



TRICARE  
MANAGEMENT ACTIVITY

OFFICE OF THE ASSISTANT SECRETARY OF DEFENSE  
HEALTH AFFAIRS

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**MB&RB**

**CHANGE 152  
6010.55-M  
MAY 17, 2012**

**PUBLICATIONS SYSTEM CHANGE TRANSMITTAL  
FOR  
TRICARE REIMBURSEMENT MANUAL (TRM), AUGUST 2002**

The TRICARE Management Activity has authorized the following addition(s)/revision(s).

**CHANGE TITLE: OUTPATIENT PROSPECTIVE PAYMENT SYSTEM (OPPS) AND RELATED CHANGES**

**CONREQ: 15216**

**PAGE CHANGE(S): See page 2.**

**SUMMARY OF CHANGE(S):** This change revises the methodology in calculating the General Temporary Military Contingency Payment Adjustment (TMCPA) percentage for qualifying hospitals OPPS year four and after, allows non-network TMCPA payments to be underwritten following the applicable financing rules of the contract. Clarifies the observation stay policy for non-OPPS facilities, clarifies that the Partial Hospitalization Program (PHP) and Substance Use Disorder Rehabilitation Facility (SUDRF) per diem reimbursement applies to other providers who are exempt (except for Maryland hospitals) from the TRICARE OPPS, adds new valid procedure code modifiers, and changes the default code for reporting observation stay services.

**EFFECTIVE DATE: As indicated, otherwise upon direction of the Contracting Officer.**

**IMPLEMENTATION DATE: Upon direction of the Contracting Officer.**

**This change is made in conjunction with Aug 2002 TPM, Change No. 157, and Aug 2002 TSM, Change No. 95.**

*Ann N. Fazzini*

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**ATTACHMENT(S): 46 PAGE(S)  
DISTRIBUTION: 6010.55-M**

**CHANGE 152  
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**REMOVE PAGE(S)**

**CHAPTER 7**

Section 2, pages 1 - 3

Section 3, pages 1 and 2

**CHAPTER 13**

Section 3, pages 17 - 56

**INSERT PAGE(S)**

Section 2, pages 1 - 3

Section 3, pages 1 and 2

Section 3, pages 17 - 57

## PSYCHIATRIC PARTIAL HOSPITALIZATION PROGRAM (PHP) REIMBURSEMENT

ISSUE DATE: July 14, 1993

AUTHORITY: [32 CFR 199.14\(a\)\(2\)\(ix\)](#)

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### I. APPLICABILITY

A. This policy is mandatory for reimbursement of services provided by either network or non-network providers. However, alternative network reimbursement methodologies are permitted when approved by the TRICARE Management Activity (TMA) and specifically included in the network provider agreement.

B. Reimbursement of PHPs prior to implementation of the reasonable cost method for Critical Access Hospitals (CAHs) and implementation of Outpatient Prospective Payment System (OPPS)), and thereafter, freestanding psychiatric PHPs **and other providers who are exempt from the TRICARE OPPS and provider PHP services.**

### II. POLICY

A. Per diem payment for psychiatric partial hospitalization services. Psychiatric partial hospitalization services authorized and provided under [32 CFR 199.4\(b\)\(10\)](#) and provided by psychiatric PHPs authorized under [32 CFR 199.4\(b\)\(3\)\(xii\)](#) are reimbursed on the basis of prospectively determined, all-inclusive per diem rates. The per diem payment amount must be accepted as payment in full for all PHP services provided. The following services and supplies are included in the per diem rate approved for an authorized PHP and are not covered even if separately billed by an individual professional provider. Effective on May 1, 2009 (implementation of OPPS), hospital-based PHP services are reimbursed under the hospital OPPS as described in [Chapter 13, Section 2, paragraph G](#).

1. Board. Includes use of the partial hospital facilities such as food service, supervised therapeutically constructed recreational and social activities, etc.

2. Patient assessment. Includes the assessment of each individual accepted by the facility, and must, at a minimum, consist of a physical examination; psychiatric examination; psychological assessment; assessment of physiological, biological and cognitive processes; developmental assessment; family history and assessment; social history and assessment; educational or vocational history and assessment; environmental assessment; and recreational/activities assessment. Assessments conducted within 30 days prior to admission to a partial program may be used if approved and deemed adequate to permit treatment planning by the PHP.

3. Psychological testing and assessment.

4. Treatment services. All services including routine nursing services, group therapy, supplies, equipment and space necessary to fulfill the requirements of each patient's individualized diagnosis and treatment plan (with the exception of the psychotherapy as indicated in [paragraph II.B.1.](#)). All mental health services must be provided by a authorized individual professional provider of mental health services. [Exception: PHPs that employ individuals with master's or doctoral level degrees in a mental health discipline who do not meet the licensure, certification and experience requirements for a qualified mental health provider but are actively working toward licensure or certification, may provide services within the all-inclusive per diem rate but the individual must work under the clinical supervision of a fully qualified mental health provider employed by the PHP.]

5. Ancillary therapies. Includes art, music, dance, occupational, and other such therapies.

6. Overhead and any other services for which the customary practice among similar providers is included as part of the institutional charges.

B. Services which may be billed separately. The following services are not considered as included within the per diem payment amount and may be separately billed when provided by an authorized individual professional provider:

1. Psychotherapy sessions. Professional services provided by an authorized individual professional provider (who is not employed by or under contract with the PHP) for purposes of providing clinical patient care to a patient in the PHP may be cost-shared when billed by the individual professional provider. Any obligation of a professional provider to provide services through employment or contract in a facility or distinct program of a facility would preclude that professional provider from receiving separate TRICARE/CHAMPUS reimbursement on a fee-for-service basis to the extent that those services are covered by the employment or contract arrangement. Psychotherapy services provided outside of the employment/contract arrangement can be reimbursed separately from the PHPs per diem. Professional mental health benefits are limited to a maximum of one session (60 minutes individual, 90 minutes family, etc.) per authorized treatment day not to exceed five sessions in any calendar week in any combination of individual and family therapy. Five sessions per week is an absolute limit, and additional sessions are not covered.

NOTE: Group therapy is strictly included in the per diem and cannot be paid separately even if billed by an individual professional provider.

2. Primary/Attending Provider. When a patient is approved for admission to a PHP, the primary or attending provider (if not contracted or employed by the partial program) may provide psychotherapy only when the care is part of the treatment environment which is the therapeutic partial program. That is why the patient is there--because that level of care and that program have been determined as medically necessary. The therapy must be adapted toward the events and interactions outlined in the treatment plan and be part of the overall partial treatment plan. Involvement as the primary or attending is allowed and covered only if he is part of the coherent and specific plan of treatment arranged in the partial setting. The treatment program must be under the general direction of the psychiatrist

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employed by the program to ensure medication and physical needs of the patients are met and the therapist must be part of the treatment team and treatment plan. An attending provider must come to the treatment plan meetings and his/her care must be coordinated with the treatment team and as part of the treatment plan. Care given independent of this is not covered.

3. Non-mental health related medical services. Those services not normally included in the evaluation and assessment of a partial hospitalization patient and not related to care in the PHP. These medical services are those services medically necessary to treat a broken leg, appendicitis, heart attack, etc., which may necessitate emergency transport to a nearby hospital for medical attention. Ambulance services may be cost-shared when billed for by an authorized provider if determined medically necessary for emergency transport.

C. Per diem rate. For any full-day PHP (minimum of six hours), the maximum per diem payment amount is 40% of the average inpatient per diem amount per case paid to both high and low volume psychiatric hospitals and units established under the mental health per diem reimbursement system. The rates shall be updated to the current year using the same factors as used under the TRICARE mental health per diem reimbursement system. A PHP of less than six hours (with a minimum of three hours) will be paid a per diem rate of 75% of the rate for full-day PHP. TRICARE will not fund the cost of educational services separately from the per diem rate. The hours devoted to education do not count toward the therapeutic half or full-day program. See [Chapter 7, Addendum B](#), for the current maximum rate limits which are to be used as is for the full-day and half-day program.

D. Other requirements. No payment is due for leave days, for days in which treatment is not provided, for days in which the patient does not keep an appointment, or for days in which the duration of the program services was less than three hours.

E. CAHs. Effective December 1, 2009, PHPs in CAHs shall be reimbursed under the reasonable cost method, reference [Chapter 15, Section 1](#).

- END -



## SUBSTANCE USE DISORDER REHABILITATION FACILITIES (SUDRFs) REIMBURSEMENT

ISSUE DATE: June 26, 1995

AUTHORITY: 32 CFR 199.14(a)(1)(ii)(E) and (a)(2)(ix)

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### I. APPLICABILITY

A. This policy is mandatory for reimbursement of services provided by either network or non-network providers. However, alternative network reimbursement methodologies are permitted when approved by the TRICARE Management Activity (TMA) and specifically included in the network provider agreement.

B. The following reimbursement methodology will be used for payment of all Substance Use Disorder Rehabilitation Facilities (SUDRFs) prior to implementation of the reasonable cost method for Critical Access Hospitals (CAHs) and implementation of Outpatient Prospective Payment System (OPPS). Thereafter, this methodology will only be used in the reimbursement of freestanding SUDRFs and other providers who are exempt from the TRICARE OPPS and provider SUDRF services.

### II. ISSUE

Reimbursement of SUDRFs. This includes reimbursement for both inpatient and partial hospitalization for the treatment of substance use disorder rehabilitation care.

### III. POLICY

A. Inpatient SUDRFs. Effective with admissions on or after July 1, 1995, authorized SUDRFs are subject to the DRG-based payment system.

B. Partial hospitalization for the treatment of substance use disorders. Substance use disorder rehabilitation partial hospitalization services are reimbursed on the basis of prospectively determined all-inclusive per diem rates. The per diem payment amount must be accepted as payment in full for all institutional services provided, including board, routine nursing services, ancillary services (includes art, music, dance, occupational and other such therapies), psychological testing and assessments, overhead and any other services for the customary practice among similar providers is included as part of the institutional charges.

C. Outpatient professional services will be reimbursed using the appropriate Healthcare Common Procedure Coding System (HCPCS) code. Payment is the lesser of the billed charge or the CHAMPUS Maximum Allowable Charge (CMAC).

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D. Family therapy provided on an inpatient or outpatient basis will be reimbursed under the CMAC for the procedure code(s) billed.

E. Cost-sharing. Effective for care on or after October 1, 1995, the cost-share for active duty dependents for inpatient substance use disorder services is \$20.00 per day for each day of the inpatient admission. The \$20.00 cost-share amount also applies to substance use disorder rehabilitation care provided in a partial hospitalization setting. The inpatient cost-share applies to the associated services billed separately by the individual professional providers. For care prior to October 1, 1995, the cost-share will be the daily rate or \$25.00, whichever is greater. For retirees and their dependents, the cost-share is 25% of the allowed amount. Since inpatient cost-sharing is being applied, no deductible is to be taken for partial hospitalization regardless of sponsor status. The cost-share for active duty dependents is to be taken from the partial hospitalization facility claim.

- END -

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(b) These queries will be run in subsequent Transitional TMCPA years to determine those network hospitals qualifying for Transitional TMCPAs.

(c) The year end adjustment will be paid approximately four months following the end of the OPPS year. Each year, subsequent adjustments will be issued to the qualifying hospitals for the prior OPPS year to ensure claims that were not Processed To Completion (PTC) the previous year are adjusted. This adjustment payment is separate from the applicable TMCPA percentage in effect during the current transitional year.

EXAMPLE: At the end of the second OPPS year, a qualifying hospital's total TRICARE OPPS payments will be increased by 15%. The hospital will also receive an additional adjustment for the first OPPS year for those claims that were not PTC and included in the prior year's payment. This subsequent adjustment would be paid at the first year's TMCPA percentage of 20%.

(d) The TMA Medical Benefits and Reimbursement (MB&R) shall verify the accuracy of the Transitional TMCPA amounts and provide the Managed Care Support Contractor (MCSC) with a copy of the report noting which hospitals in their region qualify for the Transitional TMCPAs and the amounts to pay. MB&R shall also provide a copy of the report to Contract Resource Management (CRM).

(e) The MCSCs shall submit the Transitional TMCPAs amounts on a voucher in accordance with the requirements of the TRICARE Operations Manual (TOM), [Chapter 3, Section 4](#). The voucher shall be sent electronically to [RM.Invoices@tma.osd.mil](mailto:RM.Invoices@tma.osd.mil) at the TMA CRM Office and to [OPPS.MBRB@tma.osd.mil](mailto:OPPS.MBRB@tma.osd.mil) at the MB&R before releasing payments. The vouchers should contain the following information: hospital name, address, Medicare number or provider number, Tax Identification Number (TIN), and the amount to be paid. Listings shall separate payments for prior OPPS years and the current OPPS year. Additional vouchers shall be submitted, as needed, for voided/staledated checks and/or for reissued or adjusted payments.

(f) CRM shall send an approval to the contractors to issue Transitional TMCPA payments out of the non-financially underwritten bank account based on fund availability.

(g) Hospitals that previously qualified for Transitional TMCPAs but subsequently fell below \$1.5 million revenue threshold would no longer be eligible for the adjustment. However, if a subsequent adjustment for the prior OPPS year results in a hospital exceeding the \$1.5 million revenue threshold, the hospital shall receive the Transitional TMCPA for the prior year.

(h) New hospitals that meet the \$1.5 million revenue threshold would be eligible for the Transitional TMCPA percentage adjustment in effect during the transitional year in which the revenue threshold was met.

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EXAMPLE: A hospital that meets the \$1.5 million revenue threshold in year three of the transition but failed to meet it in year one and two, would receive a percentage adjustment of 10%.

(2) General TMCPAs. The TMA Director, or designee at any time after OPSS implementation, has the authority to adopt, modify and/or extend temporary adjustments for TRICARE network hospitals located within MTF Prime Service Areas (PSAs) and deemed essential for military readiness and support during contingency operations. The TMA Director may approve a General TMCPA for hospitals that serve a disproportionate share of ADSMs and ADDs. In order for a hospital to be considered for a General TMCPA, the hospital's outpatient revenue received for services provided to TRICARE ADSMs and ADDs must have been at least 10% of the hospital's total outpatient revenue received during the previous OPSS year (May 1 through April 30); or the number of OPSS visits by ADSMs and ADDs during that same 12-month period must have been at least 50,000. Billed charges will not be used as the basis for determining a hospital's eligibility for a General TMCPA.

(c) General TMCPA Process for the First OPSS Year (May 1, 2009 through April 30, 2010); Second OPSS Year (May 1, 2010 through April 30, 2011); and Third OPSS Year (May 1, 2011 through April 30, 2012).

1 The Director, TRICARE Regional Office (DTRO), shall conduct a thorough analysis and recommend the appropriate year end adjustment to total OPSS payments for a network hospital qualifying for a General TMCPA.

2 In analyzing and recommending the appropriate year end percentage adjustment, the DTRO will ensure the General TMCPA adjustment does not exceed 95% of the amount that would have been paid prior to implementation of OPSS. Although, the maximum amount that a hospital can receive is 95% of the pre-OPSS amount, this does not infer the hospital is entitled to receive the full 95%. It is the DTRO's discretion on what percentage adjustment is appropriate to ensure access to care (ATC) in a facility requesting a General TMCPA. This applies to TRICARE beneficiaries when TRICARE is the primary payer. The MCSCs shall provide the history of pre-OPSS payments for the analysis to the DTRO.

3 Total TRICARE OPSS payments (including the TTPAs) and transitional TMCPA's, if applicable, of the qualifying hospital will be increased by the Director TMA, or designee, approved adjustment percentage by way of an additional payment after the end of the OPSS year (May 1 through April 30). At the end of the second and third OPSS years, subsequent adjustments will be issued to the qualifying hospitals for the first and second OPSS years to ensure claims that were not PTC the previous year are adjusted. This adjustment payment is separate from the applicable General TMCPA percentage approved for the current OPSS year.

EXAMPLE: Assume a hospital was approved for a General TMCPA of 5% for the first year of OPSS. At the end of the second year, the hospital will receive an adjustment of 5% for the first OPSS year for those claims that were not PTC and included in the prior

year's payment. The General TMCPA is applied to the total OPSS payment amount at year end.

4 General TMCPAs will be reviewed and approved on an annual basis; i.e., General TMCPAs will have to be evaluated on a yearly basis by the DTRO in order to determine if the hospital continues to serve a disproportionate share of ADSMs and ADDs and whether there are any other special circumstances significantly affecting military contingency capabilities. This will include a recommendation for the appropriate OPSS year end adjustment to total OPSS payments.

5 The hospital's request for a General TMCPA for the first OPSS year (May 1, 2009 through April 30, 2010); second OPSS year (May 1, 2010 through April 30, 2011); and third OPSS year (May 1, 2011 through April 30, 2012) shall include the data requirements in paragraph III.A.5.h.(2)(b), and a full 12 months of claims payment data from the OPSS year the General TMCPA is requested.

6 The TMA MB&R shall verify the accuracy of the General TMCPA amounts and provide the MCSC's with a copy of the report noting which hospitals in their region qualify for the General TMCPAs and the amounts to pay. MB&R shall also provide a copy of the report to CRM.

7 The MCSCs shall submit the General TMCPA amounts on a voucher in accordance with requirements of the TOM, Chapter 3, Section 4. The voucher shall be sent electronically to [RM.Invoices@tma.osd.mil](mailto:RM.Invoices@tma.osd.mil) at the TMA CRM Office and to [OPSS.MBRB@tma.osd.mil](mailto:OPSS.MBRB@tma.osd.mil) at the MB&R before releasing payments. The vouchers should contain the following information: hospital name, address, Medicare number or provider number, TIN, and the amount to be paid. Listings shall separate payments for prior OPSS years and the current OPSS year.

8 CRM shall send an approval to the contractors to issue General TMCPA payments out of the non-financially underwritten bank account based on fund availability.

(b) Annual Data Requirements for General TMCPAs for the First OPSS Year (May 1, 2009 through April 30, 2010); Second OPSS Year (May 1, 2010 through April 30, 2011); and Third OPSS Year (May 1, 2011 through April 30, 2012). Hospital required data submissions to the MCSC for review and consideration:

1 The hospital's percent of outpatient revenue derived from ADSM plus ADD OPSS visits; i.e., the outpatient revenue from TRICARE ADSM plus ADD visits divided by total outpatient revenue (TRICARE and non-TRICARE) derived from all other third party payers and private pay during the previous OPSS year; i.e., May 1 through April 30. Reference paragraph III.A.5.h.(2).

2 The number of OPSS visits by ADSMs and ADDs during the previous OPSS year; i.e., May 1 through April 30.

3 Hospital-specific Medicare outpatient CCR based on the hospital's most recent cost reporting period.

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4 Hospital's Medicare outpatient payment to charge ratio based on the corresponding Medicare cost reporting period.

5 The hospital's recommended percentage adjustment as supported by the above data requirement submissions.

(c) Annual MCSC Data Review Requirements for the First OPSS Year (May 1, 2009 through April 30, 2010); Second OPSS Year (May 1, 2010 through April 30, 2011); and Third OPSS Year (May 1, 2011 through April 30, 2012).

1 Data Requirements for Evaluation of Network Adequacy Necessary to Support Military Contingency Operations:

a Number of available primary care and specialist providers in the network locality;

b Availability (including reassignment) of military providers in the locations or nearby;

c Appropriate mix of primary care and specialists needed to satisfy demand and meet appropriate patient access standards (appointment/waiting time, travel distance, etc.);

d Efforts that have been made to create an adequate network, and

e Other cost effective alternatives and other relevant factors.

2 If upon initial evaluation, the MCSC determines the hospital meets the disproportionate share criteria in paragraph III.A.5.h.(2), and is essential for continued network adequacy, the request from the hospital along with the above supporting documentation shall be submitted to the TRICARE Regional Office (TRO) for review and determination.

(d) For the first OPSS year (May 1, 2009 through April 30, 2010); second OPSS year (May 1, 2010 through April 30, 2011); and third OPSS year (May 1, 2011 through April 30, 2012) the DTRO shall conduct a thorough analysis and recommend the appropriate percentage adjustments to be applied for that year; i.e., the General TMCPAs will be reviewed and approved on an annual basis. The recommendation with a cost estimate shall be submitted to the MB&R to be forwarded to the Director, TMA, or designee for review and approval. Disapprovals by the DTRO will not be forwarded to MB&R for TMA Director review and approval.

(e) General TMCPA process for OPSS Year Four and Subsequent Years (May 1, 2012 and after).

1 The hospital's request for a General TMCPA shall include the data requirements in paragraph III.A.5.h.(2)(b)1 through 4.

2 The MCSC shall conduct an initial evaluation and determine if the requesting hospital meets the disproportionate share criteria in [paragraph III.A.5.h.\(2\)](#), and is essential for continued network adequacy. The request from the hospital for a General TMCPA along with the supporting documentation in [paragraph III.A.5.h.\(2\)\(b\)1](#) through [4](#) and [paragraph III.A.5.h.\(2\)\(c\)](#), shall be submitted to the DTRO for review and determination.

3 The DTRO shall request TMA MB&R run a query of claims history to determine if the network hospital qualifies for a General TMCPA, i.e., the hospital's payment-to-cost ratio is less than 1.3 for care provided to ADSMs and ADDs during the previous OPSS year (May 1 through April 30).

4 The DTRO shall review the supporting documentation and the report from TMA MB&R, and determine if the network hospital qualifies for a General TMCPA. The recommendation for approval of a General TMCPA shall be submitted to the MB&R to be forwarded to the Director, TMA, or designee for review and approval. Disapprovals by the DTRO will not be forwarded to MB&R for TMA Director review and approval.

5 If a hospital meets the disproportionate share criteria in [paragraph III.A.5.h.\(2\)](#), and is deemed essential for network adequacy to support military contingency operations, the approved hospital's General TMCPA payment will be set so the hospital's payment-to-cost ratio for TRICARE Hospital Outpatient Department (HOPD) services does not exceed a ratio of 1.30. A hospital cannot be approved for a General TMCPA payment if it results in the hospital earning more than 30% above its costs for TRICARE beneficiaries.

6 Total TRICARE OPSS payments (including the TTPAs and the Transitional TMCPA) of the qualifying hospital will be increased by the Director TMA, or designee, by way of an additional payment after the end of the OPSS year (May 1 through April 30). Subsequent adjustments will be issued to the qualifying hospitals for the prior OPSS year to ensure claims that were not PTC the previous year are adjusted. The adjustment payment is separate from the applicable General TMCPA approved for the current OPSS year.

7 Upon approval of the General TMCPA request by the TMA Director, MB&R shall notify the TRO of the approval. The TRO shall notify the Contracting Officer (CO) who shall send a letter to the MCSC notifying them of the approval.

8 The MCSCs shall submit the General TMCPA amounts on a voucher in accordance with requirements of the TOM, [Chapter 3, Section 4](#). The voucher shall be sent electronically to [RM.Invoices@tma.osd.mil](mailto:RM.Invoices@tma.osd.mil) at the TMA CRM Office before releasing payments. The vouchers should contain the following information: hospital name, address, Medicare number or provider number, TIN, and the amount to be paid. Listings shall separate payments for prior OPSS years and the current OPSS year.

9 CRM shall send an approval to the contractors to issue General TMCPA payments out of the non-financially underwritten bank account based on fund availability.

10 General TMCPAs will be reviewed and approved on an annual basis; i.e., they will have to be evaluated on a yearly basis by the DTRO in order to determine if the hospital continues to serve a disproportionate share of ADSMs and ADDs and whether there are any other special circumstances significantly affecting military contingency capabilities.

(f) TMA Director, or designee review.

1 The Director, TMA or designee is the final approval authority.

2 A decision by the Director TMA or designee to adopt, modify, or extend General TMCPAs is not subject to appeal.

(3) Non-Network TMCPAs.

TMCPAs may also be extended to non-network hospitals on a case-by-case basis for specific procedures where it is determined that the procedures cannot be obtained timely enough from a network hospital. This determination will be based on the MCSC's and TRO's evaluation of network adequacy data related to the specific procedures for which the TMCPA is being requested as outlined under paragraph III.A.5.h.(2)(c). Non-network TMCPAs will be adjusted on a claim-by-claim basis. **The associated costs would be underwritten or non-underwritten following the applicable financing rules of the contract.**

(4) Application of Cost-Sharing.

(a) Transitional and General TMCPAs are not subject to cost-sharing.

(b) Non-network TMCPAs shall be subject to cost-sharing since they are applied on a claim-by-claim basis.

(5) Reimbursement of Transitional, **and** General TMCPA costs shall be paid as pass-through costs. The MCSC does not financially underwrite these costs.

i. Hold Harmless TRICARE Transitional Outpatient Payments (TTOPs).

(1) Effective January 1, 2010, TRICARE adopted Medicare's hold harmless provision for rural hospitals with 100 or fewer beds and all SCHs regardless of bed size. TRICARE will apply the hold harmless provision to these hospitals as long as the provision remains in effect under Medicare.

(2) TTOPs will be made to qualifying hospitals that have OPPS costs that are greater than their TRICARE allowed amounts. The 7.1% increase for SCHs, the TTPAs for ER and clinic visits, Transitional and General TMCPAs, if applicable, will be included in the allowed amounts when determining if a hospital's OPPS costs are greater than their TRICARE allowed amounts.

(3) TRICARE will use a method similar to Medicare to reimburse these hospitals their TTOPs. TRICARE will pay qualifying hospitals an amount equal to 85% of the difference between the estimated OPPS costs and the OPPS payment.

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(4) Process for TTOPs Year One (Effective January 1, 2010, through December 31, 2010) and Subsequent Years.

(a) TMA will run query reports of claims history to determine which hospitals qualify for TTOPs at year end; i.e., those hospitals whose costs exceeded their allowed amounts during the previous TTOPs year (January 1 through December 31).

(b) These query reports will be run in subsequent TTOPs years to determine those hospitals qualifying for TTOPs.

(c) The year end adjustment will be paid approximately six months following the end of the TTOPs year. Each year, subsequent adjustments will be issued to the qualifying hospitals for the prior TTOPs year to ensure claims that were not PTC the previous year are adjusted.

(d) The TMA MB&RB shall provide the MCSC with a copy of the query report noting which hospitals in their region qualify for the TTOPs and the amounts to pay. A copy of the report shall also be provided to TMA's CRM.

(e) The contractor shall process the adjustment payments per the instructions in Section G of their contracts under Invoice and Payment Non-Underwritten - Non-TEDs, Demonstrations. No payments will be sent out without approval from TMA-Aurora (TMA-A), CRM, Budget.

**B. Transitional Pass-Through for Innovative Medical Devices, Drugs, and Biologicals.**

**1. Items Subject to Transitional Pass-Through Payments.**

**a. Current Orphan Drugs.**

A drug or biological that is used for a rare disease or condition with respect to which the drug or biological has been designated under section 526 of the Federal Food, Drug, and Cosmetic Act if payment for the drug or biological as an outpatient hospital service was being made on the first date that the OPPS was implemented.

NOTE: Orphan drugs will be paid separately at the Average Sales Price (ASP) + 6%, which represents a combined payment for acquisition and overhead costs associated with furnishing these products. Orphan drugs will no longer be paid based on the use of drugs because all orphan drugs, both single-indication and multi-indication, will be paid under the same methodology. The TRICARE contractors will not be required to calculate orphan drug payments.

**b. Current Cancer Therapy Drugs, Biologicals and Brachytherapy.**

These items are drugs or biologicals that are used in cancer therapy, including (but not limited to) chemotherapeutic agents, antiemetics, hematopoietic growth factors, colony stimulating factors, biological response modifiers, biphosphonates, and a device of brachytherapy if payment for the drug or biological as an outpatient hospital service was being made on the first date that the OPPS was implemented.

c. Current Radiopharmaceutical Drugs and Biological Products.

A radiopharmaceutical drug or biological product used in diagnostic, monitoring, and therapeutic nuclear medicine procedures if payment for the drug or biological as an outpatient hospital service was being made on the first date that the OPSS was implemented.

d. New Medical Devices, Drugs, and Biologicals.

New medical devices, drugs, and biologic agents, will be subject to transitional pass-through payment in instances where the item was not being paid for as a hospital outpatient service as of December 31, 1996, and where the cost of the item is "not insignificant" in relation to the hospital OPSS payment amount.

2. Items eligible for transitional pass-through payments are generally coded under a Level II HCPCS code with an alpha prefix of "C".

a. Pass-through device categories are identified by SI H.

b. Pass-through drugs and biological agents are identified by SI G.

3. Drugs, Biologicals, and Radiopharmaceuticals With New or Continuing Pass-Through Status in CY 2009.

a. Provide payment for drugs and biologicals with pass-through status that are not part of the Part B drug Competitive Acquisition Program (CAP) at a rate of ASP + 6%, the amount authorized under section 1843(o) of the Social Security Act (SSA) rather than ASP + 4% that would be the otherwise applicable fee schedule portion associated with drug or biological.

b. Provide payment for drugs and biologicals with pass-through status that are not part of the Part B drug CAP at a rate of ASP + 6%, the amount authorized under section 1843(o) of the Act, rather than ASP + 4% that would be the otherwise applicable fee schedule portion associated with drug and biological.

c. The difference between ASP + 4% and ASP + 6%, therefore would be the CY 2009 pass-through payment amount for these drugs and biologicals.

d. Considering diagnostic radiopharmaceuticals to be drugs for pass-through purposes which will be reimbursed based on the ASP methodology; i.e., ASP + 6%.

e. Therapeutic radiopharmaceuticals with pass-through status in CY 2009 will be paid at hospital charges adjusted to cost, the same payment methodology as other therapeutic radiopharmaceuticals in CY 2009.

f. If a drug or biological that has been granted pass-through status for CY 2009 becomes covered under the Part B drug CAP (if the program is reinstated) the Centers for Medicare and Medicaid Services (CMS) will provide payment for Part B Drugs that are

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granted pass-through status and are covered under the Part B drug CAP at the Part B drug CAP rate.

g. Beneficiary copayments/cost-sharing will be based on the entire ASP of the transition pass-through drug or biological.

h. Drugs and biologicals that are continuing pass-through status or have been granted pass-through status as of January 2009 for CY 2009 are displayed in [Figure 13-3-5](#).

**FIGURE 13-3-5 DRUGS AND BIOLOGICALS WITH PASS-THROUGH STATUS IN CY 2009**

CY 2008		CY 2009		
HCPCS	HCPCS	SHORT DESCRIPTOR	SI	APC
C9238	J1953	Levetiracetam injection	G	9238
C9239	J9330	Temsirolimus injection	G	1168
C9240*	J9207	Exabepilone injection	G	9240
C9241	J1267	Doripenem injection	G	9241
C9242	J1453	Fosaprepitant injection	G	9242
C9243	J9033	Bendamustine injection	G	9243
C9244	J2785	Injection, regadenoson	G	9244
C9354	C9354	Veritas collagen matrix, cm2	G	9354
C9355	C935	Neuromatrix nerve cuff, cm	G	9355
C9356	C9356	TendoGlide Tendon prot, cm2	G	9356
C9357	Q4114	Integra flowable wound matri	G	1251
C9358	C9358	SurgiMend, 0.5cm2	G	9358
C9359	C9359	Implant, bone void filler	G	9359
J1300	J1300	Eculizumab injection	G	9236
J1571	J1571	Hepagam b im injection	G	0946
J1573	J1573	Hepagam b intravenous, inj	G	1138
J3488*	J3488	Reclast injection	G	0951
J9225*	J9225	Vantas implant	G	1711
J9226	J9226	Supprelin LA implant	G	1142
J9261	J9261	Nelarabine injection	G	0825
Q4097	J1459	Inj IVIG privitygen 500 mg	G	1214
	C9245	Injection, romiplostim	G	9245
	C9246	Inj, gadoxetate	G	9246
	C9248	Inj, clevidipine butyrate	G	9248

\* Indicates that the drug was paid at a rate determined by the Part B drug CAP methodology (prior to January 1, 2009) while identified as pass-through under the OPSS.

**4. Reduction of Transitional Pass-Through Payments for Diagnostic Radiopharmaceuticals to Offset Costs Packaged Into APC Groups.**

a. Prior to CY 2008, certain diagnostic radiopharmaceuticals were paid separately under the OPSS if their mean per day cost were greater than the applicable year's drug packaging threshold.

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b. In CY 2008, CMS payment for all non-pass-through diagnostic radiopharmaceuticals were packaged as ancillary and supportive items and service.

c. In CY 2009, continued to package payment for all non-pass-through diagnostic radiopharmaceuticals.

d. For OPSS pass-through purposes, radiopharmaceuticals are considered to be “drugs” where the transitional pass-through for the drugs and biologicals is the difference between the amount paid ASP + 4% or the Part B drug CAP rate and the otherwise applicable OPSS payment amount of ASP + 6%.

e. There is currently one radiopharmaceutical with pass-through status under OPSS.

f. New pass-through diagnostic radiopharmaceuticals with no ASP information or CAP rate will be paid at ASP + 6%, while those without ASP information will be paid based on Wholesale Acquisition Cost (WAC) or, if WAC is not available, based on 95% of the product’s most recently published Average Wholesale Price (AWP).

g. Offset Calculations.

(1) An established methodology will be employed to estimate the portion of each APC payment rate that could reasonably be attributed to the cost of an associated device eligible for pass-through payment (the APC device offset).

(2) New pass-through device categories will be evaluated individually to determine if there are device costs packaged into the associated procedural APC payment rate - suggesting that a device offset amount would be appropriate.

h. Effective April 1, 2009, diagnostic radiopharmaceutical HCPCS code C9247, Iobenguane, I-123, diagnostic, per study dose, up to 10 millicuries, has been granted pass-through status under the OPSS and will be assigned SI of G.

(1) Beginning April 1, 2009, payment for HCPCS code C9247 will be made at 106% of ASP if ASP data are submitted by the manufacturer. Otherwise, payment will be made based on the product’s WAC. Further if WAC data is not available, payment will be made at 95% of the AWP.

(2) Effective for nuclear medicine services furnished on and after April 1, 2009, when HCPCS code C9247 is billed on the same claims with a nuclear medicine procedure, the amount of payment for the pass-through diagnostic radiopharmaceutical reported with HCPCS code C9247 will be reduced by the corresponding nuclear medicine procedure’s portion of its APC payment (offset amount) associated with diagnostic radiopharmaceutical; i.e., the payment for HCPCS code C9247 will be reduced by the estimated amount of payment that is attributable to the predecessor radiopharmaceutical that is package into payment for the associated nuclear medicine procedure reported on the same claim as HCPCS code C9247.

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(3) When C9247 is billed on a claim with one or more nuclear medicine procedures, the OPPS Pricer will identify the offset amount or amounts that apply to the nuclear medicine procedures that are reported on the claim.

(4) Where there is a single nuclear medicine procedure reported on the claim with a single occurrence of C9247, the OPPS Pricer will identify a single offset amount for the procedure billed and adjust the offset by the wage index that applies to the hospital submitting the bill.

(5) Where there are multiple nuclear medicine procedures on the claim with a single occurrence of the pass-through radiopharmaceutical, the OPPS Pricer will select the nuclear medicine procedure with the single highest offset amount, and will adjust the selected offset amount by the wage index of the hospital submitting the claim.

(6) When a claim has more than one occurrence of C9247, the OPPS Pricer will rank potential offset amounts associated with the units of nuclear medicine procedures on the claim and identify a total offset amount that takes into account the number of occurrences of the pass-through radiopharmaceutical on the claims and adjust the total offset amount by the wage index of the hospital submitting the claim.

(7) The adjusted offset will be subtracted from the APC payment for the pass-through diagnostic radiopharmaceutical reported with HCPCS code C9247.

(8) The offset will cease to apply when the diagnostic radiopharmaceutical expires from pass-through status.

5. Transitional Pass-Through Device Categories.

a. Excluded Medical Devices.

Equipment, instruments, apparatuses, implements or items that are generally used for diagnostic or therapeutic purposes that are not implanted or incorporated into a body part, and that are used on more than one patient (that is, are reusable), are excluded from pass-through payment. This material is generally considered to be a part of hospital overhead costs reflected in the APC payments.

b. Included Medical Devices.

(1) The following implantable items may be considered for the transitional pass-through payments:

(a) Prosthetic implants (other than dental) that replace all or part of an internal body organ.

(b) Implantable items used in performing diagnostic x-rays, diagnostic laboratory tests, and other diagnostic tests.

NOTE: Any Durable Medical Equipment (DME), orthotics, and prosthetic devices for which transitional pass-through payment does not apply will be paid

under the DMEPOS fee schedule when the hospital is acting as the supplier (paid outside the PPS).

c. Pass-Through Payment Criteria for Devices.

Pass-through payments will be made for new or innovative medical devices that meet the following requirements:

(1) They were not recognized for payment as a hospital outpatient service prior to 1997 (i.e., payment was not being made as of December 31, 1996). However, the medical device shall be treated as meeting the time constraint (i.e., payment was not being made for the device as of December 31, 1996) if either:

(a) The device is described by one of the initial categories established and in effect, or

(b) The device is described by one of the additional categories established and in effect, and

1 An application under the Federal Food, Drug, and Cosmetic Act has been approved; or

2 The device has been cleared for market under section 510(k) of the Federal Food, Drug, and Cosmetic Act; or

3 The device is exempt from the requirements of section 510(k) of the Federal Food, Drug, and Cosmetic Act under section 510(l) or section 510(m) of the Act.

(2) They have been approved/cleared for use by the Food and Drug Administration (FDA).

(3) They are determined to be reasonable and necessary for the diagnosis or treatment of an illness or injury or to improve the functioning of a malformed body part.

(4) They are an integral and subordinate part of the procedure performed, are used for one patient only, are surgically implanted or inserted via a natural or surgically created orifice on incision, and remain with that patient after the patient is released from the hospital outpatient department.

(a) Reprocessed single-use devices that are otherwise eligible for pass-through payment will be considered for payment if they meet FDA's most recent regulatory criteria on single-use devices.

(b) It is expected that hospital charges on claims submitted for pass-through payment for reprocessed single-use devices will reflect the lower cost of these devices.

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NOTE: The FDA published guidance for the processing of single-use devices on August 14, 2000 - "Enforcement Priorities for Single-Use Devices Reprocessed by Third Parties and Hospitals".

(5) They are not equipment, instruments, apparatuses, implements, or such items for which depreciation and financing expenses are recovered as depreciable assets.

(6) They are not materials and supplies such as sutures, clips, or customized surgical kits furnished incidental to a service or procedure.

(7) They are not material such as biologicals or synthetics that may be used to replace human skin.

(8) No existing or previously existing device category is appropriate for the device.

(9) The associated cost is not insignificant in relation to the APC payment for the service in which the innovative medical equipment is packaged.

(10) The new device category must demonstrate that utilization of its devices provide substantial clinical improvement for beneficiaries compared with currently available treatments, including procedures utilizing devices in existing or previously existing device categories.

d. Duration of Transitional Pass-Through Payments.

(1) The duration of transitional pass-through payments for devices is for at least two, but not more than three years. This period begins with the first date on which a transitional pass-through payment is made for any medical device that is described by the category.

(2) The costs of devices no longer eligible for pass-through payments will be packaged into the costs of the procedures with which they are normally billed.

e. General Coding and Billing Instructions and Explanations.

(1) Devices Implanted, Removed, and Implanted Again, Not Associated With Failure (Applies to Transitional Pass-Through Devices Only):

(a) In instances where the physician is required to implant another device because the first device fractured, the hospitals may bill for both devices - the device that resulted in fracture and the one that was implanted into the patient.

(b) It is realized that there may be instances where an implant is tried but later removed due to the device's inability to achieve the necessary surgical result or due to inappropriate size selection of the device by the physician (e.g., physician implants an anchor to bone and the anchor breaks because the bone is too hard or must be replaced with a larger anchor to achieve a desirable result). In such instances, separate reimbursement will be provided for both devices. This situation does not extend to devices that result in failure or

are found to be defective. For failed or defective devices, hospitals are advised to contact the vendor/manufacturer.

NOTE: This applies to transitional pass-through devices only and not to devices packaged into an APC.

(2) Kits. Manufacturers frequently package a number of individual items used in a particular procedure in a kit. Generally, to avoid complicating the category list unnecessarily and to avoid the possibility of double coding, codes for such kits have not been established. However, hospitals are free to purchase and use such kits.

(a) If the kits contain individual items that separately qualify for transitional pass-through payment, these items may be separately billed using applicable codes. Hospitals may not bill for transitional pass-through payments for supplies that may be contained in kits.

(b) HCPCS codes that describe devices without pass-through status and that are packaged in kits with other items used in a particular procedure, hospitals may consider all kit costs in their line-item charge for the associated device/device category HCPCS code that is assigned SI of N for packaged payment (i.e., hospitals may report the total charge for the whole kit with the associated device/device category HCPCS code. Payment for device/device category HCPCS codes without pass-through status is packaged into payment for the procedures in which they are used, and these codes are assigned SI of N. In the case of a device kit, should a hospital choose to report the device charge alone under a device/device category HCPCS code with SI of N, the hospital should report charges for other items that may be included in the kit on a separate line on the claim.

(3) Multiple Units. Hospitals must bill for multiple units of items that qualify for transitional pass-through payments, when such items are used with a single procedure, by entering the number of units used on the bill.

(4) Reprocessed Devices. Hospitals may bill for transitional pass-through payments only for those devices that are "single use." Reprocessed devices may be considered "single use" if they are reprocessed in compliance with the enforcement guidance of the FDA relating to the reprocessing of devices applicable at the time the service is delivered.

f. Current Device Categories Subject to Pass-Through Payment. Two device categories were established for pass-through payment as of January 1, 2007, HCPCS code C1821 (interspinous process distraction device (implantable)) and HCPCS code L8690 (auditory osseointegrated device, includes all internal and external components), will be active categories for pass-through payment for two years as of January 1, 2007, i.e., these categories will expire from pass-through payment as of December 31, 2008.

g. Reduction of Transitional Pass-Through Payments to Offset Costs Packaged into APC Groups.

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(1) Each new device category will be reviewed on a case-by-case basis to determine whether device costs associated with the new category were packaged into the existing APC structure.

(2) If it is determined that, for any new device category, no device costs associated with the new category were packaged into existing APCs, the offset amount for the new category would be set to \$0 for CY 2008.

h. Calculation of Transitional Pass-Through Payment for a Pass-Through Device.

(1) Device pass-through payment is calculated by applying the statewide CCR to the hospital's charges on the claim and subtracting any appropriate pass-through offset. Statewide CCRs are based on the geographical CBSA (two digit = rural, five digit = urban).

(2) The following are two examples of the device pass-through calculations, one incorporating a device offset amount applicable to CY 2003 and the other only applying the CCR (offsets set to \$0 for CY 2005).

(3) The offset adjustment is applied only when a pass-through device is billed in addition to the APC<sup>2</sup>.

Example #1 Transitional Pass-Through Payment Calculation with Offset:

Device: (C1884 - Embolization Protective System)

Device cost = Hospital charge converted to cost = \$1,200.00

Associated procedure: HCPCS Level I<sup>2</sup> code 92982 (APC0083)

Payment rate = \$3,289.42

Coinsurance amount = \$657.88 (standard ADFM who has met his/her yearly deductible)

Total offset amount to be applied for each APC that contains device costs = \$802.06

NOTE: The total offset from the device amount is wage-index adjusted and the multiple procedure discount factor is adjusted before it is subtracted from the device cost. (Refer to [paragraph III.B.5.h.\(4\)](#) for detailed application of discounting factors to offset amounts.) This example assumes a wage index of 1.0000.

Device cost adjusted by total offset amount:

$\$1,200 - \$802.06 = \$397.94$

TRICARE program payment (before wage index adjustment) for APC 0083:

$\$3,289.42 - \$657.88 = \$2,631.54$

TRICARE payment for pass-through device C1884 = \$397.94

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Beneficiary cost-share liability for APC 0083 = \$657.88

Total amount received by provider for APC 0083 and pass-through device C1884:

\$2,631.54	TRICARE program payment for HCPCS Level I <sup>3</sup> code 92982 when used with device code C1884
657.88	Beneficiary coinsurance amount for HCPCS Level I <sup>3</sup> code 92982
<u>397.94</u>	Transitional pass-through payment for device
\$3,687.36	Total amount received by the provider

Example #2 Transitional Pass-Through Payment Calculation without Offset

Device: (C1884 - Embolization Protective System)

Device cost = Hospital charge converted to cost = \$1,500.00

Associated procedure: HCPCS Level I<sup>3</sup> code 92982 (APC0083)

Payment rate = \$3,289.42

Coinsurance amount = \$657.88 (standard ADFM who has met his/her yearly deductible)

Total offset amount to be applied for each APC that contains device costs = \$0.

NOTE: The total offset from the device amount is wage-index adjusted and the multiple procedure discount factor is adjusted before it is subtracted from the device cost. (Refer to [paragraph III.B.5.h.\(4\)](#) for detailed application of discounting factors to offset amounts.) This example assumes a wage index of 1.0000.

Device cost adjusted by total offset amount:

\$1,500 - \$0 = \$1,500

TRICARE program payment (before wage index adjustment) for APC 0083:

\$3,289.42 - \$657.88 = \$2,631.54

TRICARE payment for pass-through device C1884 = \$1,500

Beneficiary cost-share liability for APC 0083 = \$657.88

Total amount received by provider for APC 0083 and pass-through device C1884:

\$2,631.54	TRICARE program payment for HCPCS Level I <sup>3</sup> code 92982 when used with device code C1884
657.88	Beneficiary coinsurance amount for HCPCS Level I <sup>3</sup> code 92982
<u>1,500.00</u>	Transitional pass-through payment for device
\$4,789.42	Total amount received by the provider

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NOTE: Transitional payments for devices (SI=H) are not subject to beneficiary cost-sharing/copayments.

(4) Steps involved in applying multiple discounting factors to offset amounts prior to subtracting from the device cost.

STEP 1: For each APC with an offset multiply the offset by the discount percent (whether it is 50%, 75%, 100%, or 200%) and the units of service.

$$(\text{Offset} \times \text{Discount Rate} \times \text{Units of Service})$$

STEP 2: Sum the products of Step 1.

STEP 3: Wage adjust the sum of the products calculated in Step 2.

$$(\text{Step 2 Amount} \times \text{Labor \%} \times \text{Wage Index}) + \text{Step 2 Amount} \times \text{Nonlabor \%}$$

STEP 4: If the units of service from the procedures with offsets are greater than the device units of service, then Step 3 is adjusted by device units divided by procedure offset units.

$$[(\text{Step 2 Amount} \times \text{Labor \%} \times \text{Wage Index}) + (\text{Step 2 Amount} \times \text{Nonlabor \%}) \times (\text{Device Units} \div \text{Offset Procedure Units})]$$

**otherwise**

$$(\text{Step 2 Amount} \times \text{Labor \%} \times \text{Wage Index}) \text{ Step 2 Amount} \times \text{Non-Labor \%}$$

EXAMPLE: If there are two procedures with offsets but only one device, then the final offset is reduced by 50%.

STEP 5: If there is only one line item with a device, then the amount calculated in Step 4 is subtracted from the line item charge adjusted to cost.

$$[\text{Step 4 Amount} - (\text{Line Item Charge} \times \text{State CCR})]$$

If there are multiple devices, then the amount from Step 4 is allocated to the line items with devices based on their charges.

$$(\text{Line Item Device Charge} \div \text{Sum of Device Charges})$$

**C. Drugs, Biologicals, and Radiopharmaceuticals Without Pass-Through Status.**

1. Radiopharmaceuticals, drugs, and biologicals which do not have pass-through status, are paid in one of three ways:

- a. Packaged payment, or
  - b. Separate payment (individual APCs), or
  - c. Allowable charge.
2. The cost of drugs and radiopharmaceuticals are generally packaged into the APC payment rate for the procedure or treatment with which the products are usually furnished:
- a. Hospitals do not receive separate payment for packaged items and supplies; and
  - b. Hospitals may not bill beneficiaries separately for any such packaged items and supplies whose costs are recognized and paid for within the national OPPS payment rate for the associated procedure or services.
3. Although diagnostic and therapeutic radiopharmaceutical agents are not classified as drugs or biologicals, separate payment has been established for them under the same packaging threshold policy that is applied to drugs and biologicals; i.e., the same adjustments will be applied to the median costs for radiopharmaceuticals that will apply to non-pass-through, separately paid drugs and biologicals.

D. Criteria for Packaging Payment for Drugs, Biologicals and Radiopharmaceuticals.

1. Generally, the cost of drugs and radiopharmaceuticals are packaged into the APC payment rate for the procedure or treatment with which the products are usually furnished. However, packaging for certain drugs and radiopharmaceuticals, especially those that are particularly expensive or rarely used, might result in insufficient payments to hospitals, which could adversely affect beneficiary access to medically necessary services.
2. Payments for drugs and radiopharmaceuticals are packaged into the APCs with which they are billed if the median cost per day for the drug or radiopharmaceutical is less than \$60. Separate APC payment is established for drugs and radiopharmaceuticals for which the median cost per day exceeds \$60.
3. An exception to the packaging rule is being made for injectable oral forms of antiemetics, listed in [Figure 13-3-6](#).

**FIGURE 13-3-6 ANTIEMETICS EXEMPTED FROM CY 2008 \$60 PACKAGING THRESHOLD**

HCPCS CODE	SHORT DESCRIPTOR
J1260	Dolasetron mesylate
J1626	Granisetron HCl Injection
J2405	Ondansetron HCl Injection
J2469	Palonosetron HCl
Q0166	Granisetron HCl 1 mg oral
Q0179	Ondansetron HCl 8 mg oral
Q0180	Dolasetron Mesylate oral

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4. Continuing to package payment for all non-pass-through diagnostic radiopharmaceuticals and contrast agents, regardless of their per day costs for CY 2009.

5. Payment for Drugs, Biologicals, and Radiopharmaceuticals Without Pass-Through Status That Are Not Packaged.

a. "Specified Covered Outpatient Drugs" Classification.

(1) Special classification (i.e., "specified covered outpatient drug") is required for certain separately payable radiopharmaceutical agents and drugs or biologicals for which there are specifically mandated payments.

(2) A "specified covered outpatient drug" is a covered outpatient drug for which a separate APC exists and that is either a radiopharmaceutical agent or drug or biological for which payment was made on a pass-through basis on or before December 31, 2002.

(3) The following drugs and biologicals are designated exceptions to the "specified covered outpatient drugs" definition (i.e., not included within the designated category classification):

(a) A drug or biological for which payment was first made on or after January 1, 2003, under the transitional pass-through payment provision.

(b) A drug or biological for which a temporary HCPCS code has been assigned.

(c) Orphan drugs.

b. Payment of Specified Outpatient Drugs, Biological, and Radiopharmaceuticals.

(1) Specified outpatient drugs and biologicals will be paid a combined rate of the ASP + 4% which is reflective of the present hospital acquisition and overhead costs for separately payable drugs and biologicals under the OPPS. In the absence of ASP data, the WAC will be used for the product to establish the initial payment rate. If the WAC is also unavailable, then payment will be calculated at 95% of the most recent AWP.

(2) Since there is no ASP data for separately payable specified radiopharmaceuticals, reimbursement will be based on charges converted to costs. Refer to [Section 2, Figure 13-2-15](#), for a list of therapeutic radiopharmaceuticals that will continue to be reimbursed under the cost-to-charge methodology up through December 31, 2009.

(a) Therapeutic radiopharmaceuticals must have a mean per day cost of more than \$60 in order to be paid separately.

(b) Diagnostic radiopharmaceuticals and contrast agents are packaged regardless of per day cost since they are ancillary and supportive of the therapeutic procedures in which they are used.

c. Designated SI.

The HCPCS codes for the above three categories of “specified covered outpatient drugs” are designated with the SI K - non-pass-through drugs, biologicals, and radiopharmaceuticals paid under the hospital OPSS (APC Rate). Refer to TMA’s OPSS web site at <http://www.tricare.mil/opss> for APC payment amounts of separately payable drugs, biologicals and radiopharmaceuticals.

6. Payment for Non-Pass-Through Drugs, Biologicals, and Radiopharmaceuticals With HCPCS Codes, But Without OPSS Hospital Claims Data.

a. These new drugs and biologicals with HCPCS codes as of January 1, 2008, but which do not have pass-through status and are without OPSS hospital claims data, will be paid at ASP + 4% consistent with its final payment methodology for other separately payable non-pass-through drugs and biologicals.

b. Payment for all new non-pass-through diagnostic radiopharmaceuticals will be packaged.

c. In the absence of ASP data, the WAC will be used for the product to establish the initial payment rate for new non-pass-through drugs and biologicals with HCPCS codes, but which are without OPSS claims data. If the WAC is also unavailable, payment will be made at 95% of the product’s most recent AWP.

d. SI K will be assigned to HCPCS codes for new drugs and biologicals for which pass-through application has not been received.

e. Payment for new therapeutic radiopharmaceuticals with HCPCS codes as of January 1, 2008, but which do not have pass-through status, will be assigned SI H and continue to be reimbursed under the cost-to-charge methodology up through December 31, 2009.

f. In order to determine the packaging status of these items for CY 2008 an estimate of the per day cost of each of these items was calculated by multiplying the payment rate for each product based on ASP + 4%, by a estimated average number of units of each product that would typically be furnished to a patient during one administration in the hospital outpatient setting. Items for which the estimated per day cost is less than or equal to \$60 will be packaged. For drugs currently covered under the CAP the payment rates calculated under that program that were in effect as of April 1, 2008 will be used for purposes of packaging decisions.

7. Drugs and Biologicals Not Eligible for Pass-Through Status and Receiving Separate Non-Pass-Through Payment.

a. Payment will be based on median costs derived from CY claims data for drugs and biologicals that have been:

(1) Separately paid since implementation of the OPSS under Medicare, but were not eligible for pass-through status; and

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(2) Historically packaged with the procedures with which they were billed, even though their median cost per day was above the \$60 packaging threshold.

b. Payment based on median costs should be adequate for hospitals since these products are generally older or low-cost items.

8. Payment for New Drugs, Biologicals and Radiopharmaceuticals Before HCPCS Codes Are Assigned.

a. The following payment methodology will enable hospitals to begin billing for drugs and biologicals that are newly approved by the FDA and for which a HCPCS code has not yet been assigned by the National HCPCS Alpha-Numeric Workgroup that could qualify them for pass-through payment under the OPPS:

(1) Hospitals should be instructed to bill for a drug or biological that is newly approved by the FDA by reporting the National Drug Code (NDC) for the product along with a new HCPCS code C9399, "Unclassified Drug or Biological."

(2) When HCPCS code C9399 appears on the claim, the OCE suspends the claim for manual pricing by the contractor.

(3) The new drug, biological and/or radiopharmaceutical will be priced at 95% of its AWP from a schedule of allowable charges based on the AWP, and process the claim for payment.

(4) The above approach enables hospitals to bill and receive payment for a new drug, biological or radiopharmaceutical concurrent with its approval by the FDA.

b. Hospitals will discontinue billing C9399 and the NDC upon implementation of a HCPCS code, SI, and appropriate payment amount with the next quarterly OPPS update.

9. Package payment for any biological without pass-through status that is surgically inserted or implanted (through a surgical incision or a natural orifice) into the payment for the associated surgical procedure.

a. As a result, HCPCS codes C9352, C9353, and J7348 are packaged and assigned SI of N.

b. Any new biologicals without pass-through status that are surgically inserted or implanted will be packaged beginning in CY 2009.

10. Drugs and non-implantable biologicals with expiring pass-through status.

a. CY 2009 payment methodology of packaged or separate payment based on their estimated per day costs, in comparison with the CY 2009 drug packaging threshold.

b. Packaged drugs and biologicals are assigned SI of N and drugs and biologicals that continue to be separately paid as non-pass-through products are assigned SI of K.

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E. Drug Administration Coding and Payment.

1. The following HCPCS Level I drug administration codes will be assigned to their respective APCs for payment:

**FIGURE 13-3-7 CROSSWALK FROM HCPCS LEVEL I<sup>1</sup> CODES FOR DRUG ADMINISTRATION TO DRUG ADMINISTRATION APCs**

HCPCS LEVEL I <sup>1</sup> CODE	DESCRIPTION	SI	APC
90769	Subcutaneous infusion for therapy or prophylaxis (specify substance or drug); initial, up to 1 hour, including pump set-up and establishment of subcutaneous infusion pump	S	0440
90770	Subcutaneous infusion for therapy or prophylaxis (specify substance or drug); each additional hour (List separately in addition to code for primary procedure)	S	0437
90771	Subcutaneous infusion for therapy or prophylaxis (specify substance or drug); additional pump set-up with establishment of new subcutaneous infusion site(s) (List separately in addition to code for primary procedure)	S	0438
90772	Therapeutic, prophylactic or diagnostic injection (specify substance or drug); subcutaneous or intramuscular	S	0437
90773	Therapeutic, prophylactic or diagnostic injection (specify substance or drug); intra-arterial	S	0438
90776	Therapeutic, prophylactic or diagnostic injection (specify substance or drug); each additional sequential push of the same substance/drug provided in a facility (List separately in addition to code for primary procedure)	N	
90779	Unlisted therapeutic, prophylactic or diagnostic intravenous or intra-arterial injection or infusion	S	0436
96401	Chemotherapy administration, subcutaneous or intramuscular; non-hormonal anti-neoplastic	S	0438
96402	Chemotherapy administration subcutaneous or intramuscular; hormonal anti-neoplastic	S	0438
96405	Chemotherapy administration; intralesional, up to and including 7 lesions	S	0438
96406	Chemotherapy administration; intralesional, more than 7 lesions	S	0438
96416	Chemotherapy administration, intravenous infusion technique; initiation of prolonged chemotherapy infusion (more than 8 hours), requiring use of portable or implantable pump	S	0441
96420	Chemotherapy administration, intra-arterial; push technique	S	0439
96422	Chemotherapy administration, intra-arterial; infusion technique, up to one hour	S	0441
96423	Chemotherapy administration, intra-arterial; infusion technique, each additional hour up to 8 hours (List separately in addition to code for primary procedure)	S	0438
96425	Chemotherapy administration, intra-arterial; infusion technique, initiation of prolonged infusion (more than 8 hours), requiring the use of a portable or implantable pump	S	0441
96440	Chemotherapy administration into pleural cavity, requiring and including thoracentesis	S	0441
96445	Chemotherapy administration into peritoneal cavity, requiring and including peritoneocentesis	S	0441

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**FIGURE 13-3-7 CROSSWALK FROM HCPCS LEVEL I<sup>1</sup> CODES FOR DRUG ADMINISTRATION TO DRUG ADMINISTRATION APCs (CONTINUED)**

HCPCS LEVEL I <sup>1</sup> CODE	DESCRIPTION	SI	APC
96450	Chemotherapy administration, into CNS (e.g., intrathecal), requiring and including spinal puncture	S	0441
96521	Refilling and maintenance of portable pump	S	0440
96522	Refilling and maintenance of implantable pump or reservoir for drug delivery, systemic (e.g., intravenous, intra-arterial)	S	0440
96523	Irrigation of implanted venous access device for drug delivery systems	Q	0624
96542	Chemotherapy injection, subarachnoid or intraventricular via subcutaneous reservoir, single or multiple agents	S	0438
96549	Unlisted chemotherapy procedure	S	0436

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2. The following non-chemotherapy HCPCS codes have also been created that are similar to CPT codes for initiation of prolonged chemotherapy infusion requiring a pump and pump maintenance and refilling codes so hospitals can bill for services when provided to patients who require extended infusions for non-chemotherapy medications including drugs for pain (see [Figure 13-3-8](#)).

**FIGURE 13-3-8 NON-CHEMOTHERAPY PROLONGED INFUSION CODES THAT REQUIRE A PUMP**

HCPCS LEVEL I <sup>1</sup> CODE	DESCRIPTION	SI	APC
C8957	Intravenous infusion for therapy /diagnosis; initiation of prolonged infusion (more than 8 hours), requiring use of portable or implantable pump	S	0441

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3. Packaged HCPCS Level I codes for drug administration should continue to be billed to ensure accurate payment in the future. These are bill changes for HCPCS Level I codes with SI of N that will be used as the basis for setting median costs for each drug administration HCPCS Level I code in the future.

4. HCPCS Level I<sup>4</sup> codes 90772-90774 each represent an injection and as such, one unit of the code may be billed each time there is a separate injection that meets the definition of the code.

5. Drugs for which the median cost per day is greater than \$60 are paid separately and are not packaged into the payment for the drug administration. Separate payment for drugs with a median cost in excess of \$60 will result in more equitable payment for both the drugs and their administration.

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F. Coding and Payment Policies for Drugs and Supplies.

1. Drug Coding.

a. Drugs for which separate payment is allowed are designated by SI K and must be reported using the appropriate HCPCS code.

b. Drugs that are reported without a HCPCS code will be packaged under the revenue center code, under OPPS: 250, 251, 252, 254, 255, 257, 258, 259, 631, 632, or 633.

c. Drugs billed using revenue code 636 (“Drugs requiring detailed coding”) require use of the appropriate HCPCS code, or they will be denied.

d. Reporting charges of packaged drugs is critical because packaged drug costs are used for calculating outlier payments and hospital costs for the procedure and service with which the drugs are used in the course of the annual OPPS updates.

2. Payment for the Unused Portion of a Drug.

a. Once a drug is reconstituted in the hospital’s pharmacy, it may have a limited shelf life. Since an individual patient may receive less than the fully reconstituted amount, hospitals are encouraged to schedule patients in such a way that the hospital can use the drug most efficiently. However, if the hospital must discard the remainder of a vial after administering part of it to a TRICARE patient, the provider may bill for the amount of the drug discarded, along with the amount administered.

b. In the event that a drug is ordered and reconstituted by the hospital’s pharmacy, but not administered to the patient, payment will be made under OPPS.

EXAMPLE 1: Drug X is available only in a 100-unit size. A hospital schedules three patients to receive drug X on the same day within the designated shelf life of the product. An appropriate hospital staff member administers 30 units to each patient. The remaining 10 units are billed to OPPS on the account of the last patient. Therefore, 30 units are billed on behalf of the first patient seen, and 30 units are billed on behalf of the second patient seen. Forty units are billed on behalf of the last patient seen because the hospital had to discard 10 units at that point.

EXAMPLE 2: An appropriate hospital staff member must administer 30 units of drug X to a patient, and it is not practical to schedule another patient for the same drug. For example, the hospital has only one patient who requires drug X, or the hospital sees the patient for the first time and does not know the patient’s condition. The hospital bills for 100 units on behalf of the patient, and OPPS pays for 100 units.

c. Coding for Supplies.

(1) Supplies that are an integral component of a procedure or treatment are not reported with a HCPCS code.

(2) Charges for such supplies are typically reflected either in the charges on the line for the HCPCS for the procedure, or on another line with a revenue code that will result in the charges being assigned to the same cost center to which the cost of those services are assigned in the cost report.

(3) Hospitals should report drugs that are treated as supplies because they are an integral part of a procedure or treatment under the revenue code associated with the cost center under which the hospital accumulates the costs for the drugs.

3. Recognition of Multiple HCPCS Codes for Drugs.

a. Prior to January 1, 2008, the OPPS generally recognized only the lowest available administrative dose of a drug if multiple HCPCS codes existed for the drug; for the remainder of the doses, the OPPS assigned a SI **B** indicating that another code existed for OPPS purposes. For example, if drug X has two HCPCS codes, one for a 1 ml dose and another for a 5 ml dose, the OPPS would assign a payable status indicator to the 1 ml dose and SI **B** to the 5 ml dose.

b. Hospitals then were required to bill the appropriate number of units for the 1 ml dose in order to receive payment under OPPS.

c. Beginning January 1, 2008, the OPPS has recognized each HCPCS code for a Part B drug, regardless of the units identified in the drug descriptor.

d. Hospitals may choose to report multiple HCPCS codes for a single drug, or to continue billing the HCPCS code with the lowest dosage descriptor available.

4. Correct Reporting of Drugs and Biologicals When Used As Implantable Devices.

a. When billing for biologicals where the HCPCS code describes a product that is solely surgically implanted or inserted, whether the HCPCS code is identified as having pass-through status or not, hospitals are to report the appropriated HCPCS code for the product.

b. Separate payment will be made for an implanted biological when it has pass-through status.

c. If the implantable device does not have pass-through status it will be packaged into the payment for the associated procedure.

5. Correct Reporting of Units for Drugs.

a. Units of drugs administered to patients should be 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.

b. For example, if the description for the drug code is 6 mg, and 6 mg of the drug was administered to the patients, the units bill should be one. If the description for the drug code is 50 mg, but 200 mg of the drug was administered, the units billed should be four.

c. Hospitals should not bill the units based on the way 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, bill 10 units even though only one vial was administered.

G. Orphan Drugs.

1. Continue to use the following criteria for identifying single indication orphan drugs that are used solely for orphan conditions:

a. The drug is designated as an orphan drug by the FDA and approved by the FDA for treatment of only one or more orphan condition(s).

b. The current United States Pharmacopoeia Drug Information (USPDI) shows that the drug has neither an approved use nor an off-label use for other than the orphan condition(s).

2. Twelve single indication orphan drugs have currently been identified as having met these criteria.

3. Payment Methodology.

a. Pay all 12 single indication orphan drugs at the rate of 88% of AWP or 106 of the ASP, whichever is higher.

b. However, for drugs where 106% of ASP would exceed 95% of AWP, payment would be capped at 95% of AWP, which is the upper limit allowed for sole source specified covered outpatient drugs.

H. Vaccines.

1. Hospitals will be paid for influenza, pneumococcal pneumonia and hepatitis B vaccines based on allowable charge methodology; i.e., will be paid the CMAC rate for these vaccines.

2. Separately payable vaccines other than influenza, pneumococcal pneumonia and hepatitis B will be paid under their own APC.

3. See [Figure 13-3-9](#) for vaccine administration codes and SIs.

**FIGURE 13-3-9 VACCINE ADMINISTRATION CODES AND STATUS INDICATORS**

HCPCS LEVEL 1 <sup>1</sup> CODE	DESCRIPTION	SI	APC
G0008	Influenza vaccine administration	S	0350
G0009	Pneumococcal vaccine administration	S	0350
G0010	Hepatitis B vaccine administration	B	--
90465	Immunization admin, under 8 yrs old, with counseling; first injection	N	--
90466	Immunization admin, under 8 yrs old, with counseling; each additional injection	N	--
90467	Immunization admin, under 8 yrs old, with counseling; first intranasal or oral	N	--
90468	Immunization admin, under 8 yrs old, with counseling; each additional intranasal or oral	N	--
90471	Immunization admin, one vaccine injection	S	0437
90472	Immunization admin, each additional vaccine injections	S	0436
90473	Immunization admin, one vaccine by intranasal or oral	N	
90474	Immunization admin, each additional vaccine by intranasal or oral	N	--

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**I. Payment Policy for Radiopharmaceuticals.**

Separately paid radiopharmaceuticals are classified as “specified covered outpatient drugs” subject to the following packaging and payment provisions:

1. The threshold for the establishment of separate APCs for radiopharmaceuticals is \$60.
2. A radiopharmaceutical that is covered and furnished as part of covered outpatient department services for which a HCPCS code has not been assigned will be reimbursed an amount equal to 95% of its AWP.
3. Radiopharmaceuticals will be excluded from receiving outlier payments.
4. Applications will be accepted for pass-through status; however, in the event the manufacturer seeking pass-through status for a radiopharmaceutical does not submit data in accordance with the requirements specified for new drugs and biologicals, payment will be set for the new radiopharmaceutical as a “specified covered outpatient drug.”

**J. Blood and Blood Products.**

1. Since the OPDS was first implemented, separate payment has been made for blood and blood products in APCs rather than packaging them into payment for the procedures with which they were administered. The APCs for these products are intended to recover the costs of the products. SI **R** was created in CY 2009 to denote blood and blood products.

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2. The OPPS provider also should report charges for processing and storage services on a separate line using Revenue Code 0390 (General Classification), 0392 (Blood Processing/Storage), or 0399 (Blood Processing/Storage; Other Blood Storage and Processing), along with appropriate blood HCPCS code, the number of units transfused, and the Line Item Date Of Service (LIDOS).

3. Administrative costs for the processing and storage specific to the transfused blood product are included in the APC payment, which is based on hospitals' charges.

4. Payment for the collection, processing, and storage of autologous blood, as described by HCPCS Level I<sup>5</sup> code 86890 and used in transfusion, is made through APC 347 (Level III Transfusion Laboratory Procedures).

5. Payment rates for blood and blood products will be determined based on median costs. Refer to [Figure 13-3-10](#) for APC assignment of blood and blood product codes.

**FIGURE 13-3-10 ASSIGNMENT OF BLOOD AND BLOOD PRODUCT CODES**

HCPCS	EXPIRED HCPCS	STATUS INDICATOR	DESCRIPTION	APC
P9010		R	Whole blood for transfusion	0950
P9011		R	Split unit of blood	0967
P9012		R	Cryoprecipitate each unit	0952
P9016		R	RBC leukocytes reduced	0954
P9017		R	Plasma 1 donor frz w/in 8 hr	9508
P9019		R	Platelets, each unit	0957
P9020		R	Platelet rich plasma unit	0958
P9021		R	Red blood cells unit	0959
P9022		R	Washed red blood cells unit	0960
P9023		R	Frozen plasma, pooled, sd	0949
P9031		R	Platelets leukocytes reduced	1013
P9032		R	Platelets, irradiated	9500
P9033		R	Platelets leukoreduced irradiated	0968
P9034		R	Platelets, pheresis	9507
P9035		R	Platelets pheresis leukoreduced	9501
P9036		R	Platelet pheresis irradiated	9502
P9037		R	Platelet pheresis leukoreduced irradiated	1019
P9038		R	RBC irradiated	9505
P9039		R	RBC deglycerolized	9504
P9040		R	RBC leukoreduced irradiated	0969
P9043		R	Plasma protein fract, 5%, 50 ml	0956
P9044		R	Cryoprecipitate reduced plasma	1009
P9048		R	Granulocytes, pheresis unit	9506

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**FIGURE 13-3-10 ASSIGNMENT OF BLOOD AND BLOOD PRODUCT CODES (CONTINUED)**

HCPCS	EXPIRED HCPCS	STATUS INDICATOR	DESCRIPTION	APC
P9051	C1010	R	Blood, L/R, CMV-NEG	1010
P9052	C1011	R	Platelets, HLA-m, L/R, unit	1011
P9053	C1015	R	Plt, pher, L/R, CMV, irradiated	1020
P9054	C1016	R	Blood, L/R, Froz/Degly/Washed	1016
P9055	C1017	R	Plt, Aph/Pher, L/R, CMV-Neg	1017
P9056	C1018	R	Blood, L/R, Irradiated	1018
P9057	C1020	R	RBC, frz/deg/wash, L/R irradiated	1021
P9058	C1021	R	RBC, L/R, CMV-Neg, irradiated	1022
P9059	C1022	R	Plasma, frz within 24 hours	0955
P9060	C9503	R	Fresh frozen plasma, ea unit	9503

6. For CY 2009, blood clotting factors will be paid at ASP + 4%, plus an additional payment for the furnishing fee that is also a part of the payment for blood clotting factors furnished in physician's offices.

**K. Adjustment to Payment in Cases of Devices Replaced with Partial Credit for the Replaced Device.**

1. Hospitals will be required to append the modifier "FC" to the HCPCS code for the procedure in which the device was inserted on claims when the device that was replaced with partial credit under warranty, recall, or field action is one of the devices in [Figure 13-3-11](#). Hospitals should not append the modifier to the HCPCS procedure code if the device is not listed in [Figure 13-3-11](#).

2. Claims containing the "FC" modifier will not be accepted unless the modifier is on a procedure code with SI **S**, **T**, **V**, or **X**.

3. If the APC to which the procedure is assigned is one of the APCs listed in [Figure 13-3-12](#), the Pricer will reduce the unadjusted payment rate for the procedure by an amount equal to the percent in [Figure 13-3-12](#) for partial credit device replacement (i.e., 50% of the device offset when both a device code listed in [Figure 13-3-11](#) is present on the claim and the procedure code maps to an APC listed in [Figure 13-3-12](#)) multiplied by the unadjusted payment rate.

4. The partial credit adjustment will occur before wage adjustment and before the assessment to determine if the reductions for multiple procedures (signified by the presence of more than one procedure on the claim with a SI of **T**), discontinued service (signified by modifier 73) or reduced service (signified by modifier 52) apply.

**L. Payment When Devices Are Replaced Without Cost or Where Credit for a Replacement Device is Furnished to the Hospital.**

1. Payments will be reduced for selected APCs in cases in which an implanted device is replaced without cost to the hospital or with full credit for the removed device. The

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amount of the reduction to the APC rate will be calculated in the same manner as the offset amount that would be applied if the implanted device assigned to the APC has pass-through status.

2. This permits equitable adjustments to the OPPS payments contingent on meeting all of the following criteria:

- a. All procedures assigned to the selected APCs must require implantable devices that would be reported if device replacement procedures are performed;
- b. The required devices must be surgically inserted or implanted devices that remain in the patient's body after the conclusion of the procedures, at least temporarily; and
- c. The offset percent for the APC (i.e., the median cost of the APC without device costs divided by the median cost of the APC with device costs) must be significant-- significant offset percent is defined as exceeding 40%.

3. The presence of the modifier "FB" ["Item Provided Without Cost to Provider, Supplier, or Practitioner or Credit Received for Replacement (examples include, but are not limited to devices covered under warranty, replaced due to defect, or provided as free samples)"] would trigger the adjustment in payment if the procedure code to which modifier "FB" was amended appeared in [Figure 13-3-11](#) and was also assigned to one of the APCs listed in [Figure 13-3-12](#). OPPS payments for implantation procedures to which the "FB" modifier is appended are reduced to 100% of the device offset for no-cost/full credit cases.

**FIGURE 13-3-11 DEVICES FOR WHICH "FC" AND "FB" MODIFIERS MUST BE REPORTED WITH THE PROCEDURE WHEN FURNISHED WITHOUT COST OR AT FULL OR PARTIAL CREDIT FOR A REPLACEMENT DEVICE**

DEVICE HCPCS CODE	SHORT DESCRIPTOR
C1721	AICD, dual chamber
C1722	AICS, single chamber
C1728	Cath, brachytx seed adm
C1764	Event recorder, cardiac
C1767	Generator, neurostim, imp
C1771	Rep Dev urinary, w/sling
C1772	Infusion pump, programmable
C1776	Joint device (implantable)
C1777	Lead, AICD, endo single coil
C1778	Lead neurostimulator
C1779	Lead, pmkr, transvenous VDD
C1785	Pmkr, dual rate-resp
C1786	Pmkr, single rate-resp
C1789	Prosthesis, breast, imp
C1813	Prostheses, penile, inflatab
C1815	Pros, urinary sph, imp

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**FIGURE 13-3-11 DEVICES FOR WHICH “FC” AND “FB” MODIFIERS MUST BE REPORTED WITH THE PROCEDURE WHEN FURNISHED WITHOUT COST OR AT FULL OR PARTIAL CREDIT FOR A REPLACEMENT DEVICE (CONTINUED)**

DEVICE HCPCS CODE	SHORT DESCRIPTOR
C1820	Generator, neuro, rechg bat sys
C1882	AICD, other than sing/dual
C1891	Infusion pump, non-prog, perm
C1895	Lead, AICD, endo dual coil
C1896	Lead, AICD, non sing/dual
C1897	Lead, neurostim, test kit
C1898	Lead, pmkr, other than trans
C1899	Lead, pmkr/AICD combination
C1900	Lead coronary venous
C2619	Pmkr, dual, non rate-resp
C2620	Pmkr, single, non rate-resp
C2621	Pmkr, other than sing/dual
C2622	Pmkr, other than sing/dual
C2626	Infusion pump, non-prog, temp
C2631	Rep dev, urinary, w/o sling
L8600	Implant breast silicone/eq
L8614	Cochlear device/system
L8685	Implt nrostm pls gen sng rec
L8686	Implt nrostm pls gen sng non
L8687	Implt nrostm pls gen dua rec
L8688	Implt nrostm pls gen dua non
L8690	Aud osseo dev, int/ext comp

**FIGURE 13-3-12 ADJUSTMENTS TO APCs IN CASES OF DEVICES REPORTED WITHOUT COST OR FOR WHICH FULL OR PARTIAL CREDIT IS RECEIVED FOR CY 2009**

APC	SI	APC GROUP TITLE	DEVICE OFFSET PERCENTAGE FOR NO-COST/FULL CREDIT CASE	DEVICE OFFSET PERCENTAGE FOR PARTIAL CREDIT CASE
0039	S	Level I Implantation of Neurostimulator	84	42
0040	S	Percutaneous Implantation of Neurostimulator Electrodes, Excluding Cranial Nerve	57	29
0061	S	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electrodes	62	31
0089	T	Insertion/Replacement of Permanent Pacemaker and Electrodes	72	36
0090	T	Insertion/Replacement of Pacemaker Pulse Generator	74	37
0106	T	Insertion/Replacement of Pacemaker Leads and/or Electrodes	43	21
0107	T	Insertion of Cardioverter-Defibrillator	89	45
0108	T	Insertion/Replacement/Repair of Cardioverter-Defibrillator Leads	89	44
0222	S	Level II Implantation of Neurological Device	85	42

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**FIGURE 13-3-12 ADJUSTMENTS TO APCs IN CASES OF DEVICES REPORTED WITHOUT COST OR FOR WHICH FULL OR PARTIAL CREDIT IS RECEIVED FOR CY 2009 (CONTINUED)**

APC	SI	APC GROUP TITLE	DEVICE OFFSET PERCENTAGE FOR NO-COST/FULL CREDIT CASE	DEVICE OFFSET PERCENTAGE FOR PARTIAL CREDIT CASE
0225	S	Implantation of Neurostimulator Electrodes, Cranial Nerve	62	31
0227	T	Implantation of Drug Infusion Devices	82	41
0229	T	Transcatheter Placement of Intravascular Shunts	84	42
0259	T	Level IV ENT Procedures	88	44
0315	S	Level III Implantation of Neurostimulator	59	29
0385	S	Level I Prosthetic Urological Procedures	69	34
0386	S	Level II Prosthetic Urological Procedures	71	36
0418	T	Insertion of Left Ventricular Pacing Elect	59	29
0425	T	Level II Arthroplasty or Implantation with Prosthesis	46	23
0648	T	Level IV Breast Surgery	77	38
0625	T	Level IV Vascular Access Procedures	76	38
0654	T	Insertion/Replacement of a Permanent Dual Chamber Pacemaker	71	36
0655	T	Insertion/Replacement/Conversion of a Permanent Dual Chamber Pacemaker	71	35
0680	S	Insertion of Patient Activated Event Recorders	71	36
0681	T	Knee Arthroplasty	71	35

4. If the APC to which the device code (i.e., one of the codes in [Figure 13-3-11](#)) is assigned is on the APCs listed in [Figure 13-3-12](#), the unadjusted payment rate for the procedure APC will be reduced by an amount equal to the percent in [Figure 13-3-12](#) times the unadjusted payment rate.

5. In cases in which the device is being replaced without cost, the hospital will report a token device charge. However, if the device is being inserted as an upgrade, the hospital will report the difference between its usual charge for the device being replaced and the credit for the replacement device.

6. Multiple procedure reductions would also continue to apply even after the APC payment adjustment to remove payment for the device cost, because there would still be the expected efficiencies in performing the procedure if it was provided in the same operative session as another surgical procedure. Similarly, if the procedure was interrupted before administration of anesthesia (i.e., there was modifier 52 or 73 on the same line as the procedure), a 50% reduction would be taken from the adjusted amount.

M. Policies Affecting Payment of New Technology Services.

1. A process was developed that recognizes new technologies that do not otherwise meet the definition of current orphan drugs, or current cancer therapy drugs and biologicals and brachytherapy, or current radiopharmaceutical drugs and biologicals products. This process, along with transitional pass-throughs, provides additional payment for a significant share of new technologies.

2. Special APC groups were created to accommodate payment for new technology services. In contrast to the other APC groups, the new technology APC groups did not take into account clinical aspects of the services they were to contain, but only their costs.

3. The SI **K** is used to denote the APCs for drugs, biologicals and pharmaceuticals that are paid separately from, and in addition to, the procedure or treatment with which they are associated, yet are not eligible for transitional pass-through payment.

4. New items and services will be assigned to these new technology APCs when it is determined that they cannot appropriately be placed into existing APC groups. The new technology APC groups provide a mechanism for initiating payment at an appropriate level within a relatively short time frame.

5. As in the case of items qualifying for the transitional pass-through payment, placement in a new technology APC will be temporary. After information is gained about actual hospital costs incurred to furnish a new technology service, it will be moved to a clinically-related APC group with comparable resource costs.

6. If a new technology service cannot be moved to an existing APC because it is dissimilar clinically and with respect to resource costs from all other APCs, a separate APC will be created for such services.

7. Movement from a new technology APC to a clinically-related APC will occur as part of the annual update of APC groups.

8. The new technology APC groups have established payment rates for the APC groups based on the midpoint of ranges of possible costs; for example, the payment amount for a new technology group reflecting a range of costs from \$300 to \$500 would be set at \$400. The cost range for the groups reflects current cost distributions, and TRICARE reserves the right to modify the ranges as it gains experience under the OPPTS.

9. There are two parallel series of technology APCs covering a range of costs from less than \$50 to \$6,000.

a. The two parallel sets of technology APCs are used to distinguish between those new technology services designated with a SI of **S** and those designated as **T**. These APCs allow assignment to the same APC group procedures that are appropriately subject to a multiple procedure payment reduction (**T**) with those that should not be discounted (**S**).

b. Each set of technology APC groups have identical group titles and payment rates, but a different SI.

c. The new series of APC numbers allow for the narrowing of the cost bands and flexibility in creating additional bands as future needs may dictate. Following are the narrowed incremental cost bands for the two series of new technology APCs:

(1) From \$0 to \$50 in increments of \$10.

(2) From \$50 to \$100 in a single \$50 increment.

(3) From \$100 through \$2,000 in intervals of \$100.

(4) From \$2,000 through \$6,000 in intervals of \$500.

10. Beneficiary cost-sharing/copayment amounts for items and services in the new technology APC groups are dependent on the eligibility status of the beneficiary at the time the outpatient services were rendered (i.e., those deductibles and cost-sharing/copayment amounts applicable to Prime, Extra and Standard beneficiary categories). (Refer to [Chapter 2, Addendum A](#) for applicable deductible cost-sharing/copayment amounts for outpatient hospital services.)

11. Process and Criteria for Assignment to a New Technology APC Group.

a. Services Paid Under New Technology APCs.

(1) Limit eligibility for placement in new technology APCs to complete services and procedures.

(2) Items, material, supplies, apparatuses, instruments, implements, or equipment that are used to accomplish a more comprehensive service or procedure would not be eligible for placement in a new technology APC.

(3) A service that qualifies for a new technology APC may be a complete, stand-alone service (for example, water-induced thermotherapy of the prostate or cryosurgery of the prostate), or it may be a service that would always be billed in combination with other services (for example, coronary artery brachytherapy).

(a) In the latter case, the new technology procedure, even though billed in combination with other, previously existing procedures, describes a distinct procedure with a beginning, middle, and end.

(b) Drugs, supplies, devices, and equipment in and of themselves are not distinct procedures with a beginning, middle and end. Rather drugs, supplies, devices, and equipment are used in the performance of a procedure.

(4) Unbundled components that are integral to a service or procedure (for example, preparing a patient for surgery or preparation and application of a wound dressing for wound care) are not eligible for consideration for a new technology.

b. Criteria for determining whether a service will be assigned to a new technology APC.

(1) The most important criterion in determining whether a technology is “truly new” and appropriate for a new APC is the inability to appropriately, and without redundancy, describe the new, complete (or comprehensive) service with any combination of existing HCPCS Level I and II codes. In other words, a “truly new” service is one that cannot be appropriately described by existing HCPCS codes, and a new HCPCS code needs to be established in order to describe the new procedure.

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(2) The service is one that could not have been adequately represented in the claims data being used for the most current annual payment update; i.e., the item is one service that could not have been billed to the Medicare program in 1996 or, if it was available in 1996, the costs of the service could not have been adequately represented in 1996 data.

(3) The service does not qualify for an additional payment under the transitional pass-through provisions.

(4) The service cannot reasonably be placed in an existing APC group that is appropriate in terms of clinical characteristics and resource costs. It is unnecessary to assign a new service to a new technology APC if it may be appropriately placed in a current APC.

(5) The service falls within the scope of TRICARE benefits.

(6) The service is determined to be reasonable and necessary.

NOTE: The criterion that the service must have a HCPCS code in order to be assigned to a new technology APC has been removed. This is supported by the rationale that in order to be considered for a new technology APC, a truly new service cannot be adequately described by existing codes. Therefore, in the absence of an appropriate HCPCS code, a new HCPCS code will be created that describes the new technology service. The new HCPCS would be solely for hospitals to use when billing under the OPPS.

#### N. Coding And Payment Of ED Visits.

1. CPT defines an ED as “an organized hospital based facility for the provision of unscheduled episodic services to patients who present for immediate medical attention. The facility must available 24 hours a day.”

2. Prior to CY 2007, under the OPSS the billing of ED CPT codes was restricted to services furnished at facilities that met this CPT definition. Based on the above definition, facilities open less than 24 hours a day could not report ED CPT codes.

3. Sections 1866(a)(1)(I), 1866(a)(1)(N), and 1867 of the Act impose specific obligations on Medicare-participating hospitals that offer emergency services. These obligations concern individuals who come to a hospital’s Dedicated Emergency Department (DED) and request examination or treatment for medical conditions, and apply to all of these individuals, regardless of whether or not they are beneficiaries of any program under the Act. Section 1867(h) of the Act specifically prohibits a delay in providing required screening or stabilization services in order to inquire about the individual’s payment method or insurance status.

4. These provisions are frequently referred to as the Emergency Medical Treatment and Labor Act (EMTALA). The EMTALA regulations define DED as any department or facility of the hospital, regardless of whether it is located on or off the main campus, that meets at least one of the following requirements:

o. It is licensed by the State in which it is located under applicable State law as an **Emergency Room (ER)** or ED;

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b. It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or

c. During the calendar year immediately preceding the calendar year in which a determination under the regulations is being made, based on a representative sample of patient visits that occurred during the calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring previously scheduled appointment.

5. There are some departments or facilities of hospitals that met the definition of a dedicated ED under the EMTALA regulations, but did not meet the more restrictive CPT definition of ED. For example, a hospital department or facility that met the definition of a DED might not have been available 24 hours a day, seven days a week.

6. To determine whether visits to EDs of facilities (referred to as Type B ED) that incur EMTALA obligations, but do not meet the more prescriptive expectations that are consistent with the CPT definition of an ED (referred to as Type A ED) have different resource costs than visits to either clinics or Type A EDs, five G codes were developed for use by hospitals to report visits to all entities that meet the definition of a DED under the EMTALA regulations, but that are not Type A EDs. These codes are called "Type B ED visit codes." EDs meeting the definition of a DED under the EMTALA regulations, but which are not Type A EDs (i.e., they may meet the DED definition but are not available 24 hours a day, seven days a week).

**FIGURE 13-3-13 FINAL HCPCS CODES TO BE USED TO REPORT ED VISITS PROVIDED IN TYPE B EDs**

HCPCS CODE	SHORT DESCRIPTOR	LONG DESCRIPTOR
G0380	Level 1 Hosp Type B Visit	Level 1 hospital ED visit provided in a Type B ED. (The ED must meet at least one of the following requirements: (1) It is licensed by the State in which it is located under applicable State law as an ER or ED; (2) It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or (3) During the calendar year immediately preceding the calendar year in which a determination under this section is being made, based on a representative sample of patient visits that occurred during that calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring a previously scheduled appointment.)
G0381	Level 2 Hosp Type B Visit	Level 2 hospital ED visit provided in a Type B ED. (The ED must meet at least one of the following requirements: (1) It is licensed by the State in which it is located under applicable State law as an ER or ED; (2) It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or (3) During the calendar year immediately preceding the calendar year in which a determination under this section is being made, based on a representative sample of patient visits that occurred during that calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring a previously scheduled appointment.)

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**FIGURE 13-3-13 FINAL HCPCS CODES TO BE USED TO REPORT ED VISITS PROVIDED IN TYPE B EDs**

HCPCS CODE	SHORT DESCRIPTOR	LONG DESCRIPTOR
G0382	Level 3 Hosp Type B Visit	Level 3 hospital ED visit provided in a Type B ED. (The ED must meet at least one of the following requirements: (1) It is licensed by the State in which it is located under applicable State law as an ER or ED; (2) It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or (3) During the calendar year immediately preceding the calendar year in which a determination under this section is being made, based on a representative sample of patient visits that occurred during that calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring a previously scheduled appointment.)
G0383	Level 4 Hosp Type B Visit	Level 4 hospital ED visit provided in a Type B ED. (The ED must meet at least one of the following requirements: (1) It is licensed by the State in which it is located under applicable State law as an ER or ED; (2) It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or (3) During the calendar year immediately preceding the calendar year in which a determination under this section is being made, based on a representative sample of patient visits that occurred during that calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring a previously scheduled appointment.)
G0384	Level 5 Hosp Type B Visit	Level 5 hospital ED visit provided in a Type B ED. (The ED must meet at least one of the following requirements: (1) It is licensed by the State in which it is located under applicable State law as an ER or ED; (2) It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or (3) During the calendar year immediately preceding the calendar year in which a determination under this section is being made, based on a representative sample of patient visits that occurred during that calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring a previously scheduled appointment.)

7. The five new Type B ED visit codes for services provided in a Type B ED will be assigned to the five newly established Clinical Visit APCs 0604, 0605, 0606, 0607, and 0608.

8. For CY 2007, the five CPT E/M ED visit codes for services provided in a Type A ED were assigned to the five newly-created ED Visit APCs 0609, 0613, 0614, 0615, and 0616.

9. The definition of Type A and Type B EDs was not modified for CY 2008 because its current definition accurately distinguished between these two types of ED.

10. For CY 2008, Type A ED visits will continue to be paid based on the five ED Visit APCs, while Type B ED visits would continue to be paid based on the five Clinic Visit APCs.

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11. A new G code (G0390 - Trauma response team activation associated with hospital critical care services) was also created (effective January 1, 2007) to be used in addition to CPT<sup>6</sup> procedure codes 99291 and 99292 to address the meaningful cost difference between critical care when billed with and without trauma activation.

a. If critical care is provided without trauma activation, the hospital will bill with either CPT<sup>6</sup> procedure code 99291, receiving payment for APC 0617.

b. However if trauma activation occurs, the hospital would be called to bill one unit of "G" code (G0390), report with revenue code 68x on the same date of service, thereby receiving payment for APC 0618.

12. The CPT Evaluation and Management (E/M) codes and other HCPCS codes currently assigned to the clinic visit APCs have been mapped in [Figure 13-3-14](#) to 11 new APCs; five for clinic visits; five for ED visits; and one for critical care services, based on median costs and clinical consideration.

**FIGURE 13-3-14 ASSIGNMENT OF CPT E/M CODES AND OTHER HCPCS CODES TO NEW VISIT APCs FOR CY 2007**

APC TITLE	APC	HCPCS	SHORT DESCRIPTOR
Level 1 Hospital Clinic Visits	0604	92012	Eye exam, established pat
		99201	Office/outpatient visit, new (Level 1)
		99211	Office/outpatient visit, est (Level 1)
		99241	Office consultation
		G0101	CA screen; pelvic/breast exam
		G0245	Initial foot exam Pt lops
		G0379	Direct admit hospital observ
Level 2 Hospital Clinic Visits	0605	92002	Eye exam, new patient
		92014	Eye exam and treatment
		99202	Office/outpatient visit, new (Level 2)
		99212	Office/outpatient visit, est (Level 2)
		99213	Office/outpatient visit, est (Level 3)
		99243	Office consultation (Level 3)
		99242	Office consultation (Level 2)
		99273	Confirmatory consultation (Level 3)
		99272	Confirmatory consultation (Level 2)
		99431	Initial care, normal newborn
		G0246	Follow-up eval of foot pt lop
		G0344	Initial preventive exam
Level 3 Hospital Clinic Visits	0606	90862	Medication management
		92004	Eye exam, new patient
		99203	Office/outpatient visit, new (Level 3)
		99214	Office/outpatient visit, est (Level 4)

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**FIGURE 13-3-14 ASSIGNMENT OF CPT E/M CODES AND OTHER HCPCS CODES TO NEW VISIT APCs FOR CY 2007 (CONTINUED)**

APC TITLE	APC	HCPCS	SHORT DESCRIPTOR
Level 3 Hospital Clinic Visits (Continued)	0606	99274	Confirmatory consultation (Level 4)
		99244	Office consultation (Level 4)
		M0064	Visit for drug monitoring
Level 4 Hospital Clinic Visits	0607	99204	Confirmatory consultation (Level 1)
		99215	Office/outpatient visit, est (Level 5)
		99245	Office consultation (Level 5)
		99275	Confirmatory consultation (Level 5)
Level 5 Hospital Clinic Visits	0608	99205	Office/outpatient visit, new (Level 5)
		G0175	OPPS service, sched team conf
Level 1 Type A Emergency Visits	0609	99281	Emergency department visit
Level 2 Type A Emergency Visits	0613	99282	Emergency department visit
Level 3 Type A Emergency Visits	0614	99283	Emergency department visit
Level 4 Type A Emergency Visits	0615	99284	Emergency department visit
Level 5 Type A Emergency Visits	0616	99285	Emergency department visit
Critical Care	0617	99291	Critical care, first hour
Trauma Activation	0618	G0390	Trauma Respon. w/hosp criti

O. OPSS PRICER.

1. Common PRICER software will be provided to the contractor that includes the following data sources:

- a. National APC amounts
- b. Payment status by HCPCS code
- c. Multiple surgical procedure discounts
- d. Fixed dollar threshold
- e. Multiplier threshold
- f. Device offsets
- g. Other payment systems pricing files (CMAC, DMEPOS, and statewide prevalings)

2. The following data elements will be extracted and forwarded to the outpatient PRICER for line item pricing.

- a. Units;
- b. HCPCS/Modifiers;

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- c. APC;
  - d. Status payment indicator;
  - e. Line item date of service;
  - f. Primary diagnosis code; and
  - g. Other necessary OCE output.
3. The following data elements will be passed into the PRICER by the contractors:
- a. Wage indexes (same as DRG wage indexes);
  - b. Statewide CCRs as provided in the CMS Final Rule and listed on TMA's OPPS web site at <http://www.tricare.mil/opps>;
  - c. Locality Code: Based on CBSA - two digit = rural and five digit = urban;
  - d. Hospital Type: Rural SCH = 1 and All Others = 0
4. The outpatient PRICER will return the line item APC and cost outlier pricing information used in final payment calculation. This information will be reflected in the provider remittance notice and beneficiary EOB with exception for an electronic 835 transaction. Paper EOBs and remits will reflect APCs at the line level and will also include indication of outlier payments and pricing information for those services reimbursed under other than OPPS methodology's, e.g., CMAC (SI of A) when applicable.
5. If a claim has more than one service with a SI of T or a SI of S within the coding range of 10000 - 69999, and any lines with SI of T or a SI within the coding range of 10000 - 69999 have less than \$1.01 as charges, charges for all lines will be summed and the charges will then be divided up proportionately to the payment rates for each line (refer to [Figure 13-3-15](#)). The new charge amount will be used in place of the submitted charge amount in the line item outlier calculator.

**FIGURE 13-3-15 PROPORTIONAL PAYMENT FOR "T" LINE ITEMS**

SI	CHARGES	PAYMENT RATE	NEW CHARGES AMOUNT
T	\$19,999	\$6,000	\$12,000
T	\$1	\$3,000	\$6,000
T	\$0	\$1,000	\$2,000
Total	\$20,000	\$10,000	\$20,000

**NOTE:** Because total charges here are \$20,000 and the first SI of T gets \$6,000 of the \$10,000 total payment, the new charge for that line is  $\$6,000/\$10,000 \times \$20,000 = \$12,000$ .

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P. TRICARE Specific Procedures/Services.

1. TRICARE specific APCs have been assigned for half-day PHPs.
2. Other procedures that are normally covered under TRICARE but not under Medicare will be assigned SI of **A** (i.e., services that are paid under some payment method other than OPSS) until they can be placed into existing or new APC groups.

Q. Validation Reviews.

OPSS claims are not subject to validation review.

R. Hospital-Based Birthing Centers.

Hospital-based birthing centers will be reimbursed the same as freestanding birthing centers except the all inclusive rate consisting of the CMAC for CPT<sup>7</sup> procedure code 59400 and the state specific non-professional component, will lag two months (i.e., April 1 instead of February 1).

IV. EFFECTIVE DATE            May 1, 2009.

- END -

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