully abide by the statutes, regulations and program instructions of the Medicare program.

End users do not need to be listed on the CMS-855S. However, the AO or BAO will need to approve an end user's request for access to the bidding system.

Take Action Now

Be sure that the data you are submitting is current and in accordance with that submitted to the NSC. In particular, this concerns the AO's name, date of birth, social security

number (SSN), and mailing address. If any of these data elements have changed since your last submission to the NSC, then you should **promptly** complete a change of information on the CMS 855-S.

CMS urges that suppliers do it now. The NSC processing time to complete a change of information on the CMS-855S is approximately 45 days and all submissions are processed in the order in which they are received.

OVERVIEW OF AO IACS REGISTRATION PROCESS

For an AO, the verification of his/her last name, date of birth, and SSN must be validated against the data maintained by NSC. The NSC received this AO data when the supplier completed their most recent CMS-855S Medicare Enrollment Application. The AO's last name is listed in section 15 and the AO's date of birth and SSN in section 6A of the CMS-855S. If the data does not match, the registration will be rejected.

Following successful registration, as an added measure of security, the AO's User ID and password is then mailed in a separate correspondence to the mailing address listed in section 2A2 of the CMS-855S Medicare Enrollment Application.

The BAO goes through a similar process and an AO for the organization must approve a BAO's request for access before a User ID and password will be emailed to the BAO.

Do I Need a Backup-Authorized Official Role?

The establishment of a BAO is highly recommended to avoid any disruption in the bidding process. The AO's role is instrumental to bidding, as the AO's role must be active to avoid all other users of the organization from losing access to the bidding system. If the AO leaves the organization, the BAO role can be changed to an AO role by the competitive bidding implementation contractor (CBIC).

You will want to verify that the CMS-855S Medicare Enrollment Application for your organization has two or more AOs listed.

ADDITIONAL INFORMATION

For more information on the DMEPOS competitive bidding program, visit the CMS Web site <http://www.cms.hhs.gov/CompetitiveAcqforDMEPOS/>.

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AMBULANCE FEE SCHEDULE FACT SHEET NOW AVAILABLE

The *Ambulance Fee Schedule Fact Sheet*, which provides general information about the ambulance fee schedule, is now available in print format from the Centers for Medicare & Medicaid Services *Medicare Learning Network*. To place your order, visit <http://www.cms.hhs.gov/mlngeninfo/>, scroll down to "Related Links Inside CMS" and select "MLN Product Ordering Page."

Source: CMS Provider Education Resource 200803-14

ANNOUNCING THE RELEASE OF THE REVISED CMS-855 MEDICARE ENROLLMENT APPLICATIONS

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

All Medicare physicians, providers, and suppliers

BACKGROUND

The Centers for Medicare & Medicaid Services (CMS) issued revised CMS-855 Medicare enrollment applications in March 2008. With the exception of providers enrolling as a specialty hospital on the CMS-855A, Medicare contractors will continue to accept the 2006 version of the Medicare enrollment application through June 2008. **Providers and suppliers should begin to use the new Medicare enrollment applications immediately.** Initially, these applications will be available only from the CMS provider enrollment Web site. The link for that CMS Web site is listed in the *Additional Information* section of this article.

Over the last year, CMS has received numerous comments and suggestions regarding the proposed revisions to the Medicare enrollment applications. CMS reviewed the comments and adopted many of the suggested revisions. Also, CMS incorporated a number of enhancements and changes (see *Key Points* below) to clarify the enrollment process and to reduce the burden imposed on the provider and supplier communities.

KEY POINTS

This special edition outlines the significant revisions to the Medicare enrollment applications and they are as follows:

Application-Specific Changes for Physicians and Nonphysician Practitioners (CMS-855I)

- Removed the requirement in section 17 that providers attached their national provider identifier (NPI) notification that is received from the National Plan and Provider Enumeration System.

Application-Specific Changes for Clinics/Group Practices and Certain Other Suppliers (CMS-855B)

- Removed the supplier type "Voluntary Health/Charitable Agency" from section 2A.
- Clarified reporting timeframes throughout the CMS-855B.
- Added additional information about the NPI-legacy association and expanded the number of NPI – legacy combinations that a provider may enter in section 4A from one to five.
- Removed the requirement in section 17 that providers attach their NPI notification that is received from the National Plan and Provider Enumeration System.
- Required that an independent diagnostic testing facility (IDTF) submit copies of its comprehensive liability insurance policy in section 17.
- Added a list of the new IDTF standards found in 42 CFR 410.33(g) on a separate page in attachment 2.

- Added instructions that explain the IDTF liability insurance requirements in 42 CFR 410.33(g)(6) to attachment 2.

Application-Specific Changes for Institutional Providers (CMS-855A)

- Revised section 2A2 to include a specific box that specialty hospitals must check when completing the application. Instructions explaining the definition of a "specialty hospital" were also added to the form.
- Clarified the term "primary practice location" in the instructions in section 4. (The clarification did not change any data elements on the form.)
- Added additional information about the NPI-legacy association and expanded the number of NPI – legacy combinations that a provider may enter in section 4A from one to five.
- Removed the data element "Medicare Year-End Cost Report Date" from section 2.
- Removed the requirement in section 17 that providers attach their NPI notification that is received from the National Plan and Provider Enumeration System.

Application-Specific Changes for DMEPOS Suppliers (CMS-855S)

- Added supplier standards 22 – 25 to the list of DMEPOS supplier standards found on page 31.

ADDITIONAL INFORMATION

For additional information regarding the Medicare enrollment process, including the mailing address and telephone number for the carrier or FI serving your area, visit the CMS Web site <http://www.cms.hhs.gov/MedicareProviderSupEnroll>.

MLN Matters special edition article SE0612 contains helpful information about the Medicare enrollment process. You may review that article on the CMS Web site at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/SE0612.pdf>.

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CLINICAL LABORATORY FEE SCHEDULE - IMPLEMENTATION OF SECTION 113 MEDICARE, MEDICAID AND STATE CHILDREN'S HEALTH INSURANCE PROGRAM LEGISLATION

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Clinical laboratories billing Medicare contractors (carriers, fiscal intermediaries, or Part A/B Medicare administrative contractors [A/B MACs]) for services to Medicare beneficiaries.

PROVIDER ACTION NEEDED

This article is based on change request (CR) 5987 which alerts clinical laboratories that, effective for tests furnished on or after April 1, 2008, the Medicaid and State Children's Health Insurance Program (MMSCHIP) Extension Act of 2007 sets payment for code 83037 and 83037QW (Hemoglobin; glycosylated [A1c] by device) by crosswalking it to be the same as 83036 (glycosylated [A1c]). Make certain your billing staffs are aware of this change.

BACKGROUND

The MMSCHIP Extension Act of 2007 passed in December 2007 and included Section 113. Section 113 of the legislation set the price for any diagnostic test for HbA1C that is labeled by the Food and Drug Administration (FDA) for home use equal to the payment rate for a glycosylated hemoglobin test (identified as of October 1, 2007, by Healthcare Common Procedure Coding System [HCPCS] code 83036 [and any succeeding codes]). The legislation is effective for tests furnished on or after April 1, 2008.

- For calendar year (CY) 2006, the *Current Procedural Terminology (CPT)* established new code 83037 Hemoglobin; glycosylated (A1C) by device cleared by the FDA for home use. CPT code 83036, glycosylated (A1c), already existed and was priced at \$13.56 on the clinical laboratory fee schedule.
- For CY 2006, CMS determined that code 83037 should be paid via carrier gap filling.
- For CY 2007, CMS set the payment for code 83037 by crosswalking it to code 82985 (glycosylated protein).
- For tests furnished on or after April 1, 2008, the payment for 83037 or 83037QW will be the same as the payment on the clinical laboratory fee schedule for 83036.

Your Medicare contractor will adjust claims for services on or after April 1, 2008, processed prior to implementation of this change if you bring such claims to the contractor's attention.

ADDITIONAL INFORMATION

To see the official instruction (CR 5987) issued to your Medicare contractor visit <http://www.cms.hhs.gov/Transmittals/downloads/R331OTN.pdf> on the CMS Web site.

If you have questions, please contact your Medicare contractor at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5987

Related Change Request (CR) #: 5987

Related CR Release Date: April 11, 2008

Effective Date: April 1, 2008

Related CR Transmittal #: R331OTN

Implementation Date: May 12, 2008

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CLINICAL LABORATORY FEE SCHEDULE FACT SHEET

The Clinical Laboratory Fee Schedule Fact Sheet, which provides general information about the clinical laboratory fee schedule, coverage of clinical laboratory services, and how payment rates are set, is now available in downloadable format from the Centers for Medicare & Medicaid Services Medicare Learning Network at http://www.cms.hhs.gov/MLNProducts/downloads/clinical_lab_fee_schedule_fact_sheet.pdf.

Source: Provider Education Resources Listserv, Message 200804-06

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Join our eNews mailing list and receive urgent and other critical information issued by First Coast Service Options, Inc. (FCSO), your Medicare carrier. By signing up, you will receive automatic e-mail notification when new or updated information is posted to the provider education Web site. It's very easy to do. Simply go to our Web site <http://www.fcso.com>, select Medicare Providers, Connecticut or Florida, click on the "eNews" link located on the upper-right-hand corner of the page and follow the prompts.

CONTINUING EDUCATION CREDITS AVAILABLE FOR COCA CONFERENCE CALLS

Please remember that if you did not participate in the live COCA conference call, you can still get continuing education (CE) credit! Simply download the PowerPoint and follow along with the audio file, both of which are posted on the COCA Web site and then complete the on-line evaluation within a year <http://www2a.cdc.gov/TCEOnline/>.

Objectives

After this activity, the participants will be able to:

1. Describe safe injection and other basic infection control practices, and be able to recognize and correct unsafe practices
2. Understand the need for monitoring health care personnel practices in your facility relating to injection safety and basic infection control
3. Describe the potential consequences of syringe reuse and other unsafe practices
4. Locate related CDC infection control guidance and educational materials

Continuing Education guidelines require that the attendance of all who participate in COCA conference calls be properly documented. ALL CE credits (CME, CNE, CEU and CHES) for COCA conference calls are issued online through the CDC Training & Continuing Education Online system <http://www2a.cdc.gov/TCEOnline/>.

Those who participate in the COCA conference calls and who wish to receive CE credit and will complete the online evaluation by April 26, 2008, will use the course code EC1265. Those who wish to receive CE credit and will complete the online evaluation between April 27, 2008, and March 27, 2009, will use course code WD1265. CE certificates may be printed immediately upon completion of your online evaluation. A cumulative transcript of all CDC/ATSDR CE's obtained through the CDC Training & Continuing Education Online System will be maintained for each user.

Purpose

To enhance clinicians knowledge of emerging or re-emerging threats, their effects on human populations and medical evaluation and management of these threats.

CME: CDC is accredited by the Accreditation Council for Continuing Medical Education (ACCME) to provide continuing medical education for physicians. CDC designates this educational activity for a maximum of 1 Category 1 credit toward the AMA Physician's Recognition Award. Physicians should only claim credit commensurate with the extent of their participation in the activity.

CNE: This activity for 1.0 contact hours is provided by CDC, which is accredited as a provider of continuing education in nursing by the American Nurses Credentialing Center's Commission on Accreditations.

CEU: CDC has been reviewed and approved as an authorized provider by the International Association for Continuing Education and Training (IACET), 8405 Greensboro Drive, Suite 800, McLean, VA 22102. CDC has awarded 0.1 CEU to participants who successfully complete this program.

CHES: CDC is a designated provider of continuing education contact hours (CECH) in health education by the National Commission for Health Education Credentialing, Inc. This program is a designated event for the CHES to receive 1 Category I Contact Hour(s) in health education. CDC provider number GA0082.

Source: Provider Education Resources Listserv, Message 200803-15

CR 5550 CLARIFICATION—SIGNATURE REQUIREMENTS

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Physicians and other providers who bill Medicare contractors (carriers, fiscal intermediaries, regional home health intermediaries, Part A/B Medicare administrative contractors, including durable medical equipment Medicare administrative contractors) for care provided to Medicare beneficiaries in hospice.

WHAT YOU NEED TO KNOW

CR 5971, from which this article is taken, clarifies the instructions on signature requirements for the certification of terminal illness for hospice. It provides that Medicare contractors will accept a facsimile of an original written or electronic signature in documenting the certification of terminal illness for hospice.

Make sure that your billing staffs are aware that, to document the certification of terminal illness for hospice, a facsimile of an original written or electronic signature is acceptable.

BACKGROUND

CR 5971, from which this article is taken, clarifies the instructions in *Medicare Program Integrity Manual* chapter 3 (Verifying Potential Errors and Taking Corrective Actions), subsection 3.4.1.1B (Signature Requirements) that address the signature requirements for the certification of terminal illness for hospice, that were provided in CR 5550 (Various Medical Review Clarifications).

Subsection 3.4.1.1B of the manual notes that Medicare contractors require a legible identifier for services provided/ordered. It further requires that when this documentation is for medical review purposes, the only acceptable method of documenting the provider signature is by written or an electronic signature. Stamp signatures are not acceptable to sign an order or other medical record documentation for medical review purposes.

CR 5971 provides that there is an exception to this requirement.

It announces that a facsimile of an original written or electronic signature is acceptable for the certification of terminal illness for hospice. Please be sure to note however, that while a signature facsimile is acceptable in this instance; it and

CR 5550 Clarification—Signature Requirements, continued

hard copies of a physician's electronic signature must be present in the patient's medical record.

ADDITIONAL INFORMATION

You may find more information about the signature requirements for the certification of terminal illness for hospice by going to CR 5971, located at <http://www.cms.hhs.gov/Transmittals/downloads/R248PI.pdf> on the CMS Web site. You will find updated *Medicare Program Integrity Manual* chapter 3 (Verifying Potential Errors and Taking Corrective Actions), subsection 3.4.1.1B (Signature Requirements) as an attachment to this CR.

If you have any questions, please contact your Medicare contractor at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5971

Related Change Request (CR) #: 5971

Related CR Release Date: March 28, 2008

Effective Date: September 3, 2007

Related CR Transmittal #: R248PI

Implementation Date: April 28, 2008

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MEDICARE PHYSICIAN FEE SCHEDULE FACT SHEET NOW AVAILABLE

The revised *Medicare Physician Fee Schedule Fact Sheet* (January 2008), which provides general information about the Medicare physician fee schedule, is now available in print format from the Centers for Medicare & Medicaid Services *Medicare Learning Network*. To place your order, visit <http://www.cms.hhs.gov/mlngeninfo/>, scroll down to "Related Links Inside CMS" and select "MLN Product Ordering Page."

Source: CMS Provider Education Resource 200803-13

EXCEPTION TO 60-DAY LIMIT ON SUBSTITUTE PHYSICIAN BILLING ARRANGEMENTS FOR PHYSICIANS CALLED TO ACTIVE DUTY IN THE ARMED FORCES RESERVES

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Physician members of a reserve component of the armed forces who bill Medicare carriers or Medicare administrative contractors (A/B MAC) for services provided to Medicare beneficiaries.

Physicians called to active duty in the armed forces who wish to bill for substitute physician services during the physician's absence.

WHAT YOU NEED TO KNOW

Change request 5985, from which this article is taken, announces a six-month extension of the exception to the 60-day limit on substitute physician billing for physicians called to active duty in the armed forces. This means that a physician who is called to active duty may continue to bill for substitute physician services furnished from January 1, 2008 through June 30 2008, which may be beyond the 60-day limit.

Make sure that your billing staffs are aware of this change.

BACKGROUND

Section 1842(b)(6)(D)(iii) of the Social Security Act (the Act) and *Medicare Claims Processing Manual* chapter 1 (General Billing Requirements), sections 30.2.10 (Payment Under Reciprocal Billing Arrangements - Claims Submitted to Carriers) and 30.2.11 (Physician Payment Under Locum Tenens Arrangements - Claims Submitted to Carriers) state that when a physician is unavailable to provide services, a substitute physician's services (either on a reciprocal or locum tenens basis) are not to be provided for a period longer than 60 continuous days.

On August 3, 2007, public law 110-54 amended the Act to provide an exception to this 60-day limit for physicians who are ordered to active duty in the armed forces.

By striking "January 1, 2008" and inserting "July 1,

2008," Section 116 of the "Medicare, Medicaid, and State Children's Health Insurance Program (SCHIP) Extension Act of 2007" (signed on December 29, 2007) extended this exception for another six months.

CR 5985, from which this article is taken, updates these sections in *Medicare Claims Processing Manual* to reflect this change in the law.

Effective January 1, 2008, physicians called to active duty will be able to bill for substitute physician services furnished from January 1, 2008, through June 30 2008.

ADDITIONAL INFORMATION

You may find more information about the exception to the 60-day limit on substitute physician billing arrangements for physicians called to active duty in the armed forces reserves by going to CR 5985, located at <http://www.cms.hhs.gov/transmittals/downloads/R1486CP.pdf> on the Centers for Medicare & Medicaid Services (CMS) Web site. You will find the updated *Medicare Claims Processing Manual*, chapter 1 (General Billing Requirements), sections 30.2.10 and 30.2.11 as an attachment to that CR.

If you have any questions, please contact your carrier or A/B MAC at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5985

Related Change Request (CR) #: 5985

Related CR Release Date: April 4, 2008

Effective Date: January 1, 2008

Related CR Transmittal #: R1486CP

Implementation Date: May 5, 2008

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LABORATORY COMPETITIVE BIDDING DEMONSTRATION

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals. This information was previously published in the December 2007 Medicare B Update! pages 45-47.

Note: This article was changed on March 20, 2008, to reflect a change that was made to related change request 5389 on March 19, 2008. The CR was modified to delete a business requirement instructing carriers and A/B MACs deny claims with dates of service between April 1, 2007 and March 31, 2010 inclusive and with modifier "90" submitted by laboratories for demonstration-covered services provided to beneficiaries residing in the CBA. Since that requirement was deleted, language regarding that denial requirement was deleted from the article. The CR release date, transmittal number and Web address for accessing CR 5389 were also changed. All other information remains the same. However, it is important to note that a more current article, MM5772, is now available regarding this demonstration at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM5772.pdf> on the CMS Web site. Where there is disagreement between this article and related CR 5359 and MM5772 and related CR 5772, the information in CR 5772 is more current and takes precedence over CR 5359.

PROVIDER TYPES AFFECTED

Physicians and hospitals (TOB 14x only) who bill Medicare carriers, fiscal intermediaries (FIs), or Part A/B Medicare administrative contractors (A/B MACs) for clinical laboratory tests performed for Medicare Part B beneficiaries who live within the competitive bidding demonstration area (CBA) sites.

BACKGROUND

Section 302(b) of the Medicare Prescription Drug Improvement and Modernization Act of 2003 (MMA) requires the Centers for Medicare & Medicaid Services (CMS) to conduct a demonstration project on the application of competitive acquisition for payment of most clinical laboratory services that would otherwise be payable under the Medicare Part B fee schedule.

Under this statute, pap smears and colorectal cancer screening tests are excluded from this demonstration. Requirements under the Clinical Laboratory Improvement Amendments (CLIA), as mandated in section 353 of the Public Health Service Act, are applicable.

The payment basis determined for each CBA will be substituted for payment under the existing clinical laboratory fee schedule. Multiple winners are expected in each CBA.

Key Points

This article and change request (CR) 5359 provides instructions for the implementation of a laboratory competitive bidding demonstration. The requirements specified in this article and CR 5359 are in preparation for the implementation of the demonstration in the first CBA on April 1, 2007.

- The project will cover demonstration tests for all Medicare Part B beneficiaries who live in the demonstration sites, as determined by the ZIP code of the beneficiary's residence.
- Hospital inpatient testing is covered by Medicare Part A and is therefore exempt from the demonstration.
- Physician office laboratory (POL) testing and hospital outpatient testing are not included in the demonstration, except where the physician office or hospital laboratory functions as an independent laboratory performing testing for a beneficiary who is not a patient of the physician or hospital outpatient department.
- CMS will continue to pay POL patient and hospital outpatient laboratory services in accordance with the existing clinical laboratory fee schedule.

Required Bidders

Laboratory firms with \$100,000 or more in annual Medicare Part B (fee-for-service) payments as of calendar year (CY) 2005 for "demonstration tests" provided to beneficiaries residing in the CBAs (regardless of where the

laboratory firm is located) will be required to bid in the demonstration.

These laboratory firms will be referred to as "required bidders."

Passive Laboratories

Small laboratories or laboratory firms with less than \$100,000 in annual Medicare Part B (fee-for-service) payments for demonstration tests provided to beneficiaries residing in the CBAs will not be required to bid in the demonstration. These laboratories are considered "passive" laboratories." Passive laboratories will be paid the laboratory competitive bidding demonstration fee schedule for demonstration tests provided to beneficiaries residing in the CBA.

During the demonstration period, CMS will monitor the volume of services performed by passive laboratories to ensure that their annual payments under Medicare Part B for demonstration tests provided to beneficiaries residing in the demonstration sites do not exceed the annual ceiling of \$100,000.

Passive laboratory firms exceeding the annual ceiling of \$100,000 by \$25,000 or more will be:

- Terminated from the demonstration project.
- Will not be paid anything by Medicare for demonstration tests provided to beneficiaries residing in the CBAs (regardless of where the laboratory firm is located) for the duration of the demonstration.
- Laboratories or laboratory firms providing clinical laboratory services exclusively to beneficiaries with end-stage renal disease (ESRD) residing in the CBA will not be required to bid in the demonstration. These laboratories are considered "passive-ESRD" laboratories. Passive-ESRD laboratories will be paid the laboratory competitive bidding demonstration fee schedule for Part B demonstration tests provided to ESRD beneficiaries residing in the CBA. During the demonstration period (April 1, 2007 through March 31, 2010, inclusive), passive-ESRD laboratories that expand their business to provide clinical laboratory services to non-ESRD beneficiaries residing in the CBA will be terminated from the competitive bidding demonstration.

Winners

Both required and non-required bidders that bid and win will be paid the laboratory competitive bidding demonstration fee schedule for demonstration tests provided to beneficiaries residing in the CBAs (regardless of where the laboratory firm is located). These laboratories will be labeled "winners."

Non-Winners

Both required and non-required bidders that bid and do not win will not be paid anything by Medicare (neither under the Part B clinical laboratory fee schedule nor under the

Laboratory Competitive Bidding Demonstration, continued

competitively bid price) for demonstration tests provided to beneficiaries residing in the CBAs (regardless of where the laboratory firm is located) for the duration of the demonstration. These laboratories will be labeled "non-winners."

Similarly, required bidders that do not bid will not be paid anything by Medicare for demonstration tests provided to beneficiaries residing in the CBAs (regardless of where the laboratory firm is located) for the duration of the demonstration.

Non-winner laboratories that furnish a demonstration test to a Medicare beneficiary residing in the CBA during the demonstration have no appeal rights when Medicare payment for the test is denied. Moreover, non-winner laboratories may not charge the beneficiary for Part B laboratory services.

Demonstration-Covered Laboratory Tests

Only the laboratory that performs the test may bill for the service and only winning or passive laboratories are eligible to receive the laboratory competitive bidding demonstration fee schedule payment for services covered under the demonstration.

Although non-winner laboratories may not bill either Medicare or the beneficiary for any demonstration-covered services, such laboratories may refer such services to a winner laboratory or a passive laboratory.

For all other tests (i.e., those not covered under the demonstration or for tests for beneficiaries not residing in the service area), all laboratories will be paid according to the clinical laboratory fee schedule and in accordance with Medicare payment policies.

Demonstration Sites

There are two demonstration sites and each site runs for three years with a staggered start of one year. The demonstration uses metropolitan statistical areas (MSAs) to define the CBAs.

The residence status of beneficiaries will be determined by information in the Medicare system as of the date the claim is processed. The residence of the beneficiary receiving services must be in the same CBA as determined by review of a beneficiary's ZIP code of residence.

CMS will provide the contractors with a list of ZIP codes included in each MSA, which will be used to determine whether a beneficiary's residence is included in one of the CBAs.

The demonstration will set (competitively bid) fees in the demonstration areas for all tests paid under the Medicare Part B clinical laboratory fee schedule, with the exception of pap smears, colorectal cancer screening tests, and new tests added to the Medicare Part B clinical laboratory fee schedule during the course of the demonstration. Demonstration fees will be set for each service payable under the demonstration in each of the CBAs.

Only CLIA-certified laboratories will be allowed to participate in the demonstration.

Implementation

CR 5359 is being implemented in multiple phases. The requirements specified in this instruction are for the implementation of the demonstration in the first CBA (CBA1).

During the first quarter of 2007, CMS will provide Medicare carriers, FIs, and A/B MACs with a national ZIP code pricing file identifying the zip codes included in the first CBA. Also, in that same timeframe, CMS will provide to the carriers, FIs, and A/B MACs a list of the laboratories eligible to participate in the first CBA demonstration ("winners" and passive laboratories) and a list of those laboratories not selected to participate in CBA1.

For covered demonstration laboratory services in CBA1 with dates of service between April 1, 2007 and March 31, 2010, Medicare will pay the laboratory competitive bidding demonstration fee schedule amounts for laboratory services on that schedule. For services not on the

demonstration schedule, Medicare will pay based on the clinical laboratory fee schedule.

Claims submitted by non-winner laboratories for dates of service of April 1, 2007 through March 31, 2010, for Medicare beneficiaries in CBA1 will be denied using:

- Reason code 96 (non-covered charges)
- Remark code M114 (*This service was processed in accordance with rules and guidelines under the Medicare Clinical Laboratory Services Competitive Bidding Demonstration Project. If you would like more information regarding this project, you may contact your local contractor.*)
- Remark code N83 (No appeal rights. Administrative decision based on the provisions of a demonstration project).

Medicare will pay claims during the demonstration period submitted by non-demonstration laboratories for beneficiaries residing in the CBA who receive services outside of those areas (e.g., "snow birds") according to the laboratory competitive bidding demonstration.

Non-winning laboratories should know that advance beneficiary notices (ABNs) and notices of beneficiary exclusion from Medicare benefits (NEMBs) are not to be used to transfer liability to beneficiaries when services under the demonstration are obtained at non-winner laboratories.

Line items for demonstration services and for non-demonstration services may be submitted on the same claim.

A subsequent CR will be issued with requirements to implement the demonstration in the second CBA (CBA2).

Medicare contractors will be prepared to begin processing claims under the laboratory competitive bidding demonstration in the first CBA on April 1, 2007. The tentative start date for the demonstration in the second CBA is April 1, 2008.

Remember that required and non-required bidders that bid and lose will be paid nothing under the Part B clinical laboratory fee schedule and will have no appeal rights for demonstration tests provided to beneficiaries residing in the CBAs, regardless of the location of the laboratory itself.

IMPLEMENTATION

The implementation date for this instruction is April 2, 2007.

ADDITIONAL INFORMATION

The official instructions issued to your Medicare carrier, FI, or A/B MAC regarding this change may be found at <http://www.cms.hhs.gov/Transmittals/downloads/R57DEMO.pdf> on the CMS Web site.

If you have questions, please contact your Medicare carrier, FI, or A/B MAC at their toll-free number which may be found at: <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5359
Related Change Request (CR) #: 5359
Related CR Release Date: March 19, 2008
Effective Date: April 1, 2007
Related CR Transmittal #: R57DEMO
Implementation Date: April 2, 2007

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OUTPATIENT MAINTENANCE DIALYSIS—END-STAGE RENAL DISEASE FACT SHEET

The Outpatient Maintenance Dialysis – End-Stage Renal Disease Fact Sheet, which provides general information about outpatient maintenance dialysis for end-stage renal disease, the composite payment rate system, and separately billable items and services, is now available in downloadable format from the Centers for Medicare & Medicaid Services Medicare Learning Network at <http://www.cms.hhs.gov/MLNProducts/downloads/ESRDpaymfctst08-508.pdf>.

Visit the Medicare Learning Network – It's Free!

Source: Provider Education Resources Listserv, Message 200804-04

NOTICE OF INTEREST RATE FOR MEDICARE OVERPAYMENTS AND UNDERPAYMENTS

Medicare Regulation 42 CFR section 405.378 provides for the assessment of interest at the higher of the current value of funds rate (five percent for calendar year 2008) or the private consumer rate (PCR) as fixed by the Department of the Treasury.

The Department of the Treasury has notified the Department of Health & Human Services that the PCR has been changed to 11.375 percent, effective April 18, 2008. The PCR will remain in effect until a new rate change is published. The following table lists previous interest rates.

Period	Interest Rate
January 18, 2008 – April 17, 2008	12.125%
October 19, 2007 – January 17, 2007	12.5%
July 20, 2007 – October 18, 2007	12.625%
April 20, 2007 – July 19, 2007	12.375%
January 19, 2007 – April 19, 2007	12.5%
October 18, 2006 – January 18, 2007	12.375%

Source: CMS Pub. 100-06, Transmittal 138, CR 5752

USE OF PROFESSIONAL SOCIETY PRACTICE PARAMETERS IN PROPERLY PROVIDING ALLERGEN IMMUNOTHERAPY TO MEDICARE BENEFICIARIES

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

All physicians and providers who submit Medicare claims for providing allergen immunotherapy to Medicare beneficiaries.

PROVIDER ACTION NEEDED

This special edition (SE) article SE0812 is in response to recommendations made by the Department of Health & Human Services (DHHS), Office of Inspector General (OIG) in its 2006 "Allergen Immunotherapy for Medicare Beneficiaries" report. This article provides guidance and resources, developed by professional societies, for Medicare contractors and providers in determining the most appropriate provisions of allergen immunotherapy to Medicare beneficiaries.

BACKGROUND

The Centers for Medicare & Medicaid Services (CMS) does not have a national coverage determination (NCD) that identifies national coverage criteria for allergen immunotherapy services. Currently coverage of allergen immunotherapy services is at the local Medicare contractor's discretion. The February 2006 "Allergen Immunotherapy for Medicare Beneficiaries" report released by the DHHS, OIG identified areas of Medicare vulnerability due to insufficient available information regarding coverage, coding, and documentation requirements for physicians who provide allergen immunotherapy to Medicare beneficiaries.

KEY POINTS

- Because allergen immunotherapy may vary significantly based on geographical locations, CMS, with the support of the Joint Council of Allergy, Asthma, and Immunology (JCAAI), decided that an NCD was not the best mechanism for providing coverage to beneficiaries throughout the nation.

- CMS determined that contractor discretion for coverage of allergen immunotherapy is most appropriate.
- Without an NCD, Medicare contractors may develop their own local coverage determinations (LCDs).
- While these policies may vary between contractors, **CMS strongly encourages physicians who provide allergen immunotherapy to closely follow practice parameters agreed upon and endorsed by the professional societies that represent allergy, asthma, and immunology practitioners** like the JCAAI, as long as those parameters fall within the coverage criteria of applicable LCDs.
- Specific practice parameters are provided on the JCAAI Web site at <http://www.jcaai.org/> and are important for all practitioners and providers to study and understand.

ADDITIONAL INFORMATION

The CMS Web site provides a searchable database of LCDs, which is available on the CMS Web site at <http://www.cms.hhs.gov/mcd/search.asp>.

Providers may also wish to review the *Medicare NCD Manual*, which is available on the CMS Web site at <http://www.cms.hhs.gov/Manuals/IOM/list.asp>.

MLN Matters Number: SE0812
 Related Change Request (CR) Number: N/A
 Related CR Release Date: N/A
 Related CR Transmittal Number: N/A
 Effective Date: N/A
 Implementation Date: N/A

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WEB SITE FOR ADDITIONS AND DELETIONS OF ZIP CODES REQUIRING A PLUS FOUR ZIP CODE EXTENSION

CMS has issued the following MLN Matters article. Information for Medicare Fee-for-Service Health Care Professionals.

PROVIDER TYPES AFFECTED

Physicians, providers, and other health care providers submitting claims to Medicare fiscal intermediaries (FIs), carriers, Part A/B Medicare administrative contractors (A/B MACs) or regional home health intermediaries (RHHIs) for services paid under the Medicare physician fee schedule (MPFS) and for anesthesia services.

PROVIDER ACTION NEEDED

STOP – Impact to You

The ZIP code where services are rendered determines the payment locality for services paid under the MPFS and for anesthesia services. Certain ZIP codes fall into more than one payment locality and require a plus four ZIP code extension to ensure proper payment. (See the *MLN Matters* article at <http://www.cms.hhs.gov/MLNMattersArticles/downloads/MM5208.pdf> on the Centers for Medicare & Medicaid Services (CMS) Web site for further details regarding ZIP code reporting.)

CAUTION – What You Need to Know

The CMS will begin posting additions and deletions to the list of ZIP codes that require a plus four ZIP code extension on their Web site. A complete list of all ZIP codes requiring a plus four ZIP code extension will also be posted.

GO – What You Need to Do

Make certain your billing staffs are aware of these resources for checking plus four ZIP code extension requirements.

Key Points of CR 5970

- To access a file containing the quarterly additions and deletions to the list of ZIP Codes requiring a plus four extension refer to http://www.cms.hhs.gov/prospmedicarefeesvcvpmgen/01_overview.asp on the CMS Web site. The file is named “ZIP Code to Carrier Locality” and may be found in the *Downloads* section of this Web page.

- To access a file containing all ZIP Codes requiring a plus four extension, refer to http://www.cms.hhs.gov/prospmedicarefeesvcvpmgen/01_overview.asp on the CMS Web site. The file is named “ZIP Codes Requiring +4 Ext” and may be found in the *Downloads* section of this Web page.
- Upon release of a new quarterly update, the previous quarter’s additions and deletions are incorporated into the file name “ZIP Codes Requiring +4 Ext” file and are not included in the “ZIP Code Changes” file.

ADDITIONAL INFORMATION

To see the official instruction (CR 5970) issued to your Medicare FI, carrier, A/B MAC or RHHI refer to

<http://www.cms.hhs.gov/Transmittals/downloads/R1480CP.pdf> on the CMS Web site.

If you have questions, please contact your Medicare FI, carrier, A/B MAC or RHHI at their toll-free number, which may be found at <http://www.cms.hhs.gov/MLNProducts/downloads/CallCenterTollNumDirectory.zip> on the CMS Web site.

The toll-free number for First Coast Service Options, Inc. Medicare Part B Customer Service Center is 1-866-454-9007 (FL) or 1-888-760-6950 (CT).

MLN Matters Number: MM5970

Related Change Request (CR) #: 5970

Related CR Release Date: March 21, 2008

Effective Date: April 21, 2008

Related CR Transmittal #: R1480CP

Implementation Date: April 21, 2008

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THE CMS ONLINE MANUAL SYSTEM BROCHURE HAS BEEN UPDATED

The CMS Online Manual System: A Web-based Manual System for Medicare Contractors, Providers and State Agencies brochure has been updated and is now available to order print copies or to download as a PDF file. This brochure explains how to navigate the CMS Online Manual System. To view the PDF file, go to <http://www.cms.hhs.gov/MLNProducts/downloads/on-linebrochure.pdf>. Print copies may be ordered by visiting the *MLN* Product Ordering Page at http://cms.meridianksi.com/kc/main/kc_frame.asp?kc_ident=kc0001&loc=5 on the CMS Web site.

Source: Provider Education Resources Listserv, Message 200804-09

DELAY OF ENROLLMENT FOR EMPLOYED AUDIOLOGISTS BILLING UNDER A PHYSICIAN

On February 29, 2008, the Centers for Medicare & Medicaid Services (CMS) issued change request 5717 (CR) titled “Update to Audiology Policies.” Transmittal 1470 of that change request provided clarifications to the *Medicare Claims Processing Manual* (Pub. 100-02). In Pub 100-02, chapter 12, section 30.3, the manual instructions state, “. . . the audiologist’s NPI is required on all claims for services furnished by audiologists.” Use of the NPI in the primary identifier field on a claim requires Medicare enrollment.

Note that CMS is instructing contractors to, prior to October 1, 2008, continue to process claims without the NPI of the audiologist. All other instructions in CR 5717 remain unchanged.

CMS will require the use of the NPI on claims for diagnostic test services furnished by audiologists on or after October 1, 2008. Audiologists are encouraged to obtain an NPI and enroll as soon as possible.

Note: This delay only applies to audiologists who are employed by and billing under a physician’s NPI. Independent audiologist are not entitled to this delay and therefore subject to the mandatory NPI requirements.

Source: CMS Joint Signature Memorandum 08252, April 3, 2008

MEDICARE LEARNING NETWORK BOOKMARK NOW AVAILABLE

The MLN bookmark lists: the topics covered by the educational products and services of the MLN, the various product types available to the learner, as well as the Web address for the MLN. This product is appropriate for distribution at health care professional conferences, provider outreach and education activities and other appropriate types of provider/supplier events.

The MLN bookmark is available for download on the CMS Web site at <http://www.cms.hhs.gov/MLNProducts/downloads/MLNBookmrk-006960.pdf>.

You can also order hard copies of the bookmark through the MLN Product Ordering page on the Web at http://cms.meridianksi.com/kc/main/kc_frame.asp?kc_ident=kc0001&loc=5.

A version of the MLN bookmark is also available for distribution to Indian health care professionals. To view this bookmark, go to <http://www.cms.hhs.gov/MLNProducts/downloads/MLN-AIANBookmrk006954.pdf> or to order hard copies go to http://cms.meridianksi.com/kc/main/kc_frame.asp?kc_ident=kc0001&loc=5 on the Web.

Source: CMS Provider Education Resource 200804-14

ARE YOU SUBMITTING CLAIMS USING MODIFIER 52?

If you are submitting claims to Medicare using modifier 52, you should pay special attention to this article. First Coast Service Options Inc. (FCSO) has identified that a significantly high number of providers are submitting modifier 52 for the incorrect reason. We have verified that many providers are using the modifier to indicate a reduction in the charge amount to patients. There were many different reasons given for reducing the charge amount to the patient, but this is not the correct use of the modifier.

The *Current Procedural Terminology (CPT)* definition of modifier 52 is “Reduced services”: under certain circumstances a service or procedure is partially reduced or eliminated at the physician’s discretion. Under these circumstances the services can be identified by its usual procedure number and the addition of modifier 52, signifying the service is reduced. This provides a means of reporting reduced services without disturbing the identification of the basic service.

Modifier 52 is used to indicate that a service or procedure, which is described by a *CPT/HCPCS* code, was not completed or rendered in its entirety. The Medicare fee schedules reimburse for these services when they have been completed according to the descriptor of the services. Therefore, by using modifier 52 you are advising Medicare that you did not complete the procedure/service as described by the *CPT/HCPCS* code. You are advising Medicare to consider whether or not your reimbursement should be reduced as a result of not performing or completing the procedure. Although you may have reduced your charge amount because of the incomplete procedure/service, the modifier is not used to indicate a reduction in your billed amount.

FCSO will begin requesting supporting records and documentation on all claims submitted with modifier 52. Records will be requested to determine the medical necessity of services billed, and to determine if a reduction in the Medicare allowance is appropriate. Documentation should clearly indicate why the modifier is being used. Records must identify what portion of the service was completed and/or not completed. In order to avoid unnecessary development and records request, please ensure that you are correctly using the modifier on all claims.

Most commonly found examples of incorrect usage:

- If a *CPT/HCPCS* code descriptor indicates “unilateral or bilateral” and the service is performed unilaterally, it is not appropriate to bill using modifier 52. The key word is “or”, meaning the same code applies in both cases.
- If a *CPT/HCPCS* diagnostic code indicates “with contrast material” and a patient receives the test without contrast, it is not appropriate to bill using modifier 52. The code indicating without contrast is to be used.
- If you are attaching modifier 52 to *CPT* code 97014 (electrical stimulation) to indicate a reduction in minutes, this is not appropriate, electrical stimulation is not a timed code.

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OVERPAYMENT REDETERMINATION REQUEST FORM—INSTRUCTIONS

The “Request for Overpayment Redetermination of a Medicare Part B Claim” form on the following pages simplifies and standardizes filing requirements for redeterminations. The overpayment redetermination form:

- allows the provider of services to clearly specify the reason(s) he or she disagrees with the overpayment determination (section 6).
- provides space for a comprehensive and detailed explanation of any additional information that should be considered when the overpayment is reviewed (section 8).

Completion of these two sections is critical to correct processing of your overpayment redetermination request. Using this form will make handling requests for overpayment redeterminations easier and more efficient for providers’ offices.

If all related information (dates of service, procedure codes, etc.) is filled in on the form as requested, copying and mailing of additional medical records may be significantly reduced (the current requirements for documentation for certain redetermination types have not changed). Follow the instructions on the reverse side of the form and submit your request to the address indicated in section 1.

Overpayment Redetermination Request for Medicare Part B Claim

NOTICE - Anyone who misrepresents or falsifies essential information requested by this form may upon conviction be subject to fine and imprisonment under Federal Law.

Print legibly and complete all information.

① **Carrier's Name and Address** Medicare Part B Overpayment Review
P. O. Box 45248
Jacksonville, Florida 32232-5248

② Name of Patient	③ Medicare Health Insurance Claim Number (9 digits followed by an alpha/numeric suffix)
--------------------------	---

④ **I do not agree with the determination you made on ICN** _____

⑤ **Accounts receivable number** _____

⑥ **The reason(s) I disagree with the determination is/are:** (Please check those that apply)

Service/Claim underpaid/reduced Service/Claim overpaid

Duplicate Service(s) overutilized and/or not medically necessary

OTHER: (Please be specific) _____

⑦ **For services in question, please provide:**

Date(s) of Service:	Quantity Billed:	Modifier:	Procedure Code:
_____	_____	_____	_____
_____	_____	_____	_____
_____	_____	_____	_____
_____	_____	_____	_____

⑧ **Additional information to consider including specific diagnosis, illness and/or condition:**

⑨ **Attachments to consider:** (Please check all that apply)

Medical Records Copy of Claim

Ambulance Run Sheet Certificate of Medical Necessity

Office Records/Progress Notes

OTHER: (Please be specific) _____

⑩ Signature of Claimant or Representative:	Telephone Number:
(Print) _____	_____
(Written) _____	_____

**Instructions for Completing the
Overpayment Redetermination Request Form**

Block ① - Carrier's Name and Address

All requests for redetermination of Medicare Part B overpayments should be mailed to address indicated in this block.

Block ② - Name of Patient

Enter the patient's last name, first name, and middle initial, if any, as shown on the patient's Medicare card.

Block ③ - Medicare Health Insurance Claim Number

Enter the patient's Medicare Health Insurance Claim Number (HICN) as shown on the patient's Medicare card.

Block ④ - I do not agree with the determination you made on ICN:

Indicate the 13 digit Internal Control Number (ICN) assigned to the claim submitted for reimbursement. This number can be found on the Provider Claim Summary (PCS), Provider Remittance Notice (PRN) or Medicare Summary Notice (MSN).

Block ⑤ - Indicate the 13-digit accounts receivable number.

This number can be found on the header of the Overpayment request letter or at the top of the Health Data Insight Request form.

Block ⑥ - The reason(s) I disagree with the determination is/are:

Check appropriate item(s) why you disagree with the decision made on the claim being submitted for redetermination. If OTHER is checked, please provide specific information.

Block ⑦ - For services in question, please provide:

Please indicate on each line the date(s) of service and procedure code that you are requesting be reviewed. A consecutive date range may be used per line; however, it should be for only one procedure code.

➤➤Example of this would be: 01/01/2007 - 01/12/2007 99232 <<

Block ⑧ - Additional information to consider including specific diagnosis, illness and/or condition:

Provide any additional information that was not originally provided when the claim was submitted for processing.

Block ⑨ - Attachments to consider:

Check which attachments are being included with this form for consideration with the review of the claim being appealed.

Block ⑩ - Signature of Claimant or Representative:

Signature of claimant or his representative and telephone number.

Reminder 📄 You may provide a cover letter or attachments for multiple review cases.

LOCAL COVERAGE DETERMINATIONS

UNLESS OTHERWISE INDICATED, ARTICLES APPLY TO BOTH CONNECTICUT AND FLORIDA

This section of the *Medicare B Update!* features summaries of new and revised local coverage determinations (LCDs) developed as a result of either local medical review or comprehensive data analysis initiatives. These initiatives are designed to ensure the appropriateness of medical care and that the carrier's LCDs and review guidelines are consistent with accepted standards of medical practice. In accordance with publication requirements specified by the Centers for Medicare & Medicaid Services (CMS), carriers no longer include full-text local coverage determinations (LCDs) to providers in the *Update!* Summaries of revised and new LCDs are provided instead. Providers may obtain full-text LCDs on our provider education Web sites, <http://www.fcso.com>. Final LCDs, draft LCDs available for comment, LCD statuses, and LCD comment/response summaries may be printed from the Part B Medical Policy section.

EFFECTIVE AND NOTICE DATES
Effective dates are provided in each LCD, and are based on the date of service (unless otherwise noted in the LCD). Medicare contractors are required to offer a 45-day notice period for LCDs; the date the LCD is posted to the Web site is considered the notice date.

ELECTRONIC NOTIFICATION
To receive quick, automatic notification when new LCDs are posted to the Web site, subscribe to our FCSSO eNews mailing list. It's very easy to do; go to our Web site <http://www.fcso.com>, select Medicare Providers, Connecticut or Florida,, click on the "eNews" link located on the upper-right-hand corner of the page and follow the prompts.

MORE INFORMATION
For more information, or, if you do not have Internet access, to obtain a hardcopy of a specific LCD, contact Medical Policy at:

Medical Policy and Procedures
PO Box 2078
Jacksonville, FL 32231-0048

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ADVANCE BENEFICIARY NOTICE

Modifier GZ must be used when physicians, practitioners, or suppliers want to indicate that they expect that Medicare will deny an item or service as not reasonable and necessary and they **have not had** an advance beneficiary notification (ABN) signed by the beneficiary.

Modifier GA must be used when physicians, practitioners, or suppliers want to indicate that they expect that Medicare will deny a service as not reasonable and necessary and they **do have** on file an ABN signed by the beneficiary.

All claims not meeting medical necessity of a local coverage determination must append the billed service with modifier GA or GZ.

NEW LCDs

51784: ANORECTAL MANOMETRY AND EMG OF THE URINARY AND ANAL SPHINCTERS—NEW LCD

This new local coverage determination (LCD) for anorectal manometry and EMG of the urinary and anal sphincters was developed to outline the indications and limitations for these diagnostic tests. Medical review of claims showed that providers were performing these diagnostic tests on a frequent basis for patients who were receiving “pelvic floor therapy” or “pelvic floor rehabilitation” for services that are more appropriately described by CPT code 90911 (*Biofeedback training, perineal muscles, anorectal or urethral sphincter, including EMG and/or manometry*) or for services that fall under the physical therapy benefit. This new LCD applies to CPT codes 51784 (*Electromyography studies (EMG) of anal or urethral sphincter, other than needle, any technique*), 51785 (*Needle electromyography studies (EMG) of anal or urethral sphincter, any technique*) and 91122 (*Anorectal manometry*) and includes indications, limitations, utilization guidelines and documentation requirements. The LCD also outlines the ICD-9-CM codes that support medical necessity for CPT codes 51784/51785 and for CPT code 91122.

EMG of the anal or urinary sphincters (CPT code 51784 or 51785) are diagnostic tests that measure muscle activity and are used to assist in evaluating a diagnosis of fecal or urinary incontinence, dysfunctional elimination of the bowel and bladder and neurogenic bladder dysfunction leading to functional abnormalities of the muscular sphincter. Anorectal manometry (CPT code 91122) is a diagnostic test that measures the anal sphincter pressures and provides assessment of rectal sensation, rectoanal reflexes and rectal compliance. It is used to help evaluate a diagnosis of fecal incontinence or anorectal elimination dysfunction.

EFFECTIVE DATE

This new LCD is effective for services rendered on or after June 30, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

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87181: SUSCEPTIBILITY STUDIES—NEW LCD

Susceptibility testing is performed by growing the pure bacterial isolate in the presence of varying concentrations of several antimicrobials and then examining the amount of growth to determine which antimicrobials at which concentrations inhibit the growth of the bacteria. If there is more than one pathogen, the laboratory will report results for each one. The test results of antimicrobial susceptibility testing should be combined with clinical information and experience when selecting the most appropriate antibiotic for the patient.

The new local coverage determination (LCD) for susceptibility studies was developed to define indications and limitations of coverage and/or medical necessity, documentation requirements, ICD-9-CM codes that support medical necessity, and coding guidelines for CPT codes 87181 through 87190. The diagnoses listed under the “ICD-9 Codes that Support Medical Necessity” section of the LCD represent payable diagnosis codes added by First Coast Service Options, Inc. (FCSO). In addition, diagnosis codes listed in the “Covered Code List” of the “CMS Medicare National Coverage Determinations (NCD)”, section 190.12 – Urine Culture, Bacterial” that include CPT codes 87184 and 87186 are listed in the “Coding Guidelines” attachment of the LCD. These ICD-9-CM codes are also payable for CPT codes 87181 – 87190 when indicated.

EFFECTIVE DATE

This new LCD is effective for services rendered on or after June 30, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

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SIGN UP TO OUR eNEWS ELECTRONIC MAILING LIST

Join our eNews mailing list and receive urgent and other critical information issued by First Coast Service Options, Inc. (FCSO), your Medicare carrier. By signing up, you will receive automatic e-mail notification when new or updated information is posted to the provider education Web site. It's very easy to do. Simply go to our Web site <http://www.fcso.com>, select Medicare Providers, Connecticut or Florida, click on the “eNews” link located on the upper-right-hand corner of the page and follow the prompts.

REVISION TO THE LCDs

BOTULINUM TOXINS: BOTULINUM TOXINS—REVISION TO THE LCD

The local coverage determination (LCD) for botulinum toxins was last revised on December 22, 2006. Since that time, the “Other Comments” section of the ‘Coding Guidelines’ attachment for this LCD has been revised to add the following language:

Botulinum toxins are best described as a family of toxins produced by the anaerobic organism *Clostridia Botulinum*. There are seven distinct serotypes designated as type A, B, C-1, D, E, F and G. Each serotype is produced by a specific strain of *Clostridium Botulinum*. In this country two preparations of botulinum are available and they are manufactured from two different strains of bacteria: Type A (BOTOX®) is made from the Hall strain and has been used for more than two decades and, Type B (Myobloc™) is made from the bean strain and was approved by the FDA in December 2000.

Whether a botulinum toxin is produced from the same or a different serotype producing strain, they undergo different manufacturing processes, which yield differences in the size and weight of the molecules. Because of this, botulinum toxin A and B, as well as other type A products available internationally, are not interchangeable. They are chemically, pharmacologically and clinically distinct.

Please note the FDA labeling in each product’s package insert states: “Units of biological activity cannot be converted into units of any other botulinum toxin or any toxin assessed with any other specific assay method.”

A revision has also been made to separate the Florida and Connecticut LCDs into individual distinct LCDs.

EFFECTIVE DATE

This LCD revision is effective for services rendered on or after April 22, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

IDTF: INDEPENDENT DIAGNOSTIC TESTING FACILITY—CODING GUIDELINE REVISION

The local coverage determination (LCD) for independent diagnostic testing facility (IDTF) was effective for services rendered on or after February 29, 2008. Since that time, the “Credentialing Matrix” in the “Coding Guidelines” attachment has been revised. The “Technician Qualification Requirements” for CPT codes 95805 – 95811 have been revised to delete the requirement of “State License: Respiratory Therapist” and the “Technician Qualification Requirements” have been revised for all magnetic resonance imaging procedures listed in the “Credentialing Matrix” to delete the requirement of “State License: General Radiographer”.

EFFECTIVE DATE

This revision to the “Coding Guidelines” attachment is effective for services rendered on or after April 22, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

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J1561: INTRAVENOUS IMMUNE GLOBULIN—REVISION TO THE LCD

The local coverage determination (LCD) for intravenous immune globulin (IVIG) was last revised January 1, 2008. Since that time, the Centers for Medicare & Medicaid Services (CMS) issued change request 5980 (April Update to the 2008 Medicare Physician Fee Schedule Database [MPFSDB]), transmittal 1482, dated March 21, 2008, which included the addition of HCPCS code Q4097 (Injection, immune globulin (Privigen), intravenous, non-lyophilized (e.g., liquid), 500 mg). Therefore, the LCD has been revised to include the addition of HCPCS code Q4097 to the “CPT/HCPCS Codes” section of the LCD.

EFFECTIVE DATE

This revision is effective for claims processed on or after April 7, 2008 for services rendered on or after April 1, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

SIGN UP TO OUR eNEWS ELECTRONIC MAILING LIST

Join our **eNews** mailing list and receive urgent and other critical information issued by First Coast Service Options, Inc. (FCSO), your Medicare carrier. By signing up, you will receive automatic e-mail notification when new or updated information is posted to the provider education Web site. It’s very easy to do. Simply go to our Web site <http://www.fcso.com>, select Medicare Providers, Connecticut or Florida, click on the “eNews” link located on the upper-right-hand corner of the page and follow the prompts.

J9310: RITUXIMAB (RITUXAN®)—REVISION TO THE LCD

The local coverage determination (LCD) for rituximab (Rituxan®) was last updated on January 7, 2008. Since that time, a revision was made to update language for approved indications based on the Food and Drug Administration (FDA) drug label, for rituximab – J9310.

Language was updated for Non-Hodgkin's lymphoma, and a new indication concerning progression of structural damage was added to rheumatoid arthritis (RA) under the "Indications and Limitations of Coverage and/or Medical Necessity" section of the LCD. The indication for RA now states Rituximab (Rituxan®) is FDA approved "In combination with methotrexate to reduce signs and symptoms and to slow the progression of structural damage in adult patients with moderately-to severely-active rheumatoid arthritis who have had an inadequate response to one or more TNF antagonist therapies." In addition, the "Sources of Information and Basis for Decision" section of the LCD was updated, and drug information was added under the "Utilization Guidelines" section of the LCD.

EFFECTIVE DATE

This revision to the LCD is effective for services rendered on or after January 25, 2008. The full-text of this LCD is available through our provider education Web site <http://www.fcso.com>.

NCSVCS: THE LIST OF MEDICARE NONCOVERED SERVICES—REVISION TO THE LCD

The local coverage determination (LCD) for the list of Medicare noncovered services (NCSVCS) was last revised February 29, 2008. Since that time, the Centers for Medicare & Medicaid Services (CMS) issued change request 5980 (April Update to the 2008 Medicare Physician Fee Schedule Database [MPFSDB]), transmittal 1482, dated March 21, 2008, which included changing the MPFSDB status indicator for HCPCS code J7307 (Etonogestrel [contraceptive] implant system, including implant and supplies) to an "I" (Not valid for Medicare purposes). Therefore, HCPCS code J7307 was deleted from the "Local Noncoverage Decisions-Devices" section of the LCD.

EFFECTIVE DATE

This revision is effective for claims processed on or after April 7, 2008 for services rendered on or after January 1, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

NCSVCS: THE LIST OF MEDICARE NONCOVERED SERVICES—REVISION TO THE LCD

The local coverage determination (LCD) for the list of Medicare noncovered services was last revised on April 7, 2008 for services rendered on or after January 1, 2008. Since that time, the LCD has been revised to add the following CPT codes to the list of procedures under the "CPT/HCPCS Codes", "Local Noncoverage Decisions, Procedures" section of the LCD, as these procedures are considered experimental and investigational:

0042T Cerebral perfusion analysis using computed tomography with contrast administration, including post-processing of parametric maps with determination of cerebral blood flow, cerebral blood volume, and mean transit time

0088T Submucosal radiofrequency tissue volume reduction of tongue base, one or more sites, per session (ie, for treatment of obstructive sleep apnea syndrome)

0170T Repair of anorectal fistula with plug (eg, porcine small intestine submucosa [SIS])

0181T Corneal hysteresis determination, by air impulse stimulation, bilateral, with interpretation and report

64999 Percutaneous neuromodulation using a percutaneous electrode array (PEA) (eg, BioWave)

CPT code 22899 (*Unlisted procedure, spine*) is included in the Surgery/Musculoskeletal System section of the American Medical Association (AMA) CPT 2007 Professional Edition and refers to an unlisted surgical procedure. However, this service is non-surgical and should be reported using CPT code 97799 (*Unlisted physical medicine/rehabilitation service or procedure*) to represent Vertebral Axial Decompression (VAX-D)/Intervertebral Differential Dynamics (IDD). Therefore, CPT code 22899 (*Vertebral Axial Decompression (VAX-D)*) was replaced with CPT code 97799 (*Vertebral Axial Decompression/Intervertebral Differential Dynamics [IDD]*) under the 'CPT/HCPCS Codes', 'National Noncoverage Decisions, Procedures' section of the LCD, as this procedure is considered experimental and investigational.

This service is performed for symptomatic relief of pain associated with lumbar disk problems. This treatment combines pelvic and/or cervical traction connected to a special table that permits the traction application. However, there is insufficient scientific data to support the benefits of this technique. Therefore, VAX-D is not covered by Medicare (CMS Manual System, Pub 100-03, *Medicare National Coverage Determinations (NCD) Manual*, chapter 1, part 2, section 160.16, Vertebral Axial Decompression [VAX-D]).

Because this NCD specifically refers to a treatment which "combines pelvic and/or cervical traction connected to a special table that permits the traction application", any similar device would fall under this category of a noncovered benefit. Therapeutic Tables (e.g., VAX-D, Decompression Reduction Stabilization (DRS) System, Accu-Spina System, DRX-3000, DRX9000, SpineMED Decompression Table, and Lordex Traction Unit) are used for this service and are motorized traction devices used to stretch the lower back.

EFFECTIVE DATE

This LCD revision is effective for services rendered on or after June 30, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

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VISCO: VISCOUSUPPLEMENTATION THERAPY FOR KNEE—REVISION TO THE LCD

The local coverage determination (LCD) for viscosupplementation therapy for knee was last revised January 1, 2008. Since that time, it has come to the attention of First Coast Service Options, Inc. (FCSO) that the LCD required clarification regarding the number of allowable courses of treatment. Therefore, the statement located at the third bullet under the "Limitations" section of the LCD has been revised to indicate the following:

- If the first course of treatment produces relief, a subsequent course of treatment may be reasonable if symptoms return. A second course of treatment will be allowed six months after the last injection of a previous course of treatment.

The revision of this verbiage clarifies the conditions under which a second course of treatment will be allowed. It is also intended to remove the impression that FCSO will allow multiple courses of treatment.

EFFECTIVE DATE

This revision is effective for services rendered on or after April 01, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcsso.com> on or after this effective date.

92567: TYMPANOMETRY—REVISION TO THE LCD

This local coverage determination (LCD) was last revised effective October 1, 2007. Since that time, revisions were made to the "Indications and Limitations of Coverage and/or Medical Necessity", and "Documentation Requirements" sections of the LCD.

This revision was based on CMS change request (CR) 5717, transmittals 84 and 1470, dated February 29, 2008. This CR included the updates made to the language in the *Medicare Benefit Policy Manual* (Pub.100-02, chapter 16, section 100) and *Medicare Claims Processing Manual* (Pub. 100-04, chapter 12, section 30.3). The language in the manuals was revised to clarify and improve understanding of the policies for audiology services.

EFFECTIVE DATE

This LCD revision is effective for claims processed on or after April 7, 2008, for services rendered on or after April 1, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcsso.com> on or after this effective date.

98925: OSTEOPATHIC MANIPULATIVE TREATMENT—REVISION TO THE LCD

The local coverage determination (LCD) for osteopathic manipulative treatment (OMT) was effective on September 30, 2007. Since that time, the LCD has been revised. Language has been added to expand the utilization guidelines and documentation guidelines pertaining to frequency, duration and documentation of services.

OMT is a distinct manual procedure employed by a physician that aims to optimize a patient's health and function. OMT is the therapeutic application of manually guided forces by an osteopathic physician to improve physiologic function and/or support homeostasis that have been altered by somatic dysfunction.

EFFECTIVE DATE

This revision to the LCD is effective for services rendered on or after June 30, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcsso.com> on or after this effective date.

ADDITIONAL INFORMATION

J1440: G-CSF (FILGRASTIM, NEUPOGEN®): CLARIFICATION ON CORRECT ADMINISTRATION OF NEUPOGEN—ARTICLE REVISION

First Coast Service Options, Inc. (FCSO) has discovered, through medical review and subsequent data analysis, that providers are inappropriately administering Neupogen® (J1440 and J1441) to patients who are receiving a chemotherapeutic agent.

Neupogen® is not a cancer chemotherapy agent. It is a class II hematopoietic growth factor that acts on progenitor cells. Because Neupogen® acts only on progenitor cells that are already committed to one pathway, it increases only the neutrophil count. The local coverage determination (LCD) for Neupogen® outlines the Food and Drug Administration (FDA) approved indications and the off-label indications FCSO will cover when the medical necessity criteria are met.

Under the "limitations" section of the LCD, it is outlined that Neupogen should not be given within 24 hours before or after a dose of a chemotherapeutic agent, as rapidly dividing myeloid cells are potentially sensitive to these agents. This instruction is also outlined in the FDA approved label. This rule applies to any indication in the LCD that requires the administration of a chemotherapeutic agent.

An example of inappropriate administration found during medical review of claims shows that providers are administering Neupogen® the day before, the day of and the day after chemotherapy administration. In the cases reviewed, patients received one injection of Neupogen® less than 12 hours before chemotherapy, then received an injection

J1440: G-CSF (Filgrastim, Neupogen®): Clarification on Correct Administration of Neupogen—Article Revision, continued

immediately following chemotherapy infusion and received a Neupogen® injection, the next day, less than 12 hours after the chemotherapy infusion.

FCSO would like to reiterate to providers that the continued practice of inappropriate administration of Neupogen® might lead to medical review of documentation. FCSO does not expect to see Neupogen® billed the day before, the day of or the day after chemotherapy administration. If providers bill Neupogen® the day before or the day after chemotherapy administration, the medical record must show that Neupogen® was not given less than 24 hours before and/or less than 24 hours after chemotherapy and that this requirement is documented in the medical record. Claims that cannot support this requirement may be denied as not medically necessary.

FCSO strongly encourages providers to review the current LCD for Neupogen® to ensure their patients meet the coverage criteria outlined for each indication and that all other documentation and utilization requirements are met. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date. Providers who feel the language in the LCD is not appropriate can refer to FCSOs reconsideration process located on our Web site.

THERSVCS: THERAPY AND REHABILITATION SERVICES—CODING GUIDELINE REVISION

The “coding guidelineS” attachment for the therapy and rehabilitation services local coverage determination (LCD) was last revised on January 1, 2008. Since that time the “coding guidelines” attachment has been revised. Language has been added based on change request (CR) 5717, transmittals 84 and 1470, dated February 29, 2008. This CR outlines updated audiology policies. Language regarding speech-language pathology services has been added according to the revised manual language for this CR. A complete discussion of the updated audiology policies may be found in Pub. 100-02, *Medicare Benefit Policy Manual*, chapter 15, section 80.3 and in Pub 100-04, *Medicare Claims Processing Manual*, chapter 12, section 30.3.

EFFECTIVE DATE

Revisions are effective for claims processed on or after April 7, 2008 for services rendered on or after April 1, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

CONNECTICUT ONLY - NEW LCD

64470: PARAVERTEBRAL FACET JOINT BLOCKS—NEW LCD

Medicare will consider facet joint blocks to be reasonable and necessary for chronic pain (persistent pain for three (3) months or greater) suspected to originate from the facet joint. Facet joint blocks are one of the methods used to document/confirm suspicions of posterior element biochemical pain of the spine.

A paravertebral facet joint represents the articulation of the posterior elements of one vertebra with its neighboring vertebrae. For purposes of this local coverage determination (LCD), the facet joint is noted at a specific level, by the vertebrae that form it (e.g., C4-5, L2-3). It is further noted that there are two (2) facet joints at each level, left and right. During a paravertebral facet joint block procedure, a needle is placed in the facet joint or along the medial branches that innervate the joints under fluoroscopic guidance and a local anesthetic and/or steroid is injected.

This local coverage determination (LCD) was developed to include indications and limitations of coverage, ICD-9-CM codes that support medical necessity, documentation requirements, utilization guidelines, and coding guidelines.

EFFECTIVE DATE

This LCD is effective for services rendered on or after June 30, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

SIGN UP TO OUR eNEWS ELECTRONIC MAILING LIST

Join our **eNews** mailing list and receive urgent and other critical information issued by First Coast Service Options, Inc. (FCSO), your Medicare carrier. By signing up, you will receive automatic e-mail notification when new or updated information is posted to the provider education Web site. It's very easy to do. Simply go to our Web site <http://www.fcso.com>, select Medicare Providers, Connecticut or Florida, click on the “eNews” link located on the upper-right-hand corner of the page and follow the prompts.

CONNECTICUT ONLY - REVISION TO THE LCDs

62263: EPIDURAL—REVISION TO THE LCD

This local coverage determination (LCD) was last revised effective June 4, 2007. Since that time, a major revision was made to the “Indications and Limitations of Coverage and/or Medical Necessity”, “ICD-9 Codes that Support Medical Necessity”, “Documentation Requirements”, and “Utilization Guidelines” sections of the LCD.

The previous LCD was completely stricken and the Contractor’s Determination Number was changed from 62263 to 62310. Major changes to the LCD include:

- Procedure codes 00277, 62263, 62264, 62280, 62281, 62282, 62318, and 62319 and the indication for acute pain were removed.
- Procedure codes 62310, 62311, 64479, 64480, 64483, and 64484 and the indications and limitations for chronic pain was addressed.
- The list of ICD-9-CM codes that support medical necessity were revised.
- Language was added requiring fluoroscopic guidance to be reported with transforaminal epidural injections.
- Utilization Guidelines were added.
- Previous coding guidelines were deleted and replaced with coding information related to fluoroscopic guidance and diagnosis code V58.61.

EFFECTIVE DATE

This LCD revision is effective for services rendered on or after June 30, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

Italicized and/or quoted material is excerpted from the American Medical Association Current Procedural Terminology. CPT codes, descriptions and other data only are copyrighted 2007 American Medical Association (or other such date of publication of CPT). All rights reserved. Applicable FARS/DFARS apply.

92552: AUDIOMETRY—REVISION TO THE LCD

This local coverage determination (LCD) was last revised effective October 1, 2007. Since that time, revisions were made to the “Indications and Limitations of Coverage and/or Medical Necessity”, “Documentation Requirements”, and “Utilization Guidelines” sections of the LCD.

This revision was based on CMS change request (CR) 5717, transmittals 84 and 1470, dated February 29, 2008. This CR included the updates made to the language in the *Medicare Benefit Policy Manual* (Pub.100-02, chapter 16, section 100) and *Medicare Claims Processing Manual* (Pub. 100-04, chapter 12, section 30.3). The language in the manuals was revised to clarify and improve understanding of the policies for audiology services.

EFFECTIVE DATE

This LCD revision is effective for claims processed on or after April 7, 2008, for services rendered on or after April 1, 2008. The full text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

CONNECTICUT ONLY - DRAFT LCDs

77371: DRAFT LCD FOR STEREOTACTIC RADIOSURGERY (SRS)

The draft local coverage determination (LCD) for stereotactic radiosurgery (SRS) was posted for comment on January 22, 2007. During the comment period, First Coast Services Options, Inc. (FCSO) received a large number of comments related to the coverage and coding of these services. Since that time, it has been determined that FCSO will not be finalizing this draft LCD.

77373: DRAFT LCD FOR STEREOTACTIC BODY RADIATION THERAPY (SBRT)

The draft local coverage determination (LCD) for stereotactic body radiation therapy (SBRT) was posted for comment on January 22, 2007. During the comment period, First Coast Services Options, Inc. (FCSO) received a large number of comments related to the coverage and coding of these services. Since that time, it has been determined that FCSO will not be finalizing this draft LCD.

FLORIDA ONLY - REVISION TO THE LCD

92541: VESTIBULAR FUNCTION TESTS—REVISION TO THE LCD

The local coverage determination (LCD) for vestibular function tests was last revised on May 15, 2007. Since that time, the LCD has been revised. Language has been added based on change request (CR) 5717, transmittals 84 and 1470, dated February 29, 2008. This CR outlines updated audiology policies. Language regarding orders and language regarding the definition of a qualified audiologist has been revised according to the new manual language for this CR. A statement on where to locate a complete discussion of audiology policies was also added to the LCD and the coding guideline. This information may be found in Pub. 100-02, *Medicare Benefit Policy Manual*, chapter 15, section 80.3 and in Pub. 100-04, *Medicare Claims Processing Manual*, chapter 12, section 30.3.

EFFECTIVE DATE

This revision to the LCD is effective for claims processed on or after April 7, 2008, for services rendered on or after April 1, 2008. The full-text of this LCD is available through our provider education Web site at <http://www.fcso.com> on or after this effective date.

FLORIDA ONLY - ADDITIONAL INFORMATION

CPT CODE 78580—PULMONARY PERFUSION IMAGING, PARTICULATE

The Statistical and Medical Data Analysis department conducted an analysis of the Medicare Part B claims data for CPT code 78580 (*Pulmonary perfusion imaging, particulate*). The analysis was performed to gain an understanding of why the Florida carrier paid 500 percent more than the national average based on July through December 2006 national Medicare data. Paid claims data for the time period of January to July 2007 revealed the carrier's payments continued to grow reaching 600 percent more than the national average.

Results of the analysis show cardiologists are drivers of this anomaly as they are billing CPT code 78580 on the same date of service as CPT code 78465 (*Myocardial perfusion imaging: tomographic [SPECT], multiple studies [including attenuation correction when performed], at rest and/or stress [exercise and/or pharmacologic] and redistribution and/or rest injection, with or without quantification*). Inquiries to Florida physicians revealed practitioners are utilizing CPT code 78580 to represent the

calculation of the heart lung ratio obtained during the processing of a SPECT myocardial perfusion procedure.

The Society of Nuclear Medicine (SNM) addressed this issue April 2007, stating the phrase "with quantification" would include calculation of the heart-lung ratio if obtained during a myocardial SPECT perfusion procedure.

First Coast Service Options, Inc. concurs with the SNM interpretation. It is therefore, not appropriate to bill for a lung scan with or without any modifier for the calculation of the heart-lung ratio during the processing of a SPECT myocardial perfusion procedure. In addition, information related to the above was added to the coding guidelines for local coverage determination (LCD) L5960 (78460 Cardiovascular Nuclear Imaging Studies) effective for services rendered on or after April 22, 2008.

Italicized and/or quoted material is excerpted from the American Medical Association Current Procedural Terminology. CPT codes, descriptions and other data only are copyrighted 2007 American Medical Association (or other such date of publication of CPT). All rights reserved. Applicable FARS/DFARS apply.

FLORIDA ONLY - DRAFT LCDs

77371: DRAFT LCD FOR STEREOTACTIC RADIOSURGERY (SRS)

The draft local coverage determination (LCD) for stereotactic radiosurgery (SRS) was posted for comment on January 22,

2007. During the comment period, First Coast Services Options, Inc. (FCSO) received a large number of comments related to the coverage and coding of these services. Since that time, it has been determined that FCSO will not be finalizing this draft. A new draft LCD will be considered in the future.

77373: DRAFT LCD FOR STEREOTACTIC BODY RADIATION THERAPY (SBRT)

The draft local coverage determination (LCD) for stereotactic body radiation therapy (SBRT) was posted for comment on

January 22, 2007. During the comment period, First Coast Services Options, Inc. (FCSO) received a large number of comments related to the coverage and coding of these services. Since that time, it has been determined that FCSO will not be finalizing this draft. A new draft LCD will be considered in the future.

CONNECTICUT EDUCATIONAL RESOURCES

UPCOMING PROVIDER OUTREACH AND EDUCATION EVENTS
MAY 2008 – JUNE 2008

EVALUATION & MANAGEMENT - "INPATIENT HOSPITAL SERVICES" TELECONFERENCE

Topic: Inpatient Hospital Services
When: May 20, 2008
Time: 11:30 a.m. – 1:00 p.m.
Type of Event: Teleconference

HOT TOPICS TELECONFERENCE

Topic: Recent Medicare Changes, New/Revised Local Coverage Determinations (LCDs) and How to Avoid Top Claim Denials and Comprehensive Error Rate Testing (CERT) Errors
When: May 21, 2008
Time: 11:30 a.m. – 1:00 p.m.
Type of Event: Teleconference

ASK THE CONTRACTOR TELECONFERENCE

Topic: Overview of New Medicare Competitive Bidding Program for DMEPOS
When: June 10, 2008
Time: 11:30 a.m. – 1:00 p.m.
Type of Event: Teleconference

TWO EASY WAYS TO REGISTER

Online – Simply log on to your account on our provider training Web site at www.fcsomedicaretraining.com and select the course for which you wish to register. Class materials will be available under "My Courses" no later than one day before the event. If you need assistance, please contact our FCSO Medicare training help desk by calling 866-756-9160 or sending an e-mail to fcsohelp@geolearning.com.

- To locate this course on the provider training Web site:
• Click "Course Catalog" from the top navigation bar, then click the "Catalog" link in the middle of the page
• Type a keyword in the search box for the course you are interested in (such as "ASC" or "Hot Topics") and hit the "Search" button.
• In the short list of courses that will appear, click the link for the course you're interested in and then click the "Preview Schedule" button at the bottom of the class description page.
• On the Instructor-Led Training (ILT) Schedule page, locate the line that has the course you are interested in and click the "Register" link in the Options column.
• First-time user? Please set up an account using the instructions located at www.connecticutmedicare.com/Education/108651.asp in order to register for a class and obtain materials.

Fax – If you would like to participate in any of these events and do not have access to the Internet, please leave a message on our Registration Hotline at (203) 634-5527 in order to have a paper registration form faxed to you.

Please Note:

- Pre-registration is required for all teleconferences, webcasts and in-person educational seminars.
• Dates and times are subject to change prior to event advertisement.

Registrant's Name: _____
Registrant's Title: _____
Provider's Name: _____
Telephone Number: _____ Fax Number: _____
Email Address: _____
Provider Address: _____
City, State, ZIP Code: _____

There's always something going on in Provider Outreach & Education! Keep checking our Web site at www.fcso.com and listening to information on our Registration Hotline at (203) 634-5527 for details about upcoming events.

Don't have time to attend an event? Check out our provider training Web site at www.fcsomedicaretraining.com for self-paced Web-based training classes.

FLORIDA EDUCATIONAL RESOURCES

UPCOMING PROVIDER OUTREACH AND EDUCATION EVENTS

MAY 2008 – JUNE 2008

HOT TOPICS: MEDICARE UPDATES TELECONFERENCE

When: May 15, 2008
 Time: 11:30 a.m. – 12:30 p.m.
 Type of Event: Teleconference

EVALUATION & MANAGEMENT – "INPATIENT HOSPITAL SERVICES" TELECONFERENCE

When: May 20, 2008
 Time: 11:30 a.m. – 1:00 p.m.
 Type of Event: Teleconference

ASK THE CONTRACTOR TELECONFERENCE

Topic: Overview of New Medicare Competitive Bidding Program for DMEPOS
 When: June 10, 2008
 Time: 11:30 a.m. – 1:00 p.m.
 Type of Event: Teleconference

EVALUATION & MANAGEMENT – "CONSULTATION SERVICES" TELECONFERENCE

When: June 17, 2008
 Time: 11:30 a.m. – 1:00 p.m.
 Type of Event: Teleconference

TWO EASY WAYS TO REGISTER

Online – Simply log on to your account on our provider training Web site at www.fcsomedicaretraining.com and select the course you wish to register for. Class materials will be available under "My Courses" no later than one day before the event.

PLEASE NOTE:

- Pre-registration is required for all teleconferences, Webcasts and in-person educational seminars.
- Dates and times are subject to change prior to event advertisement.

Registrant's Name: _____
 Registrant's Title: _____
 Provider's Name: _____
 Telephone Number: _____ Fax Number: _____
 Email Address: _____
 Provider Address: _____
 City, State, ZIP Code: _____

More educational events (teleconferences, webcasts, etc.) are being planned to help providers with hot issues. Keep checking our Web site, www.fcsso.com, or listening to information on the FCSO Provider Education Registration Hotline, 1-904-791-8103, for details and newly scheduled events.

Online – Simply log on to your account on our provider training Web site at www.fcsomedicaretraining.com and select the course you wish to register for. Class materials will be available under "My Courses" no later than one day before the event.

Fax – Providers without Internet access can leave a message on our Registration Hotline at 904-791-8103 requesting a fax registration form. Class materials will be faxed to you the day of the event.

TIPS FOR USING THE FCSO PROVIDER TRAINING WEBSITE

The best way to search and register for Florida events on www.fcsomedicaretraining.com is by clicking on the following links in this order:

- "Course Catalog" from top navigation bar
- "Catalog" in the middle of the page
- "Browse Catalog" on the right of the search box
- "FL – Part B or FL – Part A" from list in the middle of the page.

Select the specific session you're interested in, click the "Preview Schedule" button at the bottom of the page. On the Instructor-Led Training (ILT) Schedule page, locate the line that has the course you are interested in and click the "Register" link in the Options column.

If you need assistance, please contact our FCSO Medicare training help desk by calling 866-756-9160 or sending an email to fcsohelp@geolearning.com.

CONNECTICUT MEDICARE PART B MAIL DIRECTORY

Connecticut Medicare Part B welcomes any questions that you may have regarding the Medicare Part B program. Always be sure to clearly explain your question or concern. This will help our staff to know exactly what issues to address when developing a response to your inquiry.

Please submit your questions to the appropriate department. This will ensure that your concerns are handled in a proper and timely manner. This can be achieved by including an Attention Line below the address on the envelope. Listed below is a directory of departments that includes the issues that you would address to their attention.

With the exception of Redeterminations and Medicare EDI, please submit all correspondence with the appropriate attention line to:

**Attention: (insert dept name)
Medicare Part B CT
P.O. Box 45010
Jacksonville, FL 32232-5010**

Attention: Correspondence

The Correspondence attention line is used for inquiries pertaining to general issues regarding Medicare Part B. Some examples of these issues are deductibles, assignment, and beneficiary address changes. Do not use words such as *REVIEW* or *RECHECK* when sending general correspondence.

Attention: Financial Services

Use this attention line to return duplicate payments or overpayment refunds.

Attention: Fraud and Abuse

If you encounter what you believe is suspected, potential, or possible fraud or abuse of the Medicare program, we encourage you to contact this department.

Attention: Medical Review

Questions regarding LMRPs/LCDs and correct documentation for evaluation and management services are handled by this department. Documentation for off-label chemotherapy use should also be submitted to the Medical Review Department.

Attention: MSP

Write to the Medicare Secondary Payer (MSP) department when submitting an Explanation of Benefits from a primary insurance, Exhaust letters from Auto Liability claims, and MSP calculation review requests.

Attention: Pricing/Provider Maintenance

Address your envelope to this department to apply for a new provider number, change a business or billing address of a provider, or to make any changes in the status of a provider. This department also handles fee schedule requests and inquiries, participation requests, and UPIN requests.

Attention: Resolutions

Use the Resolutions attention line when inquiring or submitting information

regarding dates of death, incorrect Medicare (HIC) numbers, incorrect beneficiary information, etc.

MAILING ADDRESS EXCEPTIONS

We have established special P.O. boxes to use when mailing your redeterminations and hearings requests, paper claims, or to contact Medicare EDI:

Redeterminations/Appeals

Please mail only your requests for redeterminations to this P.O. Box. *DO NOT* send new claims, general correspondence, or other documents to this location; doing so will cause a delay in the processing of that item. If you believe the payment or determination is incorrect and want a claim to be reconsidered, then send it to the attention of the review department. Requests for redeterminations must be made within 120 days of the date of the Medicare Summary Notice. These requests should not include redetermination requests on Medicare Secondary Pay calculations. Claims that are denied for return/reject need to be resubmitted and should **not** be sent as a redetermination. These resubmitted claims should be sent in as new claims.

Medicare Part B CT Appeals
First Coast Service Options, Inc.
P.O. Box 45041
Jacksonville, FL 32232-5041

Electronic Media Claims (EMC)/ The Electronic Data Interchange (EDI)

The EDI department handles questions and provides information on electronic claims submission (EMC).

Medicare Part B CT Medicare EDI
P.O. Box 44071
Jacksonville, FL 32231-4071

Claims

The Health Insurance Portability and Accountability Act (HIPAA) requires electronic submission of most types of Medicare claims. We realize, however, that on occasion it is necessary to submit a paper claim. When this happens, submit your claims on the approved red-and-white Form CMS-1500 to:

Medicare Part B CT Claims
P.O. Box 44234
Jacksonville, FL 32231-4234

Freedom of Information (FOIA)

Freedom of Information Act Requests
Post Office Box 2078
Jacksonville, Florida 32231

CONNECTICUT MEDICARE PHONE NUMBERS

BENEFICIARY SERVICES

1-800-MEDICARE (toll-free)
1-866-359-3614 (*hearing impaired*)
First Coast Service Options, Inc.

PROVIDER SERVICES

Medicare Part B
1-888-760-6950
FAX : 1-904-361-0695
E-mail Address:
AskCTMedicare@fcso.com

Appeals

1-866-535-6790, option 1

Medicare Secondary Payer

1-866-535-6790, option 2

Provider Enrollment

1-866-535-6790, option 4

Interactive Voice Response

1-866-419-9455

Electronic Data Interchange (EDI) Enrollment

1-203-639-3160, option 1

PC-ACE® PRO-32

1-203-639-3160, option 2

Marketing and Reject Report Issues

1-203-639-3160, option 4

Format, Testing, and Remittance Issues

1-203-639-3160, option 5

Electronic Funds Transfer Information

1-203-639-3219

Hospital Services

National Government Services
Medicare Part A
1-888-855-4356

Durable Medical Equipment NHC

DME MAC Medicare Part B
1-866-419-9458

Railroad Retirees

Palmetto GBA
Medicare Part B
1-877-288-7600

Quality of Care

Qualidign (Peer Review Organization)
1-800-553-7590

OTHER HELPFUL NUMBERS

Social Security Administration
1-800-772-1213

**To Report Lost or
Stolen Medicare Cards**
1-800-772-1213

**Health Insurance Counseling
Program (CHOICES)/Area Agency on
Aging**
1-800-994-9422

**Department of Social Services/
ConnMap**
1-800-842-1508

**ConnPACE/
Assistance with Prescription Drugs**
1-800-423-5026 or 1-860-832-9265
(Hartford area or from out of state)

MEDICARE WEB SITES

PROVIDER
Connecticut
<http://www.connecticutmedicare.com>

**Centers for Medicare & Medicaid
Services**
<http://www.cms.hhs.gov>

BENEFICIARY
**Centers for Medicare & Medicaid
Services**
<http://www.medicare.gov>

FLORIDA MEDICARE PART B

MAIL DIRECTORY

CLAIMS SUBMISSIONS

Routine Paper Claims

Medicare Part B
P. O. Box 2525
Jacksonville, FL 32231-0019

Participating Providers

Medicare Part B Participating Providers
P. O. Box 44117
Jacksonville, FL 32231-4117

Chiropractic Claims

Medicare Part B Chiropractic Unit
P. O. Box 44067
Jacksonville, FL 32231-4067

Ambulance Claims

Medicare Part B Ambulance Dept.
P. O. Box 44099
Jacksonville, FL 32231-4099

Medicare Secondary Payer

Medicare Part B Secondary Payer Dept.
P. O. Box 44078
Jacksonville, FL 32231-4078

ESRD Claims

Medicare Part B ESRD Claims
P. O. Box 45236
Jacksonville, FL 32232-5236

COMMUNICATION

Redetermination Requests

Medicare Part B Claims Review
P.O. Box 2360
Jacksonville, FL 32231-0018

Fair Hearing Requests

Medicare Hearings
P.O. Box 45156
Jacksonville FL 32232-5156

Freedom of Information Act

Freedom of Information Act Requests
Post Office Box 2078
Jacksonville, Florida 32231

Administrative Law Judge Hearing

Q2 Administrators, LLC
Part B QIC South Operations
P.O. Box 183092
Columbus, Ohio 43218-3092
Attn: Administration Manager

Status/General Inquiries

Medicare Part B Correspondence
P. O. Box 2360
Jacksonville, FL 32231-0018

Overpayments

Medicare Part B Financial Services
P. O. Box 44141
Jacksonville, FL 32231-4141

DURABLE MEDICAL EQUIPMENT (DME)

DME, Orthotic or Prosthetic Claims

Cigna Government Services
P.O. Box 20010
Nashville, Tennessee 37202

ELECTRONIC MEDIA CLAIMS (EMC)

EMC Claims, Agreements and Inquiries

Medicare EDI
P. O. Box 44071
Jacksonville, FL 32231-4071

MEDICARE PART B ADDITIONAL DEVELOPMENT

Within 40 days of initial request:
Medicare Part B Claims
P. O. Box 2537
Jacksonville, FL 32231-0020

**Over 40 days of initial request:
Submit the charge(s) in question,
including information requested, as
you would a new claim, to:**

Medicare Part B Claims
P. O. Box 2525
Jacksonville, FL 32231-0019

MISCELLANEOUS

Provider Participation and Group
Membership Issues; Written Requests for
UPINs, Profiles & Fee Schedules:
Medicare Enrollment
P. O. Box 44021
Jacksonville, FL 32231-4021

Provider Change of Address:

Medicare Registration
P. O. Box 44021
Jacksonville, FL 32231-4021
and

Provider Enrollment Department
Blue Cross Blue Shield of Florida
P. O. Box 41109
Jacksonville, FL 32203-1109

Provider Education

For Educational Purposes and Review of Customary/Prevailing Charges or Fee Schedule:

Medicare Part B
Provider Outreach and Education
P. O. Box 2078
Jacksonville, FL 32231-0048

For Education Event Registration:

Medicare Part B
Medicare Education and Outreach
P. O. Box 45157
Jacksonville, FL 32232-5157

Limiting Charge Issues:

For Processing Errors:
Medicare Part B
P. O. Box 2360
Jacksonville, FL 32231-0048

For Refund Verification:

Medicare Part B
Compliance Monitoring
P. O. Box 2078
Jacksonville, FL 32231-0048

Medicare Claims for Railroad

Retirees:
Palmetto GBA
Railroad Medicare Part B
P. O. Box 10066
Augusta, GA 30999-0001

Fraud and Abuse

First Coast Service Options, Inc.
Complaint Processing Unit
P. O. Box 45087
Jacksonville, FL 32232-5087

FLORIDA MEDICARE PHONE NUMBERS

PROVIDERS

Toll-Free

Customer Service:
1-866-454-9007
Interactive Voice Response (IVR):
1-877-847-4992

E-mail Address:

AskFloridaB@fcso.com

FAX: 1-904-361-0696

BENEFICIARY

Toll-Free:

1-800-MEDICARE
Hearing Impaired:
1-800-754-7820

Note: The toll-free customer service lines are reserved for Medicare beneficiaries only. Use of this line by providers is not permitted and may be considered program abuse.

For Education Event Registration (not toll-free):
1-904-791-8103

EMC

Format Issues & Testing:

1-904-354-5977 option 4

Start-Up & Front-End Edits/Rejects:

1-904-791-8767 option 1

Electronic Funds Transfer

1-904-791-8016

Electronic Remittance Advice, Electronic Claim Status, & Electronic Eligibility:

1-904-791-6895

PC-ACE Support:

1-904-355-0313

Marketing:

1-904-791-8767 option 1

New Installations:

(new electronic senders; change of
address or phone number for senders):
1-904-791-8608

Help Desk:

(Confirmation/Transmission):
1-904-905-8880 option 1

DME, ORTHOTIC OR PROSTHETIC CLAIMS

Cigna Government Services
1-866-270-4909

MEDICARE PART A

Toll-Free:
1-866-270-4909

MEDICARE WEB SITES

PROVIDER

Florida Medicare Contractor

www.floridamedicare.com

Centers for Medicare & Medicaid Services

www.cms.hhs.gov

BENEFICIARIES

Centers for Medicare & Medicaid Services

www.medicare.gov

ORDER FORM

ORDER FORM— 2008 PART B MATERIALS

The following materials are available for purchase. To order these items, please complete and **submit this form along with your check/money order** payable to FCSO with the designated account number indicated below.

Note: Payment for fee schedules cannot be combined with payment for other items; separate payments are required for purchases of items from different accounts.

ITEM	ACCOUNT NUMBER	COST PER ITEM	QUANTITY	TOTAL
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