State Operations Manual

Chapter 3 - Additional Program Activities

Table of Contents

(Rev. 198, 01-17-19)

Transmittals for Chapter 3

Adverse Actions

3000 - Adverse Actions - General
   3000A - Applicability
   3000B - State Ombudsman Programs
   3000C - CMS Authority to Terminate Medicare and Medicaid Participation
3001 - Initial Denials of Medicare Provider/Supplier Requests for Program Participation
   3001A - Authority for Adjudicating Denials
   3001B - Vacated Actions Which Are Not Denials
   3001C - Vacated Actions Which Are Denials
   3001D - RO Processing of Denials
3005 - Basis for Terminating Provider Participation - Citations and Discussion
   3005A - Medicare Provider Agreements
   3005B - Termination of Coverage of Supplier Services Subject to Certification
   3005D - Cause for Termination
   3005E - Termination of Title XIX-Only NFs, ICFs/IID, Hospitals and Psychiatric Hospitals
   3005F - Termination Action Based Upon Onsite Survey by RO or Validation Survey of a Deemed Provider or Supplier by RO or SA
   3005G - Look Behind Authority of CMS
      3005G1 - “Look-Behind” Termination or Cancellation of ICF/IID Agreement by the Secretary
      3005G2 - Old “Look-Behind” Termination of a NF or ICF/IID by the Secretary
      3005G3 - SMA Disagrees With SA Determination
3006 - Denial of Payments in Lieu of Termination of ICFs/IID
   3006A - Authority to Deny Payment for Any New Admissions for ICFs/IID
   3006B - Criteria for Imposing Denial of Payments for New Admissions
   3006C - Agency Procedures
3006D - Effect of Sanction on Status of Clients Admitted, Discharged, or on Temporary Leave and Readmitted Before or After Effective Date of Denial of Payment

3006F - Duration of Denial of Payment and Subsequent Termination of an ICF/IID

3006.1 - Sanctions for ICFs-IID - or Nonimmediate Jeopardy
   3006.1A - General
   3006.1B - Introduction
   3006.1C - Examples of Alternative Sanctions
   3006.1D - Alternatives to Termination in Nonimmediate Jeopardy Situations

3006.2 - Directed Plan of Correction (DPoC)
   3006.2A - Purpose
   3006.2B - Basis for Imposition of a DPoC
   3006.2C - Elements of a DPoC
   3006.2D - Notice of Imposition of DPoC

3006.3 - Directed In-Service Training
   3006.3A - Purpose
   3006.3B - Appropriate Resources for Directed In-Service Training Programs
   3006.3C - Further Responsibilities
   3006.3D - Notice of Imposition of Directed In-Service Training

3006.4 - State Monitoring
   3006.4A - Purpose
   3006.4B - Qualifications
   3006.4C - When to Impose State Monitoring
   3006.4D - Frequency
   3006.4E - Duration
   3006.4F - Notice of Imposition of State Monitoring
   3006.4G - Payment for and Obligation of Funds by a State Monitor

3006.5 - Achieving Continuous, Substantial Compliance
   3006.5A - Introduction
   3006.5B - Duration of a Sanction
   3006.5C - Achieving and Maintaining Substantial Compliance

3006.6 - Criteria for Review of State Plans for Approval or Disapproval of Alternative Sanctions
   3006.6A - Introduction
   3006.6B - Alternative Remedies

3008 - Services After Termination
3008.1 - Services After Termination of a Medicare Provider Agreement
3008.2 - Services for which Federal Financial Participation (FFP) May Be Temporarily Continued After Termination of a Medicaid Provider Agreement or Nonrenewal or Cancellation of an ICF/IID Provider Agreement
3008.3 - Relocating Patients Displaced by Termination or Closure
   3008.3A - General
   3008.3B - Relocation of Medicaid Patients
   3008.3C - Relocation Activities

3010 - Termination Procedures - Immediate Jeopardy to Patient Health and Safety (Medicare)
   3010A - Substantial Noncompliance With Program Requirements Which Poses Immediate Jeopardy to Patient Health or Safety
   3010B - Processing of Immediate Jeopardy Terminations

3012 - Termination Procedures – Substantial Noncompliance; No Immediate Jeopardy (Medicare)
   3012.1 - Termination of Psychiatric Hospitals
   3012.2 - Termination of Organ Procurement Organizations (OPO)
     3012.2A - Termination Procedures
     3012.2B - Reconsideration Procedures
     3012.2C - Appeal Procedures
     3012.3 – Termination of Organ Transplant Programs

3014 - RO Termination Action Based on Onsite Survey of Medicare Provider or Supplier (Excluding SNFs) Conducted by RO Staff

3016 - Intervening Actions That Do Not Postpone or Delay Termination Timetable - (Includes Credible Allegations)
   3016A - Credible Allegation of Compliance
   3016B - Informal Hearings Do Not Interrupt Timetable
   3016C - Acceleration of Timetable
   3016D - Termination Development Coinciding With Change of Ownership (CHOW) Development
   3016E - Disagreement over Deficiencies
   3016.1 - Provider Undergoes Chow During Termination Proceedings

3018 - Termination - SA Documentation Requirements
   3018A - Documentation to Support Proposed Termination
     3018A1 - Current Survey Report
     3018A2 - Previous Survey Reports
   3018B - Record of Contacts With Providers/Suppliers
   3018C - Notification to Provider/Supplier of Deficiencies and Recommendation of Termination
3020 - Additional SA Communications With Providers/Suppliers
3022 - Notice of Termination (Medicare)
3024 - RO Termination Processing Sequence - Noncompliance With CoPs or Conditions for Coverage (Excluding SNFs)
3026 - Significance of Documentary Evidence in Determining Noncompliance
  3026A - Statement of Deficiencies
  3026B - Plan of Correction (PoC)
  3026C - Revisit Reports and Subsequent Statements of Deficiencies
  3026D - SA Certification (Completed Certification and Transmittal (Form CMS-1539))
  3026E - Survey Reports
  3026F - Documents of Collateral Evidence
  3026G - Notice of Termination
3028 - Documentation Guide List - Termination for Noncompliance With §§1866(b)(2)(A) and (C)
  3028A - Documentation Appropriate to All Cases
  3028B - Additional Documentation - Charging for Covered Services and/or Refusing to Refund Incorrect Collections
  3028C - Additional Documentation - Failure to File Cost Reports
  3028D - Additional Documentation - Failure to Make Satisfactory Overpayment Arrangements
  3028E - Additional Documentation - Admission Policies and Practices
3030 - Provider Agreement Terminations - Noncompliance with §§1866(b)(2)(A) and (C)
  3030A - Cause for Termination
  3030B - Preparing Termination Cases
  3030C - Preliminary Notice to Provider
  3030D - Violation of §§1866(b)(2)(A) and (C)
3032 - Termination for Violations of §§1866(a)(1)(E), (F), (G), and (H)
3034 - Public Notice - Involuntary Termination
  3034A - For Providers
  3034B - For Suppliers
  3034C - For Hospitals, CAHs, and SNFs
  3034D - For HHAs and Hospices
  3034E - For Other Providers and Suppliers
3036 - Billing for Public Notice of Termination or Withdrawal
  3036A - Advertising Order, SF-1143
3038 - Rescinding or Postponing Effective Date of Termination
  3038A - Initial Action
  3038B - Criteria for Credible Allegation
3038C - Form of Public Notice of Termination Retraction for Providers
3038D - Form of Public Notice of Termination Retraction for Suppliers
3040 - Terminating Medicaid ICF/IID Eligibility Based on “Look Behind” Determination
  3040A - Termination Procedures
    3040A1 - Immediate Jeopardy
    3040A2 - No Immediate Threat to Patients’ Health and Safety
3042 - Disallowance of FFP to State Because State Fails to Follow Correct Certification Procedures for Medicaid Providers
3044 - Terminating Approval for Suppliers
3046 - Voluntary Terminations
  3046A - General
    3046B - Decision by Provider or Supplier to Remain in the Medicare Program
  3046C - Notice to Public
  3046D - Effective Date of Voluntary Termination
3047 - Notice to Intermediary or Carrier - Voluntary Termination
3048 - Notice to Provider or Supplier - Voluntary Termination
  3048A - Voluntary Termination
  3048B - Close of Business
3049 - Completing Certification and Transmittal (Form CMS-1539)
  3049A - Voluntary Terminations and Close of Business

Reconsideration, Hearings, and Appeals
3050 - Initial Determinations Versus Administrative Actions - Right to Review
3052 - Nature of Reconsideration Determination - SA Procedures
  3052A - Right to Reconsideration of Initial Denial
  3052B - Request for Reconsideration
  3052C - Acknowledgment of Reconsideration Request
  3052D - Documentation of File
3054 - Reconsideration - RO Procedures – Excluding SNFs and NFs
  3054A - Review
  3054B - RO Receipt of Request
  3054C - Acknowledgment or Reconsideration Request
  3054D - Reconsideration Determination
  3054E - RO Notice of Reconsidered Determination
    3054E1 - Denial Reversal (Approval)
    3054E2 - Denial Affirmed
    3054E3 - Acting Official for Reconsideration Denial Notices
3058 - Hearing on §1910(b) Cancellation of Medicaid Eligibility
3060 - Appeals of Adverse Actions for Medicaid Non-State Operated NFs (Non-State Operated) and ICFs/IID (Not Applicable to Federal Terminations of Medicaid Facilities)
   3060A - Informal Reconsideration
   3060B - Evidentiary Hearing
   3060C - Informal Reconsideration (Applies to ICFs/IID for Denial of Payment for New Admissions Only)
   3060D - Judicial Review
   3060E - Impartial Decision Maker (Hearing Officer)

Prosp ective Payment System (PPS)
3100 - Hospitals and Hospital Units Excluded From the Inpatient Prospective Payment System (IPPS) - Introduction
3102 - General Information on IPPS Exclusion Deemed Providers and Suppliers
3104 - Criteria for PPS-Excluded Hospitals
   3104A - Psychiatric Hospitals
   3104B - Rehabilitation Hospitals
   3104C - Children’s Hospitals
   3104D - Long-Term Care Hospitals
   3104E - Hospital within Hospitals
3106 - Criteria for Psychiatric and Rehabilitation Units
   3106A - General Criteria for Units
   3106B - Specific Criteria for Psychiatric Units
   3106C - Specific Criteria for Rehabilitation Units
3108 - SA First-Time Verification Procedures for Hospitals and Units
   3108A - Rehabilitation Hospitals and Rehabilitation Units of Hospitals
   3108B - Psychiatric Units of Hospitals
3110 - SA Reverification of PPS-Excluded Hospitals and Units
   3110A - Annual Reverification Process for Nonaccredited, PPS-Excluded, Rehabilitation Hospitals and Units
   3110B - Reverification Process for Rehabilitation Hospitals and/or Units Accredited by CARF Under CIRP or JCAHO
   3110C - Reverification Process for Psychiatric Units of Hospitals:
3112 - RO Procedures for Exclusion from PPS for Hospitals and Units
   3112.1 - RO Procedures for First-Time Exclusion of Hospitals and Units
   3112.2 - RO Verifying Continued Compliance With Exclusion Criteria by Currently Excluded Hospitals or Units
   3112.2A - Self-Attestation Procedures for PPS-Excluded Hospitals and Units
   3112.2B - RO Verifying Exclusion Eligibility of Other Facilities
3112.3 - Role of FIs in Reverification of PPS Excluded Hospitals and Units

Changes in Provider Status or Services
3200 - Action Based on Changes in Provider Organization, Services, or Action of Other Approving Agencies
3202 - Change in Size or Location of Participating SNF and/or NF
  3202A - Requirements for Distinct Part Certification
    3202A1 - Meet Distinct Part Certification
    3202A2 - Do Not Meet Distinct Part Certification
  3202B - Changes in Bed Size of Participating SNF and/or NF
  3202C - General Request Filing Requirements
  3202D - Exceptions
  3202E - Change in Designated Bed Location(s)
  3202F - RO or SA (as appropriate) Actions Upon Receipt of Written Request for Change in Bed Size/Location
  3202G - Evaluation
  3202H - Survey Considerations
3206 - Existing ESRD Facility Relocation, Expansion, or Addition of New Service
3210 - CHOW of Providers and Suppliers
  3210A - Existing PoC
  3210B - Compliance With Health and Safety Standards
  3210C - Compliance With Ownership and Financial Interest Disclosure Requirement
  3210D - Compliance With Civil Rights Requirements
  3210E - All Medicare Sanctions and Penalties
3210.1 - Determining Ownership
  3210.1A - General
  3210.1B - SA Actions to be Taken Following CHOW
  3210.1C - Certification of Accredited Providers/Suppliers Which Change Ownership
  3210.1D - CHOW Situations
  3210.1E - CHOW Analysis General Rules
3210.2 - RO Role in CHOW Determination
3210.3 - CHOWS Involving Multi-Regional Chain Organizations
3210.4 - Other Changes Related to CHOW - RO Procedures
  3210.4A - New Owner Requests Different Intermediary
  3210.4B - New Owner Sets Different Fiscal Reporting Period
  3210.4C - New Provider Number Must Be Issued
3210.5 - New Owner Refuses to Accept Assignment of the Provider Agreement
3210.5A - New Owner Refuses to Accept Assignment of Previous Owner’s Provider Agreement
3210.5B - Withdrawal After CHOW - Provider
3210.5C - CHOW and Withdrawal - Supplier

Expansion of Services
3220 - Certifications of Additional Services
  3220A - Services in Compliance
  3220B - Services Not In Compliance (HHAs, RHCs, and ESRD Facilities)
3222 - Specific Requirements for Expansion of Services
  3222A - HHA’s Request to Provide OPT Services on Its Premises
  3222B - RHC’s Request to Provide Visiting Nurse Services
3224 - Addition of Sites to an Existing Provider

Validation Surveys of Accredited Providers and Suppliers
3240 - Validation Surveys - General
3241 – Objective of Validation Surveys
3242 - Representative Sample Validation Surveys of Deemed Providers/Suppliers
3243 - Substantial Allegation Validation Surveys of Deemed Providers/Suppliers
3244 - SA Preparation for Validation Survey
3246 - Provider/Supplier Authorization for Validation Survey
3248 – Provider/Supplier Refusal to Permit Validation Survey
3252 - SA Forwarding Validation Survey Records to RO
3254 - RO Actions Following Validation Survey
  3254A – Providers/Suppliers Found in Compliance Following Validation Survey
  3254B - Providers/Suppliers Found Not In Compliance With One or More Conditions Following Validation Survey and Noncompliance Constitutes Immediate Jeopardy
  3254C - Condition-level Deficiencies That Do Not Pose Immediate Jeopardy
  3254E - Plans of Correction
  3254F - Termination
  3254G - Compliance with All Conditions After Correction of Deficiencies
3256 - RO Provision of Information to Accrediting Organizations
3257 - Reinstatement to Accrediting Organization Jurisdiction
3258 – Termination of Accreditation

Handling Public Inquiries
3300 - Confidentiality and Disclosure of Records - Citations and Applicability
3302 - Federal Freedom of Information Act (FOIA)
3304 - Multi-Program Information in SA Files
3305 - Sharing State Licensure Information With Medicare Contractors
3308 - Information That May Be Disclosed to Public
  3308A - Information Disclosable to Public Under CMS Rules That May Be Disclosed Directly by the SA
3310 - Requests for Information About Nonparticipating Institutions
3312 - Charges for Information
3314 - Time Periods for Disclosure Other Than Nursing Homes
3316 - Information Furnished to Original Source
3318 - Disclosure of Information To and From Operating Components
3319 - Monthly Quality Indicator Comparison Reports Policy
  3319A - Purpose
  3319B - Contents
  3319C - Differences From Facility Quality Indicator Profile Report
  3319D - Delivery of Report
  3319E - Release of Report
  3319F - Monthly Quality Indicator Comparison Report Guidelines
3320 - Necessary Preclearance With RO Before Releasing Confidential Information

Additional State Agency Responsibilities

3330 - HHA Toll-Free Hotline and Investigative Unit
  3330A - HHA Hotline Function
  3330B - HHA Hotline Information
  3330C - Disclosure of Information
  3330D - Record Keeping Requirements
  3330E - Public Awareness
  3330F - Hotline Investigative Unit

Response to Subpoenas Served On and Suits Against the State Agency

3350 - Subpoena for Program Records
3352 - Forthwith Subpoena
3354 - Subpoena for SA Licensure Records
3356 - Suit Against SA
Adverse Actions

3000 - Adverse Actions - General
(Rev. 1, 05-21-04)

3000A - Applicability
(Rev. 1, 05-21-04)

The Regional Office (RO) and State survey agency (SA) follow the procedures in this part if an adverse action is likely to be initiated against Medicare participating providers and suppliers. Because many Medicare providers and suppliers also participate in the Medicaid program and Federal procedures must also be followed when surveying and certifying providers that only participate in the Medicaid program, these procedures generally apply to both programs. Exceptions for Medicaid are noted. (See also Chapter 7 for specifics pertaining to SNFs and NFs.)

For Medicaid-only facilities, termination procedures are not State plan requirements. However, a State risks disallowance of Federal matching funds for failure to use Federal standards and the forms, methods, and procedures prescribed by CMS. (See 42 CFR 442.30.)

3000B - State Ombudsman Programs
(Rev. 1, 05-21-04)

To coordinate with the State ombudsman network, the SA should establish procedures to:

- Notify the State ombudsman of decisions to initiate proceedings to terminate, or nonrenew a provider agreement;
- Notify the State ombudsman of voluntary terminations and planned terminations, including dates of closure;
- Consider ombudsman information about situations in the facility and the credibility of the provider’s allegations of compliance; and
- Share Statements of Deficiencies and Plans of Correction (PoCs).

3000C - CMS Authority to Terminate Medicare and Medicaid Participation
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

1. Noncompliance with Conditions of Participation (CoPs), Conditions for Coverage, or Requirements for SNFs - The RO is delegated authority to terminate Medicare participation of any certified provider or supplier because of noncompliance with the applicable regulatory requirements, or Conditions of Participation (CoPs) or Conditions for Coverage (CfCs).
2. Violations of Medicare Provider Agreements or certified Supplier Approvals, Quality Improvement Organization (QIO) Sanctions, or Program Abuse - The Secretary’s authority to terminate provider agreements or certified supplier approvals is delegated to the Associate Regional Administrator and may be redelegated to the RO Branch Chief, but other components may also be authorized to find that termination is in order. Accordingly, the RO processes terminations on grounds other than noncompliance with the CoPs. See §3032.

3. “Look Behind” Cancellation of Medicaid Intermediate Care Facility/Individuals with Intellectual Disabilities (ICF/IID) Agreements - The ROs are authorized to cancel the approval of an ICF/IID to participate in the Medicaid program when the ICF/IID fails to comply substantially with the applicable CoPs. (See §1910(b) of the Act.)

4. Termination of Nursing Facility (NF) Medicaid Agreements - The ROs are, under certain circumstances, authorized to terminate a NF’s participation in the Medicaid program. (See §1919(h) of the Act and Chapter 7 of the SOM.)

3001 - Initial Denials of Medicare Provider/Supplier Requests for Program Participation
(Rev. 1, 05-21-04)

Denials are made only when there has been an expression, written or otherwise, of interest in participating (or in expanding the scope of existing participation) and/or an initial survey is performed. Because the RO makes the compliance decision for Medicare and approves an effective date of participation, the SA should not lead the provider/supplier to believe that the provider/supplier has been approved and can start to furnish services to Medicare beneficiaries.

An initial denial is made when, after evaluating the evidence the adjudicating office (in this case the RO) finds that the requirements of law and regulation are not met. The SA forwards recommendations for initial denials to the RO within 10 working days after the date of survey. Formal written denial notices that explain the right to appeal are issued by the RO as soon as possible.

3001A - Authority for Adjudicating Denials
(Rev. 1, 05-21-04)

The RO adjudicates all approvals or disapprovals for Medicare participation. 42 CFR 498 addresses determination and appeal procedures. Title 42 CFR Part 488 provides the basis for denying suppliers of services. The statutory authority is implied in §§1819, 1832, 1861, and 1881 of the Act which authorize the Secretary to establish CoPs or CfCs.

3001B - Vacated Actions Which Are Not Denials
(Rev. 1, 05-21-04)
If the SA is contacted by a potential provider and it schedules a survey, but the survey is canceled after finding that the party is either no longer interested in participating or in meeting program requirements, the SA notifies the RO by Form CMS-1539, indicating the lack of interest. The RO sends a written notice to the potential provider to document the reason why certification action was not completed. Despite the lack of interest, if the potential provider operates a SNF, and the SA has sufficient information, the SA prepares a §1819(a)(1) (formerly §1861(j)(1)) certification, if indicated. (See §2164.)

3001C - Vacated Actions Which Are Denials
(Rev. 1, 05-21-04)

If a potential provider or supplier is surveyed and deficiencies are cited, the SA forwards Form CMS-1539 and related documentation to the RO, even when the request for participation is withdrawn. The RO either notifies the provider or supplier of the failure to meet eligibility requirements or affirms the provider’s or supplier’s request to withdraw. The SA uses Form CMS-1539 to transmit all certification forms and pertinent documents to the RO within 45 calendar days of the survey. A §1819(a)(1) certification is included, if indicated.

NOTE: The RO, based on notification by the FI or carrier of their inability to verify environment data, may issue the provider or supplier a notice of Medicare denial of participation.

3001D - RO Processing of Denials
(Rev. 1, 05-21-04)

The RO processes the denial and sends the provider or supplier a formal notice, with a copy to the SA, the State Medicaid Agency (SMA), and the intermediary, if applicable, documenting the basis for the action.

The RO includes the following information in the formal notice:

- The date of the notice;
- The decision and reason for it (cite provisions of the law or regulations not met);
- The right to request participation in the future; and
- The procedures to follow for a formal reconsideration and a hearing before an administrative law judge (ALJ).

A denial notice must be signed by the RO official delegated to adjudicate denials.
3005 - Basis for Terminating Provider Participation - Citations and Discussion
(Rev. 1, 05-21-04)

3005A - Medicare Provider Agreements
(Rev. 1, 05-21-04)

Provider agreements and agreements with clinics that provide Outpatient Physical Therapy (OPT) and Community Mental Health Centers (CMHCs) are terminated by the RO under the authority of §1866(b) of the Act. (See 42 CFR 489.52-489.57.) Medicare providers (as defined in §2002) must substantially meet each of the applicable CoPs or requirements for participation.

3005B - Termination of Coverage of Supplier Services Subject to Certification
(Rev. 1, 05-21-04)

Sections 1832(a), 1861(g), (p), (s), and (aa) and 1881(b) of the Act authorize the Secretary to establish CfCs of supplier services and thus implicitly authorize determinations that the Conditions cease to be met. Title 42 CFR 498.3(b) provides that the Secretary makes findings, setting forth pertinent facts and conclusions, and an initial determination as to whether a supplier meets the respective Conditions. The determination can be a result of the written request by the supplier to start or expand services or to establish that it continues to meet respective CfCs. An adverse determination may involve one or more areas of services offered by a supplier. While these adverse determinations are not in the regulations as “terminations,” their effect on payment for the supplier’s services is the same as when a provider agreement is terminated. Procedures for certifying supplier noncompliance parallel those for certifying provider noncompliance.

For example, the agreement into which an Ambulatory Surgical Center (ASC) or Rural Health Clinic (RHC) enters is a specific agreement related to those suppliers, not a provider agreement.

3005D - Cause for Termination
(Rev. 1, 05-21-04)

The CMS may terminate Medicare provider participation if the provider does not comply with a CoP or Requirement for SNFs/NFs, or fails to provide an acceptable PoC for other requirements. (See 42 CFR 489.53.)

Certain causes for termination are unrelated to certification and have no impact on the SA. The CMS may terminate provider participation if:
1. The provider places restrictions on the persons it accepts for treatment, and fails either to exempt Medicare beneficiaries from the restrictions, or to apply the same restrictions to Medicare beneficiaries as to all other persons seeking care;

2. The provider refuses to permit examination of its records by or on behalf of CMS for verification of information it furnished as a basis for payment;

3. The provider has knowingly and willfully made false statements or representations of a material fact for use in a request for payment;

4. The provider has submitted, or caused to be submitted, requests for payment under Medicare, or amounts for items and services, substantially in excess of the costs incurred;

5. The provider has furnished items or services which CMS determined to be substantially in excess of the needs of individuals or of a quality that failed to meet professionally recognized standards; or

6. The provider fails to:
   - Permit photocopying of any records necessary to determine compliance;
   - Furnish information necessary for CMS to determine whether payments are or were due under Medicare and the amount due;
   - Furnish information on business transactions as required;
   - Disclose information on convicted principals;
   - Furnish ownership information;
   - Comply with civil rights requirements; or
   - Furnish notice of discharge rights.

7. A hospital or critical access hospital that has reason to believe it may have received an individual by another hospital in violation of 42 CFR 489.24 fails to report the incident.

8. A hospital fails to furnish inpatient services to TRICARE or CHAMPVA beneficiaries or to veterans.

9. A critical access hospital fails to maintain an annual average length of stay of 96 hours or less.
The SA certifies provider compliance with Medicare requirements. FIs generally are responsible for dealing with matters related to payment and coverage. However, in the course of a survey, the SA may encounter information indicative of program abuse or failure to meet requirements described in the above list. The SA communicates these areas of concern to the RO.

3005E - Termination of Title XIX-Only NFs, ICFs/IID, Hospitals and Psychiatric Hospitals
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Under 42 CFR 431.54(f), the State Medicaid Agency may “lock out,” i.e., restrict the participation of a Medicaid provider for a reasonable period if it has found that the provider has abused the Medicaid program. The SMA may take this action in response to, or independent of any SA or accrediting organization recommendation. Note that the Medicaid program, unlike the Medicare program, does not distinguish between “providers” and “suppliers” and the term “provider” applies to both for Medicaid purposes.

In order to determine that a provider has abused the Medicaid program, the SMA must find that in a significant number or proportion of cases the provider has:

- Furnished services at a frequency or amount not medically necessary (§431.54(f)(2)(i)); or
- Furnished Medicaid services of a “quality that does not meet professionally recognized standards of health care.” (§431.54(f)(2)(ii)).

Additional conditions required before imposing any restriction on a provider are found in §431.54(f).

Medicaid-only NFs and ICFs/IIDs

Medicaid regulations provide for terminations, and for ICFs/IID cancellations, but do not describe the implementing procedures. Each SMA has procedures for terminating agreements with NFs and ICFs/IID when they are not in substantial compliance with program requirements. In any Medicaid-only noncompliance situation, the SA initiates the action, prepares the necessary documents, and forwards them to the SMA, which has responsibility for the termination, nonrenewal, or cancellation of the agreement (see §7300 for the exception regarding State operated NFs). In this case, the SMA notifies CMS and the public of its action and affords the facility notice and opportunity for a hearing before an ALJ prior to termination.

Medicaid-only Hospitals and Psychiatric Hospitals

42 CFR 482.1(a)(5) notes generally that any hospital or psychiatric hospital that participates in Medicaid must meet the Medicare CoPs. Medicaid regulations at 42 CFR
440.10 (hospitals) and §440.140 and 440.160 (psychiatric hospitals) require Medicaid-participating hospitals and psychiatric hospitals to comply with the Medicare CoPs. If a SA survey of a Medicaid-only hospital or psychiatric hospital finds substantial noncompliance, i.e., condition-level noncompliance, including immediate jeopardy, with the CoPs, the SA follows the standard SOM processes for requiring a plan of correction and conducting revisits. If the hospital or psychiatric hospital fails to come into substantial compliance in a timely manner, the SA notifies the SMA, which takes appropriate action in accordance with Federal Medicaid regulations and State plan provisions.

Likewise, in the case of a hospital or psychiatric hospital that participates in Medicaid based on accreditation by a CMS-approved Medicare hospital or Medicare psychiatric hospital accreditation program, the accrediting organization (AO) must notify both CMS and the SMA if the facility’s accreditation is terminated for any reason. The SMA may in such cases request the SA to conduct a survey to determine the hospital’s or psychiatric hospital’s compliance with the CoPs, or may proceed directly to take appropriate action in accordance with Federal Medicaid regulations and State plan provisions.

If the SMA decides to restrict the provider’s participation in Medicaid, the SMA is responsible for:

- Giving notice to the provider and opportunity for appeal, in accordance with procedures established by the SMA (§431.54(f)(1));
- Notifying CMS and the general public of the provider’s restriction and its duration (§431.54(f)(3)); and
- Ensuring that restrictions do not result in denying Medicaid recipient reasonable access to services of adequate quality (§431.54(f)(4)).

If the SMA restricts a Medicaid-only hospital’s or psychiatric hospital’s Medicaid participation, it gives notice to CMS via the SA. The SA forwards the notice to the RO. The RO terminates the hospital’s or psychiatric hospital’s Medicaid-only CCN, unless the SMA notice contains a date certain when the provider’s participation in Medicaid will resume.

3005F - Termination Action Based Upon Onsite Survey by RO, or Validation Survey of a Deemed Provider or Supplier by RO or SA (Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

RO Conducts Survey:

When substantial noncompliance, including immediate jeopardy to patient health and safety, is identified in the course of a survey conducted by RO staff, the RO initiates termination procedures. Survey findings and factual development are the responsibility
of the RO. However, the SA may be asked to assist in documenting or developing aspects of the termination. The SA (and the SMA, if the provider/supplier also participates in Medicaid) is notified by the RO of the action taken.

RO or SA Validation Survey of a Deemed Provider or Supplier

Section 1864(c) of the Social Security Act provides that SAs conduct validation surveys of deemed providers and suppliers, at the direction of the CMS RO. The RO may also choose to conduct the validation survey with RO surveyors, or with a mixture of SA and RO surveyors. There are two types of validation surveys:

- Full, standard surveys of a representative sample of deemed providers and suppliers, and
- Focused substantial allegation, i.e., complaint investigation surveys.

Representative Sample Validation Survey: If the representative sample validation survey identifies either an immediate jeopardy or substantial, i.e., condition-level, noncompliance and the RO agrees with this finding, the RO initiates termination of the deemed provider or supplier, including an opportunity for the provider/supplier to make a timely correction of the deficient practices to avoid termination.

Substantial Allegation Validation Survey:

- If the survey identifies an immediate jeopardy and the RO agrees with this finding, the RO initiates termination of the deemed provider or supplier, including an opportunity for the provider/supplier to make a timely correction of the deficient practices to avoid termination.

- If the survey identifies substantial, i.e., condition-level, noncompliance and the RO agrees with this finding, the RO may either:
  - initiate termination of the deemed provider or supplier, including an opportunity for the provider/supplier to make a timely correction of the deficient practices to avoid termination; or
  - Require the SA to conduct a full survey of the provider or supplier. Termination action would be initiated if the full survey identifies substantial noncompliance.

See Chapter 5, Section 5110 for more details on post-survey procedures after a substantial allegation validation survey.

3005G - Look Behind Authority of CMS
(Rev. 1, 05-21-04)
3005G1 - “Look-Behind” Termination or Cancellation of ICF/IID Agreement by the Secretary
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

The CMS has authority under §1910(b) of the Act to terminate approval of an ICF/IID to participate in the Medicaid program when it determines that the facility fails to comply substantially with the CoPs, 42 CFR Part 483, Subpart I, or to submit an acceptable PoC.

The cancellation is prospective, usually after the provider has had the opportunity for a formal hearing before an ALJ.

If there is no immediate jeopardy to resident health and safety and CMS elects to terminate, the ICF/IID is afforded an opportunity for a pre-termination hearing before an ALJ. If the effective date of termination is held in abeyance pending an ALJ’s ruling and the ICF/IID makes a credible allegation of compliance while the hearing is pending, it is up to the RO to determine whether it is in the recipients’ and the government’s interest to resurvey the facility and dispose of the case based on the findings. If a revisit is made and the ICF/IID failed to achieve compliance, adverse action continues based on the findings of the first Federal survey and the revisit. If the ALJ affirms the CMS decision, the effective date of termination is set by the ALJ.

If there is an immediate jeopardy to resident health and safety, CMS terminates or cancels approval of the ICF/IID and affords it the opportunity for a post-termination ALJ hearing.

Following termination, ICFs/IID wanting readmission must request a survey from the RO. The RO directs the SA to do a survey unless it feels that a Federal survey is necessary. The CMS must be satisfied that the reasonable assurance provision is met before the State executes a Medicaid agreement with the ICF/IID.

3005G2 - Old “Look-Behind” Termination of a NF or ICF/IID by the Secretary
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Under 42 CFR 442.30, a provider agreement of a SNF, NF, or ICF/IID is considered invalid for purposes of providing FFP to the State unless the State has followed proper survey and certification procedures. For example, the SMA may have issued the provider agreement even though it had not certified the facility as being in compliance. Other examples of procedural error include, but are not limited to:

- The SA documents noncompliance yet certifies compliance;
- The SA certifies compliance, but all cited deficiencies are not covered by an acceptable PoC;
- The SA fails to survey against all applicable requirements; or
The SA fails to use federally approved survey and certification documents.

When procedures are not followed by either the SA or SMA, CMS considers the provider agreement void from its inception, and the State is disallowed FFP for bills related to the facility for the period covered by that Medicaid agreement. This type of adverse action, referred to as “Old Look Behind,” is covered in more detail in §3042.

3005G3 - SMA Disagrees With SA Determination
(Rev. 1, 05-21-04)

With the exception of State-operated NFs, the SA surveys and certifies compliance of Medicaid facilities with health and safety requirements to the SMA. The SMA is responsible for reviewing certifications to ensure that the SA has adhered to procedural requirements. If the SMA disagrees with the SA’s certification, it first contacts the SA to resolve the issue. If the issue cannot be resolved, it contacts the RO. To resolve the dispute, the RO conducts a Federal survey of the facility or take other action as necessary.

3006 - Denial of Payments in Lieu of Termination of ICFs/IID
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

3006A - Authority to Deny Payment for Any New Admissions for ICFs/IID
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Section 1902(i) of the Act and 42 CFR 442.118 provide the SMA with an alternative to terminating ICFs/IID that fail to meet program requirements. This sanction is the one-time denial of payment for new admissions for a period of up to 11 months after the month it was imposed, if the facility’s deficiencies do not present an immediate jeopardy to residents’ health and safety. A decision is made at the end of 11 months whether to continue participation. However, the 11-month period can be shortened if circumstances change and there is immediate jeopardy to health and safety before 11 months have passed. Alternatively, the State might rescind the denial of payments in fewer than 11 months if full compliance is achieved or if the ICF/IID has made significant, good-faith efforts and progress in achieving compliance.

3006B - Criteria for Imposing Denial of Payments for New Admissions
(Rev. 1, 05-21-04)

The SMA retains the right to establish its own criteria for imposing this sanction. However, the SMA may not use this sanction if the facility’s deficiencies pose immediate jeopardy to the health and safety of its clients.

3006C - Agency Procedures
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)
Before denying payment for new admissions, the SMA must comply with the following requirements:

- Provide the ICF/IID up to 60 calendar days to correct the cited deficiencies and comply with the CoP.

- If at the end of the specified period the ICF/IID has not achieved compliance, give the facility notice of intent to deny admissions and the opportunity for an informal hearing.

- If the ICF/IID requests a hearing and the decision of the hearing is to deny payment, the SMA must provide the facility and the public, at least 15 calendar days before the effective date of the sanction, a notice that includes the effective date of the sanction and the reasons for the denial of payment.

3006D - Effect of Sanction on Status of Clients Admitted, Discharged, or on Temporary Leave and Readmitted Before or After Effective Date of Denial of Payment
(Rev. 1, 05-21-04)

The client’s status on the effective date of the denial of payment is the controlling factor in determining whether readmitted clients are subject to the denial of payment. Guidelines are as follows:

- Clients who were admitted and discharged before the effective date of the denial of payment are considered new admissions if they are readmitted on or after the effective date. Therefore, they are subject to the denial of payment;

- Clients admitted on or after the effective date of the denial of payment are considered new admissions. If readmitted after being discharged, they continue to be considered new admissions, and are subject to the denial of payment;

- Clients admitted before and discharged on or after the effective date of the denial of payment are considered new admissions if subsequently readmitted. Therefore, they are subject to the denial of payment;

- Clients admitted before the effective date of the denial of payment who take temporary leave before, on, or after the effective date of the denial of payment are not considered new admissions upon return and therefore, are not subject to the denial of payment; and

- Clients admitted on or after the effective date of the denial of payment who take temporary leave are not considered new admissions, but continue to be subject to the denial of payment.
NOTE: The term “temporary leave” refers to clients who leave temporarily for any reason. If clients were not subject to a denial of payment when they went on temporary leave, the term indicates that upon return they are not considered new admissions for the purposes of the denial of payment. Therefore, the term “temporary leave” is used to justify a resumption of any interrupted payment upon re-entry into the facility.

The term “leave of absence” is defined as any situation where the client is absent, but not discharged, for reasons other than admission to a hospital, SNF or NF, or distinct part of a SNF or NF. The term “leave of absence” is used for the purpose of preventing duplicate payments during an absence by assuring that the absence is not due to a temporary alternate inpatient arrangement. If the client is not on a leave of absence but is actually temporarily in an alternate inpatient situation, any ongoing payment to the facility will be interrupted as mentioned above.

The client who is not subject to the denial of payment sanction and who goes on temporary leave, whether there is a leave of absence, will not be considered a new admission for the purposes of the denial of payment sanction, upon his/her return to the facility. Any interrupted payment will be resumed. In either situation, it is expected that the client will return to the facility following leave.

3006F - Duration of Denial of Payment and Subsequent Termination of an ICF/IID
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

The denial of payment for new admissions will continue for eleven months unless, before the end of that period, the SMA finds that the ICF/IID has corrected the deficiencies or is making a good faith effort to achieve compliance with the CoPs or the deficiencies are such that it is necessary to terminate the facility.

The SMA must terminate the facility’s provider agreement:

- Upon finding that the ICF/IID has been unable to achieve compliance with the CoPs during the 11-month period that payments were denied for new admissions; and

- Termination is effective the day following the last day of the denial of payment period.

3006.1 - Sanctions for ICFs-IID - or Nonimmediate Jeopardy
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

3006.1A - General
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)
The Balanced Budget Act (BBA) of 1997 provided the statutory authority for States to establish and impose sanctions that are additional to the already existing alternative sanction of denial of payment for new admissions, and which are alternative to termination in cases where the ICF/IID’s deficiencies are not determined to pose immediate jeopardy to client health and safety. This strategy recognizes that deficiencies take on greater or lesser significance depending on the specific circumstances and client outcomes in each facility, and that additional enforcement options should be available so that the enforcement consequence to the facility is effective, proportionate, and appropriate to the specifics of the noncompliance.

3006.1B - Introduction
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Section 1902(i)(1)(B) of the Act, as revised by the BBA of 1997, provides that the State may establish alternative sanctions to use as enforcement remedies for deficiencies that do not constitute immediate jeopardy to client health and safety if the State can demonstrate to CMS’ satisfaction that its alternative sanctions are effective in deterring noncompliance and correcting deficiencies (see §3006.6). One or more alternative sanctions may be imposed against private or State operated ICFs/IID instead of provider agreement termination, and may also be imposed instead of or in addition to the existing alternative sanction of denial of payment for new admissions. Examples of sanctions that may be appropriate as alternative sanctions are listed in subsection C.

3006.1C - Examples of Alternative Sanctions
(Rev.92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

States should consider establishing the following alternative sanctions in their State plan for noncompliant ICFs/IID having nonimmediate jeopardy deficiencies:

- Directed plan of correction;
- Directed in-service training; and
- State monitoring.

States are not limited to establishing and using these alternative sanctions and may submit others for CMS’ approval. When the State wants to use alternative sanctions, it must be authorized to do so under its State plan by CMS (see §3006.6). In order to be approved, the State must provide specified information to indicate that the alternative sanction and its application is not inconsistent with applicable statutory and regulatory requirements, as well as demonstrate to CMS’ satisfaction that the alternative sanction is effective in deterring noncompliance and correcting deficiencies. Many States already have experience in imposing the three intermediate sanctions specified above against nursing homes (SNF/NFs) that fail to meet participation requirements. States also have experience in imposing remedies under their State licensure authority and may also wish to submit any of those to CMS for approval as alternative remedies for ICF/IID. While
we want States to have the three specified intermediate sanctions listed above available for ICF/IID enforcement purposes, States are free to submit others for CMS approval as well.

3006.1D - Alternatives to Termination in Nonimmediate Jeopardy Situations
(Rev. 1, 05-21-04)

When the facility is found to have deficiencies that do not immediately jeopardize the health and safety of individuals served, the State may, in lieu of terminating the facility’s provider agreement:

1. Deny payment for all new Medicaid admissions to the facility after the effective date of the sanction;

2. Impose one or more of the alternative sanctions that CMS has approved; or

3. Do both (1) and (2) above. Deny payment for all new Medicaid admissions to the facility after the effective date of the sanction and impose one or more of the alternative sanctions that CMS has approved.

3006.2 - Directed Plan of Correction (DPoC)
(Rev. 1, 05-21-04)

3006.2A - Purpose
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

A DPoC is a plan that the State develops to require an ICF/IID to take action within specified time frames. The purpose of the DPoC is to achieve correction and continued compliance with the CoPs.

A DPoC differs from a traditional PoC in that the State, not the facility, develops the PoC. Achieving compliance is the provider’s responsibility, whether or not a DPoC was followed. If the facility fails to achieve substantial compliance after complying with the DPoC, the State may impose another alternative sanction (or sanctions) until the facility achieves substantial compliance or it is terminated from the Medicaid program.

3006.2B - Basis for Imposition of a DPoC
(Rev. 1, 05-21-04)

Use of a DPoC should be dependent upon causal factors identified by the SA. For example, a DPoC may be an appropriate sanction when a facility has no system in place for detecting abuse and neglect. The DPoC would specify that the facility must develop a system and must have that system in place within a specified time frame.

3006.2C - Elements of a DPoC
(Rev. 1, 05-21-04)
The DPoC includes all elements of a traditional plan of correction (see §3006.5.C), as well as when the corrective action must be accomplished, and how substantial compliance will be measured.

3006.2D - Notice of Imposition of DPoC  
(Rev. 1, 05-21-04)

A DPoC may be imposed 15 calendar days after the facility receives notice of this sanction. The date the DPoC is imposed does not mean that all corrections must be completed by that date.

3006.3 - Directed In-Service Training  
(Rev. 1, 05-21-04)

3006.3A - Purpose  
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Directed in-service training is a sanction that may be used when the SA concludes that education is likely to correct the deficiencies. This remedy requires the staff of the ICF/IID to attend in-service training program(s). The purpose of the directed in-service training is to provide knowledge required to achieve compliance and remain in compliance with the CoPs.

3006.3B - Appropriate Resources for Directed In-Service Training Programs  
(Rev.92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Facilities should use programs developed by well-established organizations of intellectual disabilities, developmental disabilities, mental health or health services education, such as special education departments in colleges or universities or schools of medicine, State departments/bureaus of mental health/ intellectual disabilities or developmental disabilities; Developmental Disabilities Councils; Federally funded State protection and advocacy agencies serving people with developmental disabilities; professional organizations with expertise in developmental disabilities, and a State may provide special consultative services for obtaining this type of training. The SA may also compile a list of resources that can provide directed in-service training and could make this list available to facilities and interested organizations. Facilities may request to use training resources internal to the organization, if the trainer was not directly involved with the area sanctioned. Examples of directed in-service training topics include, but are not limited to, client rights issues, behavior intervention, active treatment, health and safety, and outcome measures.

3006.3C - Further Responsibilities  
(Rev. 1, 05-21-04)
The facility bears the expense of the directed in-service training. After the training has been completed the SA will assess whether agency staff has demonstrated competency in the area(s) of deficiency and whether compliance has been achieved. If the facility still has not achieved substantial compliance, the State may impose one or more additional sanctions.

3006.3D - Notice of Imposition of Directed In-Service Training
(Rev. 1, 05-21-04)

Directed in-service training may be imposed 15 calendar days after the facility receives notice of this sanction. The SA will determine time frames for completion of the directed in-service training.

3006.4 - State Monitoring
(Rev. 1, 05-21-04)

3006.4A - Purpose
(Rev. 1, 05-21-04)

A State monitor oversees the correction of cited deficiencies in the facility as a safeguard against further noncompliance when a situation with a potential for jeopardizing health and safety has occurred, but has not risen to the level of immediate jeopardy.

3006.4B - Qualifications
(Rev. 1, 05-21-04)

The SA identifies state monitors as appropriate professionals to monitor cited deficiencies. A State monitor:

- Is an employee or contractor of the SA;
- Is not an employee, designee or contractor of the monitored facility;
- Does not have an immediate family member who is a client of the facility;
- Is not a person who has been terminated for cause by the facility; and
- Is not a former contractor who had a contract canceled for cause by the facility.

3006.4C - When to Impose State Monitoring
(Rev. 1, 05-21-04)

When considering whether or not to impose State monitoring for current noncompliance, the State may want to consider whether:
• The facility has a history of noncompliance which may suggest that it would benefit from external surveillance during corrections;
• The facility has had numerous complaints; or
• The State is concerned that the situation in the facility has the potential to worsen.

States are not limited to considering only these factors and are free to consider any others that would assist them in making remedy determinations.

3006.4D - Frequency
(Rev. 1, 05-21-04)

When State monitoring is imposed, the SA appoints a monitor or monitors. Monitoring may occur anytime in a facility; e.g., 24 hours a day, 7 days a week, if necessary or less often such as once a week to monitor specific areas. In all instances, monitors have complete access to all areas of the facility, as necessary, for performance of the monitoring activity. Factors used to decide how often a facility is monitored may include, but are not limited to, the following:

• The nature and seriousness of the deficiency(ies) as specified by the SA; and
• The timing and frequency of when the problems occurred; e.g., mealtimes, evening shifts, daily, etc.

Monitors may be assigned to the facility at these specific times for a specified number of days, as determined by the SA, to ensure corrective action.

3006.4E - Duration
(Rev. 1, 05-21-04)

The sanction is discontinued when the facility’s provider agreement is terminated or when the facility has demonstrated to the satisfaction of the SA that it is in substantial compliance with the Conditions of Participation.

3006.4F - Notice of Imposition of State Monitoring
(Rev. 1, 05-21-04)

Notice requirements for this sanction state that it may be imposed immediately. No notice is required because the sanction imposes no hardship or expense on the facility.

3006.4G - Payment for and Obligation of Funds by a State Monitor
(Rev. 1, 05-21-04)
The facility will not be required to pay the salary of the State monitor; nor will the State monitor have managerial authority to obligate facility funds.

3006.5 - Achieving Continuous, Substantial Compliance
(Rev. 1, 05-21-04)

3006.5A - Introduction
(Rev. 1, 05-21-04)

In order to safeguard the health, welfare and safety of individuals served within a facility, it is imperative that a facility not only attain substantial compliance in each area of identified deficiency(ies), but that it maintain/remain in continuous compliance.

3006.5B - Duration of a Sanction
(Rev. 1, 05-21-04)

A sanction is discontinued when the facility's provider agreement is terminated or when the facility has demonstrated to the satisfaction of the SA that it is in substantial compliance with the Conditions of Participation.

3006.5C - Achieving and Maintaining Substantial Compliance
(Rev. 1, 05-21-04)

The facilities must establish policies and procedures to remedy deficient practices and to ensure that correction is lasting. Facilities must take the initiative and responsibility for monitoring their own performance continuously to sustain compliance. In order for a PoC to be acceptable, it must include the following elements:

1. Core Elements of PoC:
   a. How the corrective action will be accomplished for individuals found to have been affected by the deficient practice;
   b. How the facility will identify other individuals who have the potential to be affected by the same deficient practice, and how the facility will act to protect individuals in similar situations;
   c. What measures will be put into place or systemic changes made to ensure that the deficient practice will not recur;
   d. How the facility will monitor its corrective actions/performance to ensure that the deficient practice is being corrected and will not recur, i.e. what program will be put into place to monitor the continued effectiveness of the systemic change to ensure that solutions are permanent; and
   e. When corrective action must be accomplished.
3006.6 - Criteria for Review of State Plans for Approval or Disapproval of Alternative Sanctions  
(Rev. 1, 05-21-04)

3006.6A - Introduction  
(Rev. 1, 05-21-04)

This section implements §1902(i)(1)(B) of the Act and provides guidance to the RO relative to reviewing for approval or disapproval, State plan amendments for alternative enforcement sanctions.

3006.6B - Alternative Remedies  
(Rev. 1, 05-21-04)

If a State wishes to establish alternative sanctions in addition to the already existing alternative sanction of denial of payment for new admissions, to be used in situations of nonimmediate jeopardy, the State plan should describe:

- Timing and notice requirements;
- When the remedy will be applied;
- How the alternative remedy is effective in deterring noncompliance; and
- Factors considered in selecting the remedy.

3008 - Services After Termination  
(Rev. 1, 05-21-04)

Payment may be made for a limited time for some patients after the effective dates of termination. (See §§3008.1 and 3008.2.)

3008.1 - Services After Termination of a Medicare Provider Agreement  
(Rev. 1, 05-21-04)

Effective the date the provider agreement is terminated, no payment will be made under the agreement. (See 42 CFR 489.55.) However, payment is available for up to 30 calendar days after the effective date of termination for beneficiaries admitted before the effective date of termination for:

- Inpatient hospital services;
- Psychiatric hospital services; and
- SNF services.

Also, payment is available for up to 30 calendar days for care furnished under a plan established before the effective date of termination for:

- Home Health Agency (HHA) services; and

- Hospice care.

3008.2 - Services for which Federal Financial Participation (FFP) May Be Temporarily Continued After Termination of a Medicaid Provider Agreement or Nonrenewal or Cancellation of an ICF/IID Provider Agreement
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Federal Financial Participation (FFP) may continue for up to 30 calendar days after the effective date of termination if the Medicaid beneficiaries were admitted to the entity before the effective date of termination and the State is making reasonable effort to transfer those beneficiaries to other facilities or to alternate care or to ensure that they are appropriately discharged. (See 42 CFR 441.11.) Services for which FFP may be continued are:

- Inpatient hospital services (for both dually-participating and Medicaid-only hospitals);

- Inpatient hospital services for individuals age 65 or older in institutions for mental disease (IMD) (for both dually-participating psychiatric hospitals and Medicaid-only psychiatric hospitals which are IMDs);

- NF services;

- NF services for individuals age 65 or older in IMD;

- Inpatient psychiatric services for individuals under age 21(for both dually-participating and Medicaid-only psychiatric hospitals); and

- ICF/IID services.

3008.3 - Relocating Patients Displaced by Termination or Closure
(Rev. 1, 05-21-04)

3008.3A - General
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)
There are instances when patients in Medicare and Medicaid long-term care facilities need to be transferred to other facilities. Specific actions, decisions, and events that require the relocation of patients include:

- Voluntary or involuntary termination of provider agreement;
- Expiration or renewal of an ICF/IID provider agreement;
- The provider’s inability to provide care and related services because of fire, natural disaster, loss of staff, or another reason beyond its control;
- The provider’s voluntary termination of participation in Medicaid and/or Medicare; and
- Closure of a facility.

3008.3B - Relocation of Medicaid Patients
(Rev. 1, 05-21-04)

The SMA has the primary responsibility for relocating Medicaid patients and for ensuring their safe and orderly transfer from a facility that no longer participates in Medicaid to a participating facility. This is because the State remains responsible for the care and services provided to Medicaid patients. The State’s transfer policies must:

- Consider the nature and severity of the facility’s failure to meet standards;
- Consider the availability of alternative facilities;
- Ensure that the situation is explained to the recipient and the recipient is permitted to exercise an informed choice as to whether he or she wishes to move and, if so, to which available facility.
- Provide that qualified personnel will assess patients’ medical and psychological condition and needs, including the necessity to prepare the patient for transfer;
- Provide for adequate and appropriate transportation on the day the patient is moved; and
- Apprise the receiving facility of the patient’s condition and needs.

3008.3C - Relocation Activities
(Rev. 1, 05-21-04)

The State should develop a long-term care patient relocation plan for the orderly transfer of patients. The plan should provide for:
• Decisions to be made on relocation of patients on a case-by-case basis;

• Independent current assessment of the patient’s need for institutional care and the level and type of care needed;

• Procedures they will employ to ensure that the transferring facility maintains acceptable health care standards and takes necessary precautions to minimize fire hazards;

• Description of the organization and staffing necessary to implement the plan; and

• Necessary agreements and approvals for the transfer of the patient.

The State plan should have a degree of flexibility that will enable it to be used for the relocation of any number of patients. The frequency with which States will implement their patient relocation plans will vary considerably from State to State. However, it should be developed as an ongoing, standard operating procedure that can be implemented quickly and efficiently. The plan should be developed by the State Medicaid agency in conjunction with the SA.

3010 - Termination Procedures - Immediate Jeopardy to Patient Health and Safety (Medicare)
(Rev. 1, 05-21-04)

(See §§7307 - 7309 for SNFs/NFs.)

3010A - Substantial Noncompliance With Program Requirements Which Poses Immediate Jeopardy to Patient Health or Safety
(Rev. 1, 05-21-04)

“Immediate Jeopardy” is interpreted as a crisis situation in which the health and safety of patients is at risk. Generally, it is a deficient practice that indicates the operator’s inability to furnish safe care and services, although it may not have resulted in actual harm. The threat of probable harm is real and important and could be perceived as something that will result in potentially severe temporary or permanent injury, disability, or death. Therefore, it must be perceived as something that is likely to occur in the very near future. If the patients are not protected effectively from the threat, or if the threat is not removed, there is a high probability that serious harm or injury could occur at any time or already has occurred and may occur again.

A list of operational definitions of what can constitute an immediate jeopardy to patient health and safety is presented as a guide to be used by all surveyors. (See Appendix Q.) Generally, the criteria applies to most providers and suppliers, although some criteria may apply to only certain types of providers or suppliers. The operational definitions are
not intended to be all-inclusive, nor are they intended to inhibit the professional judgment of the surveyors. Surveyors may find that an immediate jeopardy does not exist when the definitions seem to apply or that such a threat does exist even though the definitions do not address the situation or condition observed by the surveyors.

The key factor in the use of the immediate jeopardy termination authority is, as the name implies, limited to immediate and serious. The threat must be present when you are onsite and must be of such magnitude as to seriously jeopardize a patient’s health and safety. There should be no other application of immediate jeopardy terminations. Do not use these procedures to enforce compliance quickly on more routine deficiencies.

3010B - Processing of Immediate Jeopardy Terminations
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

When an immediate jeopardy to patient health or safety is documented, the SA and RO complete termination procedures within 23 calendar days. Processing times given here are the maximum allowed. Do not postpone or stop the procedure unless compliance is achieved and documented through onsite verification. If there is a credible allegation that the threat or deficiency has been corrected, the SA conducts a revisit prior to termination if possible.

Deemed Providers/Suppliers:

See Chapter 5, Section 5110.2, for the procedures and timelines applicable when an SA validation survey identifies an immediate jeopardy in a deemed provider or supplier.

Special Procedures for IJ in Psychiatric Hospital Surveys Conducted by CMS’ Contract Surveyors

When a psychiatric hospital survey is conducted by CMS contract surveyors and they identify immediate jeopardy to patient health or safety, the RO will make the final determination as to whether or not there is an immediate jeopardy situation. The CMS contract surveyors will notify the RO during survey if possible that immediate jeopardy to patient health or safety is identified. On the last day of the survey, CMS contract surveyors contact the CMS Central Office (CO) and the applicable RO to certify noncompliance and that an immediate jeopardy exists. The CMS contract surveyors discuss their findings with the provider and tell the providers that they are mailing the RO by overnight express mail completed Forms CMS-1537A and CMS-2567. A copy is also mailed to CO for review. The RO reviews the survey package (Forms CMS-1537A and CMS-2567), and if it determines noncompliance, it mails Form CMS-2567 to the provider. After doing so, the RO follows the 23 calendar day termination procedure as outlined below beginning with the fifth working day.

All Other SA Surveys with IJ Findings: 23-Day Termination Procedures
1. Date of Survey - The date of the survey is the date on which the entire survey is completed, regardless of when the exit conference is held.

2. Second Working Day - No later than 2 working days following the survey date. The SA:
   
   - Telephones the RO that it is certifying noncompliance and that an immediate jeopardy exists; and
   
   - Notifies the provider SUPPLIER (by overnight express mail, FAX or e-mail) of its deficiencies and informs the provider SUPPLIER that it is recommending termination to the RO, which will issue a formal notice. The notice advises the provider/supplier of its right to due process, the expected schedule for termination action, and that the deficiency must be corrected and verified by the SA to halt the termination. If the provider also participates in Medicaid, the SA notifies the SMA of its certification of noncompliance.

3. Third Working Day - The SA forwards all supporting documentation to the RO (e.g., statement of deficiencies, correspondence, contact reports, Form CMS-1539). The SA forwards the information by overnight mail to assure that the RO receives it in time to meet the 5-working-day deadline. Upon receipt of the SA information, the RO reviews the documents and makes its determination of noncompliance.

4. Fifth Working Day - The provider SUPPLIER and the public are then notified by the RO of the proposed termination action by the most expeditious means available. A press release to the radio and television stations serving the area in which the provider SUPPLIER or institution is located is acceptable if a newspaper notice cannot be arranged in the time allotted. Notice must be made at least 2 calendar days prior to the effective date of termination. (See 42 CFR 488.456(c).)

5. Tenth Working Day - If the SA only sent notification of the IJ deficiencies on the second working day to the provider SUPPLIER and RO, and there are other, non-IJ deficiencies (non-IJ condition and standard level), then the SA must write up another 2567 with the non-IJ deficiencies and forward copies to the provider SUPPLIER, the RO and SMA within ten working days. The SA retains a copy for its records.

6. Twenty-Third Calendar Day - The termination takes effect unless compliance is achieved or threat is removed. If the threat has been removed, but deficiencies still exist at the Condition level, the SA gives the provider SUPPLIER up to 67 more calendar days, or 90 calendar days total (23 plus 67). These dates are maximum times, and participation may be terminated earlier if processing allows. However, the RO must adhere to both the provider SUPPLIER and public notice timeframes.
If the RO disagrees based upon its review of the documentation, the RO discusses the results of the review with the SA and solicits further evidence to support the SA’s recommendation. The RO confers with the SA as to the appropriate action to be taken. Should the RO and the SA fail to agree that an immediate jeopardy exists, a revisit will be conducted by the RO and the SA together to ascertain whether or not immediate jeopardy to the patient’s health and safety exists or has been removed. If the RO and SA agree that an immediate jeopardy exists, no revisit is necessary by the RO. Under no circumstances should the RO reverse a SA recommendation that an immediate jeopardy has been removed or not removed unless the determination is made on the basis of an onsite determination by Federal surveyors.

Medicaid agreements with facilities that concurrently participate in Medicare should be terminated on the same date the Medicare agreement is terminated. For NFs that also participate as SNFs (i.e., dually-participating), the State’s timing of termination shall control if it does not occur later than six months after the last day of the survey when both CMS and the State find that a facility is not in substantial compliance and the facility’s participation should be terminated. (See 42 CFR 488.452.)

For NFs, ICFs/IID, Medicaid-only hospitals and Medicaid-only psychiatric hospitals, where State law or the State Medicaid plan permits, Medicaid-only facilities with an immediate jeopardy situation should be terminated by the State Medicaid Agency within the above time limits.

3012 - Termination Procedures – Substantial Noncompliance; No Immediate Jeopardy (Medicare)
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Deemed Providers/Suppliers:

See Chapter 5, Section 5110.3, in the case of a substantial allegation validation survey, or Section 5110.4c, in the case of a representative sample validation survey for the procedures to follow when an SA validation survey identifies substantial noncompliance in a deemed provider or supplier.

All Other Non-Long Term Care Providers/Suppliers

(See §§7310 - 7313 and §7400 for SNFs/NFs)

Failure to substantially meet one or more Conditions is a cause for termination of participation. “Substantially,” for purposes of this section, is defined as meeting the applicable CoPs or CfCs. Any provider/supplier that does not substantially meet the Conditions is considered to be limited in its capacity to furnish services at an adequate level or quality. Compliance with Conditions; i.e., condition-level deficiencies, can never be certified based upon a PoC or acceptable progress since the law specifically requires that all CoPs or CfCs must be met. If there is not an immediate jeopardy to patient health or safety, the RO and the SA use the following schedule:
1. Date of Survey - The date of the survey is the date on which the entire survey is completed regardless of when the exit conference is held.

2. Tenth Working Day – On the 10th working day, the SA sends a warning letter and the Form CMS 2567 containing the deficiencies to the provider/supplier and the RO. The SA informs the provider/supplier in writing that there is a determination of noncompliance and that it is recommending termination to be effective within 90 calendar days from the date of the survey. The recommended termination date is included in the letter. The SA informs the provider/supplier that the termination process provides an opportunity to make corrections and achieve compliance. This opportunity allows the provider/supplier ten calendar days to complete and return a plan of correction on the Form CMS 2567. The SA should state in the letter that it will make a revisit within 45 calendar days of the survey if a credible allegation of compliance is received. Termination takes effect as planned if compliance is not achieved. This notice serves as a warning letter to the provider or supplier. The SA allows the provider/supplier 10 calendar days to complete and return the plan of correction.

3. Forty-Fifth Calendar Day - If the facility has made a credible allegation of compliance (see §3016.A.), the SA conducts a revisit to determine whether compliance or acceptable progress has been achieved. Only 2 revisits are permitted; one within 45 calendar days and one between the 46th and 90th calendar days. If a second credible allegation of compliance is made prior to the effective date of termination, the SA telephones the RO and submits documentation to support the second revisit (only the second revisit is subject to RO approval). If the facility fails to make a credible allegation, no revisit is necessary.

4. Fifty-Fifth Calendar Day - If compliance has not been achieved, the SA certifies noncompliance. The SA forwards the certification and supporting documentation to the RO. The SA notifies the provider/supplier that termination is recommended and alerts the SMA if the provider/supplier is also participating in Medicaid.

5. Sixty-Fifth Calendar Day - Within 65 calendar days following the date of survey, the RO determines whether survey findings continue to support a determination of noncompliance.

6. Seventieth Calendar Day - The RO sends an official termination notice to the provider/supplier, the public, and the SMA if the provider/supplier also participates in Medicaid. Notices must be made at least 15 calendar days before the effective date of termination.
7. Ninetieth Calendar Day - Termination takes effect if compliance is not achieved. It can take effect in fewer than 90 calendar days if required procedures are completed.

NOTE: All timeframes are maximum. The RO may terminate more quickly as long as the regulatory requirements for notification of the public and provider/supplier are satisfied.

3012.1 - Termination of Psychiatric Hospitals (Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Deemed Psychiatric Hospitals

See Chapter 5, Section 5110.3 for the procedures to follow when a validation survey of a deemed status psychiatric hospital conducted by the SA identifies substantial noncompliance. For validation surveys conducted by CMS contract surveyors who assess compliance with the special psychiatric conditions (42 CFR 482.61 and 42 CFR 482.62 - the “B” tags), whether as part of a representative sample survey where the SA surveys for compliance with the regular hospital conditions (the “A” tags), or as a substantial allegation survey focused solely on the B tags, contract surveyors send their survey findings within 10 working days from the last day of survey to the CMS RO, not to SAs, with a copy to CMS CO. The CO reviews the survey findings for appropriateness and completeness of documentation and forwards them to the RO with CO’s recommendations for issuance of a determination of compliance or noncompliance. The RO makes the determination and notifies the hospital. If any of the regular hospital conditions (the A tags) or either of the two special psychiatric conditions, the B tags) is found not in substantial compliance, the 90 calendar-day termination procedures begin the date of the RO’s notice to the hospital with the survey report. The procedures to be followed after the RO issues its notice are the same as in Chapter 5, Section 5110.3, except that in some cases revisits would be conducted by the contract surveyors in addition to or instead of the SA.

Non-deemed Psychiatric Hospitals, Based on CMS Mental Health Surveyors’ Survey

The termination process for psychiatric hospitals using CMS mental health surveyors is consistent with the 90 calendar-day timeframe for other providers. However, due to the additional administrative process of sending the survey findings to the CO, day 1 of the 90 day termination timeframe begins on the date the RO receives the psychiatric survey report form findings from CO. The CMS mental health surveyors send the survey findings within 10 working days from the last day of survey to the CO, not to SAs and ROs. The CO reviews the survey findings for appropriateness and completeness of documentation and forwards them to the RO for final review and determination of compliance or noncompliance. If either of the two special psychiatric conditions is not in compliance (42 CFR 482.61 and 42 CFR 482.62), the 90 calendar-day termination procedures begin the day the RO receives the survey report.
Follow the termination procedures and timeframes below:

- **First Day** - Date of RO receipt of the survey findings from CO.

- **First – Tenth Working Day** - The RO reviews the survey report for adequacy of documentation to determine whether the documentation supports a finding of noncompliance with either psychiatric hospital requirement. (See 42 CFR 482.61 and 482.62.)

The RO notifies the CO via telephone if it does not concur with the CMS mental health surveyors’ findings regarding noncompliance with the psychiatric hospital requirements. Note that day 1 of the termination procedures begins the day the RO receives the completed psychiatric hospital survey report, not the day the RO reviewed the report for concurrence or nonconcurrence with the findings.

- **Tenth Working Day** - The RO notifies the provider of the cited deficiencies. The RO informs the provider in writing that a determination of noncompliance has been made and that termination will be effective 90 calendar days from the RO’s receipt of the survey report form (see Exhibit 180). Also the RO informs the provider that the termination process provides the opportunity to make corrections, and that if it reasonably believes that compliance has been achieved, it should notify the RO immediately. Explain that a revisit will be made within 45 calendar days from the RO’s receipt of the survey report form if a credible allegation of compliance is received. However, termination takes effect as planned if compliance is not achieved. This notice serves as a warning notice to the hospital, and it contains the proposed termination date. (The provider is to complete and return the POC to the RO within 10 calendar days.)

- **Forty-Fifth Calendar Day** - If the provider makes a credible allegation of compliance, the RO notifies CO and requests a revisit using the CMS mental health surveyors. The revisit to determine whether compliance has been achieved is to be conducted by the 45th calendar day. If a provider has not alleged compliance by the 45th calendar day, it is not precluded from making an initial credible allegation of compliance between the 46th and 90th calendar day.

- **Fifty-Fifth Calendar Day** - If a revisit has been made and compliance has not been achieved, the RO notifies the provider of the deficiencies that are not corrected and of any new deficiencies noted on the revisit.

- **Forty-Sixth - Ninetieth Calendar Day** - If the provider makes a credible allegation of compliance, the RO notifies the CO to schedule a second revisit to be conducted before the ninetieth day to determine whether compliance has been achieved.
• Seventieth Calendar Day - The RO sends an official termination notice to the hospital and a copy to the SMA if the provider also participates in the Medicaid program.

• Seventy-Fifth Calendar Day - The RO publishes the public notice.

Non-deemed Psychiatric Hospitals, Based on SA Surveys

• The procedures in Section 3012 are followed.

3012.2 - Termination of Organ Procurement Organizations (OPO)
(Rev. 112, Issued: 04-11-14, Effective, 04-11-14, Implementation: 04-11-14)

If an OPO voluntarily terminates its agreement, it must send a written notice to the CMS RO with the proposed effective date. The CMS RO approves the proposed termination date or sets a different date no later than 6 months after the proposed effective date or sets a date less than 6 months after the proposed effective date if it determines that a different date would prevent a disruption of services to the service area. When an OPO ceases organ procurement services in its service area, this will be considered a voluntary termination by the OPO. The CMS RO determines the effective date and notifies the OPO.

OPOs are involuntarily terminated when they fail to meet the requirements of certification at 42 CFR 486.303, including one or more conditions for coverage (CfC) or when they are not in substantial compliance with and any other applicable Federal regulations or provisions of titles XI, XVIII, or XIX of the Social Security Act (the Act).

The provider agreement between the OPO and the Secretary may be involuntarily terminated at anytime during the 4-year certification cycle or at the end of a cycle may not be renewed (re-certified) for another 4 year cycle. The CMS RO notifies CO, the FI/MAC and the Organ Procurement Transplantation Network (OPTN) that the OPO has been terminated and the effective date of the termination. The CMS RO follows the termination procedures detailed below.

3012.2A - Termination Procedures
(Rev. 112, Issued: 04-11-14, Effective, 04-11-14, Implementation: 04-11-14)

1. If the OPO is determined to be out of compliance with one or more CfCs, initiate termination of the provider agreement. CMS RO notifies the OPO of the pending termination date and the timeframes for submitting any additional information to support that appropriate corrections have been made and compliance achieved.

2. If the OPO achieves compliance, as determined through an onsite or desk review, send Model Letter: Organ Procurement Organization Approval. (See Exhibit 172.)
3. If the OPO fails to achieve compliance, send Model Letter: Organ Procurement Organization Notice of Termination. (See Exhibit 173.) The letter informs the OPO of:
   a. Reasons for the determination;
   b. The effective date of the determination;
   c. The deficiencies cited and the requirements not met;
   d. The OPOs right to seek reconsideration; and
   e. The timeframe for submitting additional data.

4. Open the service area for competition. (See §2812.3 and Exhibit 175.)

5. Publish a public notice in the newspapers.

6. Send copies of the public notice with a cover letter to:
   a. Association of Organ Procurement Organizations
      1364 Beverly Road, Suite 100
      McLean, VA 22101;
   b. Current Organ Procurement and Transplantation Network (OPTN)
      The current OPTN is:
      UNOS
      Post Office Box 2484
      Richmond, VA 23225;
   c. CMS Central Office
   d. Hospitals that have a working relationship and agreements with the OPO;
   e. Bordering OPOs;
   f. Medicaid/Medicare State Agencies; and
   g. FI/MAC of the terminated OPO.

3012.2B - Reconsideration Procedures
(Rev. 112, Issued: 04-11-14, Effective, 04-11-14, Implementation: 04-11-14)

The OPO may appeal the termination decision under 42 CFR 486.314.
The CMS RO mails notice to the OPO of a termination initial determination. The notice contains the reasons for the termination action, the effective date of the termination, and the OPO’s right to seek reconsideration. If the OPO is dissatisfied with the termination action, it has 15 business days from receipt of the notice of the termination action to submit its request in writing to the appropriate CMS RO to seek reconsideration of that determination from CMS. The request for reconsideration must state the issues or findings of fact with which the OPO disagrees and the reasons for the disagreement.

An OPO must seek reconsideration before it is entitled to seek a hearing before a hearing officer. If an OPO does not request reconsideration or its request is not made timely, the OPO has no right to further administrative review.

The CMS RO will make a written reconsidered determination within 10 business days of receipt of the request for reconsideration, affirming, reversing, or modifying the initial determination. An OPO dissatisfied with the CMS reconsideration decision, must file a request for a hearing before a CMS hearing officer within 40 business days of receipt of the notice of the reconsideration determination. If an OPO does not request a hearing or its request is not received timely, the OPO has no right to further administrative review.

3012.2C - Appeal Procedures
(Rev. 112, Issued: 04-11-14, Effective, 04-11-14, Implementation: 04-11-14)

If the OPO submits a request for a formal hearing, the CMS RO sends a complete administrative record that includes any additional materials submitted by the OPO, and a copy of the reconsideration decision and sends the supplemental administrative record to the CMS hearing officer.

CMS will provide the hearing officer with:

a) A copy of the CMS OPO Database report ranking of all OPO(s) utilizing the most recent data collection period, based upon compliance with the regulatory data requirements at §486.318 through §486.328.

b) Copies of all written correspondence between the OPO and the CMS RO relevant to the certification action under appeal;

c) All relevant e-mail correspondence between the OPO and the RO;

d) Any pertinent entries from a correspondence log if utilized; and

e) Relevant Survey and Certification Memoranda and guidance.
3012.3 – Termination of Organ Transplant Programs
(Rev. 190, Issued: 06-14-19, Effective: 06-14-19, Implementation: 06-14-19)

Transplant programs with one or more condition-level deficiencies other than 482.80 and 482.82 are placed on a 90 calendar-day termination track. However, if the program is found to be out of compliance with CoPs 482.80 or 482.82 and submits a request for reconsideration based on mitigating factors, the program will be given 210 days to come into compliance with these conditions. The 90 day termination track is enforced if the program does not come back into compliance regardless of whether the program is also on a 210 day termination track.

Please note that the termination of the transplant program’s Medicare approval does not affect the associated hospital’s provider agreement for participation as a Medicare-certified hospital. However, condition level findings at the hospital CoPs, may affect the hospitals provider agreement if corrections are not made timely.

HRSA and, if applicable the ESRD Network for a kidney program, are notified by the applicable RO of either a voluntary or involuntary termination of Medicare participation.

“The Heath Resources Administration (HRSA) (and if kidney program add ESRD Network) will be notified of this termination in order for them to provide assistance as indicated with potential recipient transfers to another Medicare-approved program.”

3014 - RO Termination Action Based on Onsite Survey of Medicare Provider or Supplier (Excluding SNFs) Conducted by RO Staff
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

When substantial noncompliance, including immediate jeopardy to patient health and safety, is determined by a survey team consisting of RO rather than SA surveyors, whether in the course of a Federal Look-behind survey, in response to a complaint, as part of the validation effort of a deemed provider or supplier, or to support other program needs, the RO initiates termination procedures as provided in §§3010 or 3012. The RO notifies the SA and the SMA of the action being taken. The RO completes the Certification and Transmittal, Form CMS-1539 in the Automated Survey Processing Environment (ASPEN) and ensures that the CMS National Data Base is updated to include the termination action. (See Chapter 7 for SNFs and NFs.)

3016 - Intervening Actions That Do Not Postpone or Delay Termination Timetable - (Includes Credible Allegations)
(Rev. 1, 05-21-04)

3016A - Credible Allegation of Compliance
(Rev. 1, 05-21-04)
A credible allegation is a statement or documentation:

- That is realistic in terms of the possibility of the corrective action being accomplished between the exit conference and the date of the allegation; and
- That indicates resolution of the problems.

If the provider/supplier makes an additional credible allegation that the deficiency(ies) is corrected following an earlier revisit or between the 46th and 90th calendar day prior to the effective date of termination, notify the RO by telephone. The SA submits all evidence or documentation regarding the facility’s allegation and its recommendation regarding the facility’s alleged compliance. The RO makes a determination whether a second revisit is appropriate.

The SA conducts a second revisit if one is approved by the RO. The SA forwards all supporting documentation, along with Form CMS-1539, certifying compliance/noncompliance to the RO immediately following the revisit.

Only compliance can stop a termination action.

3016B - Informal Hearings Do Not Interrupt Timetable
(Rev. 1, 05-21-04)

The process may not be postponed to accommodate informal hearings or meetings or to give the provider additional time to achieve compliance. Such discussion may, however, be conducted within the procedural time limits in §3012, as deemed appropriate by the RO. This 90 calendar-day procedure provides adequate time for the provider to achieve compliance if the decision by the RO is to wait the full time allowed and if the well being of patients is not jeopardized in the interim.

3016C - Acceleration of Timetable
(Rev. 1, 05-21-04)

The SA switches from the 90-day procedures in §3012 to the accelerated procedures in §3010 at any point when there is an immediate threat to patient health and safety.

3016D - Termination Development Coinciding With Change of Ownership (CHOW) Development
(Rev. 1, 05-21-04)

A CHOW does not affect completion of a termination action. The SA does not postpone any required termination, nor does it solicit a PoC from the new owner. Court appointed receivership is not a basis for cessation of the termination process. Following termination, the new owner may, however, request approval for participation as a new
provider, subject to reasonable assurance provisions (reasonable assurance only for Medicare). (See §2016.)

3016E - Disagreement over Deficiencies
(Rev. 1, 05-21-04)

(See also §7212 about Informal dispute Resolution for SNFs and NFs)

A provider that disagrees with any SA finding regarding a cited deficiency or an acceptable PoC should be advised to annotate its position on the PoC, and should specify why the SA’s citation is not correct. This information does not interrupt the termination process, but is publicly disclosable and is included in the documentation considered during subsequent reconsideration and hearings.

3016.1 - Provider Undergoes Chow During Termination Proceedings
(Rev. 1, 05-21-04)

If the SA learns that the provider is initiating a CHOW, it does not interrupt completion of its documentation of the certification of noncompliance. The SA continues to document noncompliance of the previous owner. The SA does not send a Form CMS-1539 to report the change.

3018 - Termination - SA Documentation Requirements
(Rev. 1, 05-21-04)

3018A - Documentation to Support Proposed Termination
(Rev. 1, 05-21-04)

All documents to support a proposed termination must be complete, accurate, and logical in sequence. Each document must be dated and signed by the preparer or indicate the date of receipt in the SA. The documentation must be supported by a complete current survey report or, in the case of an HHA, required CMS forms.

3018A1 - Current Survey Report
(Rev. 1, 05-21-04)

The SA reviews the current survey report or required forms to ensure that all items are properly completed. If there are any changes or erasures, the SA initials the item and explains the basis for the modification in the explanatory remarks column.

The SA includes the following information in the explanatory remarks column for each item “not met”:

- A description of the deficiency;
• Whether the deficiency existed during the previous survey and whether compliance was achieved, and then not sustained; and

• Current PoCs, if any.

In addition, the SA includes with the package an estimate of whether there is a prospect of compliance with all eligibility requirements within the time limits and the basis for this opinion.

3018A2 - Previous Survey Reports
(Rev. 1, 05-21-04)

The SA reviews previous survey reports for consistency. If a deficiency is reported on the current survey report that has obviously existed for some time, explain why it was not reported previously; e.g., serious structural defects, inadequate fire escapes.

The SA explains any conclusions that might be questioned, especially if certain requirements are being weighed heavily. For example:

• The majority of standards are “not met,” yet the Condition is found in compliance; or

• A Condition is found not in compliance based upon the relationship of standards or other deficiencies not being met.

3018B - Record of Contacts With Providers/Suppliers
(Rev. 1, 05-21-04)

The SA includes in documentation copies of communications and written reports of oral communications with providers/suppliers including the date of contact, the person involved, the purpose, and the content of the communication. Also, the SA includes reports of investigations of complaints.

3018C - Notification to Provider/Supplier of Deficiencies and Recommendation of Termination
(Rev. 1, 05-21-04)

The SA includes in the file a copy of the letter notifying the provider/supplier of the deficiencies found on the survey and advising that failure to correct will result in a recommendation for termination and includes copies of any other SA notices to the provider/supplier.
3020 - Additional SA Communications With Providers/Suppliers  
(Rev. 1, 05-21-04)

After the SA forwards the certification of noncompliance, it clears any further communications to the provider/supplier with the RO. Unrecorded visits, surveys, or correctional allegations that were not reported before final termination action could cause embarrassment or even result in failure to sustain the termination action. Even after final termination action, any additional contacts may be pertinent to proper handling of the case. The SA notifies the RO of any such contacts.

3022 - Notice of Termination (Medicare)  
(Rev. 1, 05-21-04)

The RO notifies the provider/supplier of its termination by letter at least 15 calendar days before the effective date of the termination. In the case of a hospital with an emergency department having deficiencies that pose an immediate jeopardy to the health or safety of individuals who present themselves to the hospital for emergency services, CMS gives the hospital a preliminary notice that its provider agreement will be terminated in 23 calendar days if it does not correct the identified deficiencies or refute the finding. CMS gives a final notice of termination, and concurrent notice to the public, at least 2, but not more than 4, calendar days before the effective date of termination of the provider agreement. For skilled nursing facilities (SNFs) and nursing facilities (NFs), CMS gives notice of termination, and concurrent notice to the public, at least 2 calendar days, one of which must be a working day, before the effective date of termination of the provider agreement, for a facility with immediate jeopardy circumstances, and at least 15 calendar days before the effective date of termination for a facility with nonimmediate jeopardy deficiencies. (42 CFR 488.456). The notice states the reasons for, and the effective date of, the termination and explains the extent to which services may continue after that date. The notice also contains information regarding the provider’s/supplier’s right to appeal the termination. (See 42 CFR 489.53) The only suppliers requiring public termination notices are RHCs (42 CFR 405.2404), ASCs (42 CFR 416.35), and FQHCs (42 CFR 405.2442). Public notices for other suppliers are optional at the discretion of the RO.

3024 - RO Termination Processing Sequence - Noncompliance With CoPs or Conditions for Coverage (Excluding SNFs)  
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Upon receipt of the SA’s unfavorable certification, the RO:

A. Establishes controls for processing the termination;

B. Performs an initial documentary review to make certain that copies of all pertinent surveys, statements of deficiencies, plans of correction (if submitted by the provider), and other necessary documents are included and that all relevant issues are resolved. When unable to determine the relationship of cited deficiencies to the quality of services or the health and safety of patients, the RO requests further
SA development. If necessary, the RO retains the file and phones the SA for the additional documentation needed;

C. Does a substantive review, resolves all substantive discrepancies and disputes, assesses the severity of the provider’s/supplier’s noncompliance, and makes its determination. The RO consults with LSC specialists in the RO, if necessary. See discussion in §3026 concerning how to treat key documents in making your determination;

D. Prepares the Termination Notice (Exhibits 181 and 182) and Newspaper Notice (Exhibit 183) and any supplemental press releases, if planned. The RO forwards a copy of its notice to the SMA, if appropriate; and

E. Inserts the effective date of termination in the notice and makes the necessary arrangements for public notice. To give both the provider and the public sufficient advance notice of termination of a provider’s agreement (at least 2 calendar days if there is immediate jeopardy or at least 15 calendar days if there is no immediate jeopardy), the RO determines the effective date of termination as follows:

- Allows sufficient time for delivery of the notice to the provider, depending on the provider’s location and the method of notification, i.e., letter, overnight mail, or electronic means.
- Determines the time needed for actual public notice by contacting the local newspaper or radio and television stations to determine their deadlines. (See §3034);
- Allows for receipt of the notice by the provider prior to publication of the public notice and assures that the public receives at least 2 calendar days if immediate jeopardy exists, otherwise 15 calendar days notice prior to the date of termination;
- Mails the termination notice to the provider (return receipt requested); and
- Notifies the SMA of action taken against Medicaid ICFs/IID and the effective dates if termination action is taken pursuant to §3000.C.3. When the termination action is taken, the RO mails the informational copies to the following offices:
  - Division of Medicare;
  - Division of Medicaid;
  - CO;
  - Intermediary;
o SA;
o SMA;
o Regional Director, Department of Health and Human Services (DHHS); and
o State Ombudsman.

3026 - Significance of Documentary Evidence in Determining Noncompliance
(Rev. 1, 05-21-04)

The RO uses the following documentation in determining compliance with the Medicare Federal CoPs or CFCs or Requirements of Participation.

3026A - Statement of Deficiencies
(Rev. 1, 05-21-04)

This statement constitutes evidence that the provider/supplier was notified of the specific deficiencies. These deficiencies are to be written as required by the Principles of Documentation. This assures that the statement provides accurate descriptions of the deficiencies and interpretations of Federal Medicare requirements that are not met. Otherwise, it might be alleged at a hearing that the termination action was based on error.

3026B - Plan of Correction (PoC)
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

When a PoC has been found to be unacceptable by the SA or RO, the PoC presents evidence that the provider or supplier is unable or unwilling to achieve compliance in a reasonable amount of time.

If a provider or supplier disagrees with a SA or RO finding of a cited deficiency, the provider or supplier may, in lieu of submitting a PoC, state on Form CMS-2567 the factual basis for disagreeing that a deficiency occurred. Whenever possible, the provider or supplier must reference the specific regulatory provision involved in the disputed issue and what factual evidence was available at the time of the survey to demonstrate compliance. It is not acceptable for the provider or supplier to provide evidence of corrective actions taken after the survey started as a basis for removal of a deficiency citation. It also is not acceptable for the provider or supplier to base its disagreement on a different interpretation of the regulatory requirements than that found in CMS guidance.

The original termination date is not changed by the provider’s or supplier’s disagreement with one or more of the deficiency citations. The RO reviews all of the documentation, including the survey findings and the documentation presented by the provider/supplier.
before making a determination. (If the RO determines that a deficiency did not exist, it is removed from Form CMS-2567.)

3026C - Revisit Reports and Subsequent Statements of Deficiencies (Rev. 1, 05-21-04)

This is the SA’s report of any revisit that was made to the provider/supplier following the survey that found the provider/supplier out of compliance.

3026D - SA Certification (Completed Certification and Transmittal (Form CMS-1539)) (Rev. 1, 05-21-04)

This is the SA’s certification as to whether the Medicare health and safety requirements were met at the time of survey. It also indicates that the SA completed the required actions and decisions.

3026E - Survey Reports (Rev. 1, 05-21-04)

Survey reports are the surveyor’s written records of findings during surveys that are primary evidence for the RO’s determination.

3026F - Documents of Collateral Evidence (Rev. 1, 05-21-04)

When obtainable, the RO adds such items as copies of pertinent provider records, correspondence, and State licensure information to the termination file to resolve or forestall conflicts of factual information elsewhere in the file and to support the adverse findings in the determination. Documentation can include verified complaint information.

3026G - Notice of Termination (Rev. 1, 05-21-04)

An adjudicative determination is consummated in an official notice of determination given to the parties whose rights are at issue. The determination becomes official when the notice (the formal termination letter) is mailed. It is essential that the notice be correct, not only in its procedural rendition, but also in the substance of the decision reported, since the receiving entity, as well as appellate authorities and courts, will treat it as the official “determination.”
3028 - Documentation Guide List - Termination for Noncompliance With §§1866(b)(2)(A) and (C) (Rev. 1, 05-21-04)

3028A - Documentation Appropriate to All Cases (Rev. 1, 05-21-04)

The RO uses the following to document recommendations for termination of a provider agreement pursuant to §§1866(b)(2)(A) and (C) of the Act:

- Copy of the letter to the provider advising it of the recommendation for termination (include the certified mail return receipt);
- Copy of Health Insurance Benefits Agreement, Form CMS-1561, filed by the provider and accepted for filing in the RO;
- Copy of the Letter of Acceptance of Agreement forwarded to the provider;
- All pertinent beneficiary complaints received;
- All pertinent violation reports made by the intermediary, Social Security office, etc.;
- Detailed reports of all pertinent intermediary efforts (i.e., dates, topic, reactions, results) and copies of related correspondence from the intermediary to the provider, including copies of any pertinent “provider letters” that may have been issued by the intermediary to the provider;
- Intermediary report on the current payment status of the provider, e.g., suspended, reduced, in current payment status. If the recommendation is based on failure to file annual cost reports, this item may be included as part of §3028.C;
- Detailed reports of all pertinent RO efforts (i.e., dates, topics, results, etc.) and copies of related correspondence from the RO to the provider; and
- All pertinent letters (or detailed records of visits or phone calls) received by the intermediary or by the RO from the provider.

3028B - Additional Documentation - Charging for Covered Services and/or Refusing to Refund Incorrect Collections (Rev. 188, Issued: 04-26-19, Effective: 04-26-19, Implementation: 04-26-19)

The term Medicare beneficiary identifier (Mbi) is a general term describing a beneficiary's Medicare identification number. For purposes of this manual, Medicare beneficiary identifier references both the Health Insurance Claim Number (HICN) and
the Medicare Beneficiary Identifier (MBI) during the new Medicare card transition period and after for certain business areas that will continue to use the HICN as part of their processes.

Additional documentation for the RO to use in making a determination of noncompliance includes:

- Name, address, Medicare beneficiary identifier, and dates of stay of any involved Medicare beneficiary known to have been furnished covered services by the provider;

- Where the services in question were furnished in a SNF, information relating to the beneficiary’s prior qualifying hospital stay, including if appropriate, copies of the Medicare billing submitted for the period of hospitalization;

- Copies of any bills, receipts, letters, that were received by the beneficiary from the provider requesting payment, including, if available, a description of the services furnished to assure that payment was requested for “covered services”;

- Copies of any checks, money orders, receipts, etc., which show payment to the provider by the beneficiary;

- Copies of any materials available which would show the payment conditions under which the beneficiary was furnished services, e.g., a contract of admittance;

- Copies of all pertinent “request for payment” forms that may have been filed by the provider for services furnished to the beneficiary during the period in question. If requests for payment were filed and the provider received program payment, the RO secures a written statement from the intermediary that shows that program payment (and the amount) was made to the provider on behalf of the beneficiary. For those cases where program payment has not been made (including cases where the provider has not filed a request for payment), secure a written opinion from the intermediary (based on available medical records and/or billings) as to the probability for making program payment;

- Copies of any requests by the beneficiary for the return of amounts paid to the provider for covered services, including any reply by the provider; and

- Authorization for the United States to act on behalf of the beneficiary. If the beneficiary has instituted any legal action to recover amounts paid to the provider, the RO does not secure the authorization. Use the following format:

Authorization
I hereby request that the United States act on my behalf to recover from (name of provider) amounts which I paid to said (hospital, nursing home, etc.) for services covered under title XVIII of the Social Security Act, popularly known as Medicare.

I hereby affirm that I am an eligible Medicare beneficiary and that as such I was a patient at said (hospital, etc.) from _______ to ____ (insert dates) and that I paid said (hospital, etc.) $____ (insert amount paid) for services rendered during this period.

Signed

Medicare beneficiary identifier

All of the information listed above should be obtained by the Division of Medicare as part of its responsibility in monitoring Medicare fiscal intermediaries.

3028C - Additional Documentation - Failure to File Cost Reports
(Rev. 1, 05-21-04)

The RO provides a listing of all pertinent health insurance checks, including check number, date, amount, disposition, drawn payable to the provider by the intermediary. (This information is to be shown under an intermediary letterhead and signed by a responsible person.) The RO includes a breakdown to show separately the period (“from to ____”) that related to these payments, including any offset against payments which may be due the provider, the total payment for the period involved, and the method of payment, e.g., periodic interim payments.

The RO reviews the data and material received from the intermediary and the additional documentation listed here and in subsection A, above, to avoid a duplication of development and documentation. If the case file shows that the provider has also failed to make satisfactory refund of overpayments, see subsection D.

3028D - Additional Documentation - Failure to Make Satisfactory Overpayment Arrangements
(Rev. 1, 05-21-04)

The RO reviews the data and material received from the intermediary and the documentation listed in subsection A to avoid a duplication of development and documentation. If the case file shows that the provider has also failed to file a cost report, see subsection C.

Section 1866(b)(2)(A) of the Act provides the authority to terminate the agreement of a provider that has failed to refund a substantial overpayment. For this purpose, when a provider has failed to make satisfactory arrangements for repayment, the sum of $1,000 or more may be viewed as a substantial overpayment. However, when considering
termination of an agreement, the RO takes into consideration the full circumstances of the particular case.

3028E - Additional Documentation - Admission Policies and Practices (Rev. 1, 05-21-04)

As provided in 42 CFR 489.53(a)(2), participation of a provider that voluntarily files an agreement to participate in the Medicare program, contemplates that the provider accepts beneficiaries for care and treatment. If a participating provider has any restrictions on the types of services it makes available and/or the type of health conditions that it accepts, or has any other criteria relating to the acceptance of persons for care and treatment, it is expected that such restrictions or criteria, if made applicable to Medicare beneficiaries, are applied in the same manner to all other persons seeking care and treatment. A provider’s admission policies and practices that are inconsistent with the provider agreement objectives set forth in this paragraph may be the basis for termination of participation by the Secretary pursuant to §1866(b)(2)(A) of the Act and 42 CFR 489.53(a)(2).

The amount of documentation for a provider’s failure to meet the above requirements is dictated by the circumstances of the particular case. Thus, the RO attempts to secure from all available sources (e.g., beneficiaries, providers, and intermediaries) any information that would be useful in making the determination.

3030 - Provider Agreement Terminations - Noncompliance with §§1866(b)(2)(A) and (C) (Rev. 1, 05-21-04)

3030A - Cause for Termination (Rev. 1, 05-21-04)

Under the provisions of §§1866(b)(2)(A) and (C) of the Act (also 42 CFR 489.53), the Secretary may terminate an agreement with a provider of services if it is determined that the provider:

- Is not complying substantially with the terms of the agreement, the provisions of title XVIII, or regulations promulgated thereunder;

- Has failed to supply information necessary to determine whether payments are or were due and the amounts of such payments;

- Refuses to permit examination of fiscal and other records (including medical records) necessary for the verification of information furnished as a basis for claiming payment under the Medicare program; or

- Refuses to permit photocopying of any records or other information necessary to determine or verify compliance with participation requirements.
3030B - Preparing Termination Cases  
(Rev. 1, 05-21-04)

The RO apprises the provider of its obligations and the consequences of continued violation before considering the need for terminating the agreement. Contacts with the provider may be made through the intermediary or by the RO staff, at the RO’s discretion. In corresponding with the provider, the RO uses certified mail with a return receipt requested.

The RO bases termination on documentation that supports a finding that the provider is not complying with the terms of the agreement or the provisions of title XVIII and implementing regulations.

3030C - Preliminary Notice to Provider  
(Rev. 1, 05-21-04)

The RO notifies the provider by letter that the findings and recommendations are being considered and that, if the findings are affirmed, the provider will receive official notice of termination of participation and the effective date on which the agreement is to be terminated. The RO advises the provider that when the official notice is released, it may be changed only if the determination is reversed upon appeal. Also the RO advises the provider to contact your office if it has taken steps to correct the violation or has definite plans for doing so.

3030D - Violation of §§1866(b)(2)(A) and (C)  
(Rev. 1, 05-21-04)

The RO notifies the provider that notice will be placed in the local newspaper(s) advising the public in accordance with the provisions of title XVIII. The RO explains the effect of the termination with respect to services furnished on or after the termination date, and advises the provider of the right to a hearing and the manner of filing for it. After the official notice of termination of participation is released to the provider, the RO proceeds with publication of the public notice. (See §§3034 and 3036.)

On the day before the public notice is published, the RO calls the intermediary and advises them of the termination. The RO cautions the intermediary not to divulge this information before the notice is published. Following publication, the RO formally notifies the SA.
3032 - Termination for Violations of §§1866(a)(1)(E), (F), (G), and (H) (Rev. 1, 05-21-04)

Under the provisions of §§1866(a)(1)(E), (F), (G), and (H) of the Act, the Secretary may terminate an agreement with a provider of services if it is determined that the provider:

- Failed to release data upon request to an organization having a QIO contract with the Secretary under Part B of title XI;

- Failed to maintain an agreement with the organization having a QIO contract with the Secretary under Part B of title XI;

- Has charged an individual for inpatient hospital services for which the individual was entitled to have payment made under Part A, pursuant to §§1886(b) and (d); or

- Failed to furnish all services (except for physician services as defined in §§1862(a)(14) and 1861(w)(1) of the Act) and items for which payment is made under §1866(a)(1)(G) of the Act.

3034 - Public Notice - Involuntary Termination (Rev. 1, 05-21-04)

The RO arranges to publish a notice in the local newspaper(s) announcing the termination and the reasons for termination of providers. Public notices are not required for suppliers, except in the case of RHCs (42 CFR 405.2404(d)), Ambulatory Surgical Centers (ASCs) (42 CFR 416.35(d)), and Federally Qualified Health Centers (FQHCs) (42 CFR 405.2442). The RO may place public notices for the termination of other suppliers at its discretion.

3034A - For Providers (Rev. 1, 05-21-04)

MEDICARE NOTICE TO THE PUBLIC:

Notice is hereby given that the agreement between the (facility/address), and the Secretary of Health and Human Services, as a provider of services in the Health Insurance for the Aged and Disabled Program (Medicare) is to be terminated at the close of (date of termination).
3034B - For Suppliers
(Rev. 1, 05-21-04)

MEDICARE NOTICE TO THE PUBLIC:

Notice is hereby given that effective at the close of (date of termination) the (name and address of facility) is no longer approved for participation in the Medicare program as a supplier of (type of services).

The RO inserts the reasons for termination in the format of the following example:

The (name of provider) does not comply with the Medicare (Condition of Participation/Requirements of Participation/Condition for Coverage) pertaining to (Condition(s) or Requirement(s) (for SNFs) out of compliance).

Add one of the paragraphs below, depending on the type of provider being terminated:

3034C - For Hospitals, CAHs, and SNFs
(Rev. 1, 05-21-04)

The Medicare program will not make payment for inpatient hospital services (or skilled nursing facility services) furnished to patients who are admitted after the close of (date of termination). For patients admitted on (date of termination), or earlier, payment may continue for up to 30 calendar days of inpatient hospital services (or skilled nursing facility services) furnished after (date of termination).

3034D - For HHAs and Hospices
(Rev. 1, 05-21-04)

The Medicare program will not make payment for (home health) (hospice) services furnished to patients whose plan of treatment was established after the close of (date of termination). For patients whose plan of treatment was established before (date of termination), payment may be made for up to 30 calendar days after date of termination.

3034E - For Other Providers and Suppliers
(Rev. 1, 05-21-04)

The Medicare program will not make payment for (type of facility) services furnished to patients after the close of (date of termination).

Arrange for publication of notice at least 15 calendar days before the effective date of termination (2 calendar days if immediate jeopardy), but not before the provider or supplier receives notice of termination. The notice advises the
provider/supplier of it’s right to due process, the expected schedule for termination action, and that the deficiency must be corrected and verified by the SA to halt the termination.

In addition to preparation of legal advertisements, the RO may decide to develop and distribute a press release (via radio/television) on the termination (or other action).

3036 - Billing for Public Notice of Termination or Withdrawal
(Rev. 1, 05-21-04)

3036A - Advertising Order, SF-1143
(Rev. 1, 05-21-04)

Before arranging for publication of the termination notice, the RO/ARA obtains authorization from the RA via the Advertising Order, SF-1143 (Exhibit 184). The RO prepares an original and three copies, completing the front portion as shown in the exhibit. The RO attaches a copy of the text of the public notice to the original and to each copy of the SF-1143. After the RA signs the form (and retains one copy) the RO:

- Types in the RO address in the bottom portion of the front of the SF-1143;
- Sends the original and one copy to the newspaper. Informs the publisher that one of the following options should be used in claiming payment:
  - The back of the original SF-1143 may be completed with either a copy of the advertisement attached or a certification made in the space provided on the form; or
  - Any billing form may be submitted to the RA with a copy of the advertisement or a certification attached to the original SF-1143; and
- Following publication of the public notice and upon receipt of the SF-1143 and attachments from the publisher, issues payment using funds available for this purpose.

3038 - Rescinding or Postponing Effective Date of Termination
(Rev. 1, 05-21-04)

3038A - Initial Action
(Rev. 1, 05-21-04)

The RO stops the processing of an involuntary termination if it is positively ascertained that the provider now complies with all requirements and that termination is no longer appropriate. The RO does not postpone a termination action for:

- Meetings or visits with the provider;
• CHOWs;
• Court appointed receivership; or
• Any other cause not explicitly required by statute, regulation, Federal court order, or procedures stated elsewhere in this chapter.

When credible evidence that the cause for the termination has been removed is received in time to halt publication of the public notice, the RO immediately contacts the newspaper to stop the scheduled notice and suspends the termination action pending RO and SA verification. If the termination action is later revived, the RO again makes arrangements for public notice.

When the RO receives credible evidence that the cause for the termination has been removed before the effective date of termination, it requests that the SA make a revisit as soon as possible or it conducts a Federal survey. The SA should make the revisit when:

• The SA conducted the initial survey; or

• Due to travel considerations, scheduling problems, or scarcity of resources, the RO has requested the SA to conduct the revisit, even though the RO conducted the initial survey.

The RO does not use allegations of compliance in the absence of credible evidence of compliance as the basis for postponing the effective date of termination.

3038B - Criteria for Credible Allegation
(Rev. 1, 05-21-04)

A credible allegation must meet the following criteria:

• The alleged corrective action should be detailed and must have removed the deficiency; and

• The corrective action was of the kind that could have been accomplished between the survey date and the date of the allegation.

If the Medicare facility makes additional credible allegations of compliance following an earlier SA revisit, or between the 46th and 90th calendar day prior to the effective date of termination, the SA will contact the RO for a determination as to whether a revisit will be made. Only two revisits are permitted, one before the 45-day period and one between the 46th and 90th day. The RO bases its determination on the validity and reasonableness of the evidence or documentation provided by the facility prior to conducting a revisit. Except in the case of SNFs and NFs, the SA can conduct a second or late revisit only after RO approval. (See §7317 for revisit policies for nursing homes.)
Notice is hereby given that (name and address of facility) has achieved compliance with the Medicare Conditions of Participation/Requirements for Participation pertaining to (CoPs or Requirements). As a result, the Secretary of Health and Human Services is continuing the agreement with (the facility) in the Medicare program.

Notice is hereby given that (name and address of facility) has achieved compliance with the Medicare Conditions for Coverage pertaining to (Conditions of Coverage). As a result (the facility) will continue to participate in the Medicare program as a supplier of (type of supplier services).

Section 1910(b)(1) of the Act authorizes CMS to terminate approval of a Medicaid ICF/IID’s eligibility to participate in the Medicaid program when CMS determines that the provider does not substantially meet the CoPs for ICFs/IID (42 CFR Part 483). The Act uses the terms “cancel” and “terminate” interchangeably. The adjudicative procedures are similar to those followed for terminating a §1866 provider agreement under §§3010 and 3012. Except in the case of immediate jeopardy situations, termination usually becomes effective after the provider has had an evidentiary hearing before an Administrative Law Judge (ALJ) and the ALJ has upheld the termination.

At the exit conference, the RO team leader should explain the findings to facility management as well as which findings apparently constitute an immediate jeopardy to client health and safety. (See Appendix Q for examples.) The RO survey team leader explains that if his/her supervisor agrees with the seriousness of the team’s onsite survey findings, the RO will notify the facility by electronic facsimile, telegram, or overnight express mail of the determination to cancel/terminate the facility’s program participation unless the immediate threat is eliminated. The RO allows no more than two working
days following the exit conference to determine whether circumstances found in the facility pose an immediate jeopardy to client health and safety and to notify the facility of your determination. The RO notifies the facility and SMA by electronic facsimile, telegram (Exhibit 185), or other expeditious means, that as a result of the finding that an immediate jeopardy to client health and safety exists, the RO is initiating termination proceedings pursuant to §1910(b)(1). The RO telephones the SA to inform them of its determination. The RO gives the facility no more than 5 working days from the date of the notification to eliminate the threat and to notify it of the remedial action taken. Also, the RO should state that if the facility does not notify the RO that the threat has been removed or compliance has been achieved by the time specified, it will assume the condition still exists and termination occurs on the proposed effective date.

If the threat is removed but deficiencies still exist at the Condition level, the RO uses the procedures for no immediate threat and gives the provider 90 calendar days from the date of survey to correct the deficiencies.

Following CMS termination based on immediate jeopardy, CMS grants up to 30 calendar days of Federal Financial Participation (FFP) for purposes of relocating patients after the effective date of termination. If an appeal is filed by the facility, the ALJ hearing is afforded after the effective date of termination and does not forestall termination from taking effect.

3040A2 - No Immediate Threat to Patients’ Health and Safety
(Rev.92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

If the ICF/IID is not in compliance with one or more of the CoPs, the RO completes the actions within the time limits prescribed.

a. Tenth Working Day - The RO notifies the provider in writing of its deficiencies and that termination action is being initiated. Included in the notices is the provider’s right to appeal this action, along with the effective date of termination. If the provider makes a credible allegation of compliance prior to the effective date of termination, the RO conducts a revisit. (See §3038.B.)
b. Seventieth Calendar Day - The RO completes all related documentation and notifies the facility, the SMA, and the SA.
c. Ninetieth Calendar Day - If compliance has not been achieved, the RO terminates participation.

If an appeal is filed by the facility, termination must be delayed pending the hearing and decision by the ALJ (see 42 CFR 498.5(j). The provider agreement remains in effect and FFP continues pending the appeal decision. If the facility makes a credible allegation of compliance during the appeal period, it is the RO’s decision whether or not it is in the recipients’ and the Federal Government’s best interests to conduct a revisit and dispose of the case based on the findings. If a revisit is made and the facility failed to achieve compliance, adverse action continues based on the findings of the first Federal survey and
the findings of the revisit. If at the time of the revisit the provider is in compliance with
the requirements forming the basis for the original termination, but has new deficiencies
that are also grounds for termination, the RO initiates a new termination process
commencing with the revisit.

If the ALJ sustains the termination action, the effective date of termination and cessation
of FFP is set by the ALJ. Further appeal by the facility to the DAB does not cause the
provider agreement to be extended, i.e., payment does not continue pending a decision by
the DAB.

Whenever possible, the RO conducts the termination notification and decision-making
process in the 90-day timeframe used for Medicare terminations. However, there are
circumstances that require that the RO give a facility extra time. For example, with
State-owned facilities, it sometimes takes longer to get a PoC because of the need for
action by other parts of State government, thus requiring additional processing time.
Keep these situations to an absolute minimum. In addition, an ALJ hearing may not be
scheduled within the usual 90-day termination timeframe.

The reasonable assurance provisions apply to ICFs/IID terminated by CMS.

3042 - Disallowance of FFP to State Because State Fails to Follow Correct
Certification Procedures for Medicaid Providers
(Rev. 1, 05-21-04)

This process applies when a SA or SMA has failed to properly apply Federal
requirements or procedures in surveying or certifying a facility or in entering into a
provider agreement, as described in 42 CFR 442.30 and 431.610(g).

The “look behind” regulation (42 CFR 442.30) referred to as “old look behind” provides
a mechanism for testing the validity of a provider agreement for FFP purposes. If
certification procedures are not followed, CMS does not accept the provider agreement
between the State and the facility as evidence that the facility was properly certified.
Whether a facility actually meets participation requirements is not the issue. The issue is
that the State failed to adhere to Federal requirements in certifying and issuing an
agreement to a facility. Therefore, CMS will discontinue FFP to the State for the facility,
even though payments were made prior to the RO determination. When the RO invokes
this aspect of “look behind,” it advises the SA and the SMA that the provider agreement
cannot be accepted, and, for FFP purposes, the agreement is void from its inception.

If the SA failed to adhere to certification procedures, the RO requests additional
information. The RO advises the agency that in the absence of a satisfactory response, the
provider agreement is invalid for FFP purposes that will result in a disallowance of FFP
for the period in question. The RO requests an explanation or evidence that would
contravene a finding that Federal requirements were not followed. The nature of the
issue (i.e., a defect in the survey versus a defect in the issuance of the provider
agreement) determines whether the RO notifies the SA or the SMA. Whichever agency
is notified, the RO sends copies of all correspondence to both agencies.

If unable to correct the difficulty through contact with the State, the RO contacts the
Division of Medicaid regarding the problem. The RO recommends the disallowance of
FFP for the State under the current provider agreement. A sample memorandum to the
Medicaid Division is in Exhibit 186.

3044 - Terminating Approval for Suppliers
(Rev. 1, 05-21-04)

If the SA certifies that a supplier is no longer in compliance with the Conditions for
Coverage, the RO notifies the supplier of the certification and processes the termination.
(See Exhibit 187.)

In supplier terminations, although there may not be a provider agreement to terminate and
the Act does not use the term, “terminate,” formal adjudicative disapproval is clearly
implied in the Act. Unless otherwise noted, procedures for provider terminations are
equally applicable to certified suppliers. SMAs are notified of supplier terminations. If
the SMA continues making payments to Medicaid suppliers, FFP is disallowed. (See
Chapter 6 for laboratories.)

The termination of coverage is effective following at least a 2 calendar-day notice if
immediate jeopardy is present, otherwise it is at least a 15 calendar-day notice to the
supplier (42 CFR 489.53(c)).

The RO notifies the carrier and the Divisions of Medicare and Medicaid of a supplier
termination action and the effective date.

Public notification is optional for suppliers other than RHCs, ASCs, FQHCs, and OPOs
(42 CFR 486.325(e)). Public notification, when undertaken, should be given in
accordance with §3034.

3046 - Voluntary Terminations
(Rev. 1, 05-21-04)

3046A - General
(Rev. 1, 05-21-04)

Under the provisions of §1866(b)(1) of the Act, a provider of services may terminate its
agreement by filing a written notice of its intention. If a Medicare provider/supplier
notifies the SA of its desire to terminate its Medicare participation or if it ceases
operations which is considered as voluntarily terminating its agreement, the SA notifies
the RO immediately. The RO accepts the proposed termination date or set a different
date. However, the termination date must not be more than 6 months from the date the
notice is filed.
The RO determines the provider’s or supplier’s reason(s) for deciding to terminate participation. Identifying the reasons for voluntary termination aids in evaluating policies and procedures and focuses on problems not previously recognized.

1 - Provider or Supplier Is Unable or Unwilling to Correct Deficiencies to Continue to Meet CoPs, Conditions for Coverage, or Participation Requirements for SNFs

In many cases, the facilities have cited as a reason for seeking termination an inability to continue to meet the Conditions of Participation or Coverage or Requirements for participation for SNFs.

2 - Provider Dissatisfied With Reimbursement

If a provider is withdrawing because of disagreement with the reimbursement formula, the RO indicates this on Form CMS-1539.

3 - CHOW

If, after a CHOW, the successor does not wish to participate, the date of termination is usually the date the previous owner ceased doing business. However, coverage of beneficiary services extends until it is learned that the successor will not continue operations under a provider agreement. Payment can continue for up to 30 calendar days after a provider is terminated for hospitals, SNFs, HHAs and hospice beneficiaries who were admitted before the effective date of termination (42 CFR 489.55). (See §3008.)

4 - Close of Business

The provider may temporarily or permanently cease all business (Medicare and non-Medicare operations). No further RO action is necessary.

3046B - Decision by Provider or Supplier to Remain in the Medicare Program
(Rev. 1, 05-21-04)

If a provider or supplier changes its mind after requesting termination, the RO secures a written statement to document the provider/supplier file to prevent any future misunderstanding. If the voluntary termination has not already taken place, the RO sends a letter to the provider rescinding its voluntary termination. Copies are sent to the SA, SMA, the intermediary, and the carrier. If the provider’s request is received after the effective date of the voluntary termination, the RO treats the request as an initial request to participate in the Medicare program.
3046C - Notice to Public  
(Rev. 1, 05-21-04)

In voluntary termination cases, the provider or supplier is obligated to notify the public of the effective termination date. An exception to the requirement for public notice is made when the RO receives retroactive notice of the close of a business. If the RO learns that the provider does not intend to comply with the requirement for a public notice, where required, the RO should assume the responsibility. The required public notice should be published in the local newspaper with the widest circulation as soon as possible after the provider receives the RO’s letter, and, if time permits, not less than 15 calendar days before the effective termination date. When a supplier of services is voluntarily terminating program participation, public notice by either the supplier or the RO’s office is optional. However, such a notice is to be published for RHCs, ASCs, and FQHCs.

3046D - Effective Date of Voluntary Termination  
(Rev. 1, 05-21-04)

The effective date of termination is the date business ceased (if there is closure) and should allow sufficient lead-time to notify CMS components and to give the public notice of the termination. If the provider’s request does not specify an acceptable termination date, the RO sets the date (42 CFR 489.52(b)). This date cannot be more than 6 months after the provider’s request is dated. If a retroactive termination date is requested, the RO honors it, provided there were no Medicare beneficiaries receiving services from the facility on or after the requested termination date.

In setting an effective termination date that is less than 6 months in the future, the RO must be assured that it would not unduly disrupt the services to the community or otherwise interfere with the effective and efficient administration of the health insurance program. In making this determination, the RO considers the availability of other facilities in the area. In the case of a closure, the effective date is the actual date of closing.

3047 - Notice to Intermediary or Carrier - Voluntary Termination  
(Rev. 1, 05-21-04)

The RO notifies the intermediary when it receives notice that a provider wants to terminate its participation. The RO keeps the intermediary informed of the status of the provider’s request. This permits the intermediary to make preliminary arrangements for final cost reports and final settlement, and to adjust any outstanding payments to avoid overpayments in accordance with intermediary instructions. If a supplier is voluntarily terminated, the RO notifies the carrier immediately.

3048 - Notice to Provider or Supplier - Voluntary Termination  
(Rev. 1, 05-21-04)
3048A - Voluntary Termination
(Rev. 1, 05-21-04)

Exhibits 188 and 189 may be used to notify the provider or supplier of approval of voluntary termination. The RO sends copies of the letter to the SA, intermediary or carrier, and, if appropriate, the SMA.

3048B - Close of Business
(Rev. 1, 05-21-04)

The RO sends a letter modeled after Exhibit 190 to the provider with copies to the SA, intermediary, and, if appropriate, the SMA. If a supplier is closing, the RO notifies the carrier and sends a letter modeled after Exhibit 191.

3049 - Completing Certification and Transmittal (Form CMS-1539)
(Rev. 1, 05-21-04)

3049A - Voluntary Terminations and Close of Business
(Rev. 1, 05-21-04)

In the upper left-hand corner The RO enters “VOLUNTARY TERMINATION--CODE 2.”

The RO completes items 1, 3, 7, 26, 28, 29, 30 (signature of RO official delegated to sign), and 32.
Reconsideration, Hearings, and Appeals

3050 - Initial Determinations Versus Administrative Actions - Right to Review
(Rev. 1, 05-21-04)

Only initial determinations are subject to reconsideration, hearing, or appeal. In general, an initial determination is a decision with respect to the following matters:

- Whether a provider or prospective provider meets or does not meet the Medicare requirements as a provider of services;

- Whether a supplier or prospective supplier meets or does not meet the appropriate Conditions for Coverage of its services;

- Whether the termination of a provider agreement or benefits agreement is in accordance with 42 CFR 489.53, the termination of a RHC agreement is in accordance with 42 CFR 405.2404, the termination of a FQHC agreement is in accordance with 42 CFR 405.2442, or the termination of an ASC is in accordance with 42 CFR 416.35;

- Whether a hospital meets or does not meet or continues to meet the requirements to qualify as an emergency services hospital;

- Whether the services of a supplier meet or continue to meet the Conditions for Coverage; or

- The effective date of the provider agreement between CMS and a provider of services, or the effective date CMS approved for a supplier of services.

3052 - Nature of Reconsideration Determination - SA Procedures
(Rev. 1, 05-21-04)

3052A - Right to Reconsideration of Initial Denial
(Rev. 1, 05-21-04)

Reconsideration is granted administratively, not statutorily, pursuant to regulations 42 CFR 498.22 through 498.25. Any prospective provider or supplier dissatisfied with an initial determination that does not qualify as a Medicare provider may submit a request that the Secretary reconsider the decision within 60 calendar days from receipt of the notice of initial determination.

Reconsideration is a review of the determination. This review results in affirmation or reversal or the determination. Further appeal rights include hearing before an ALJ and review by the DAB.
3052B - Request for Reconsideration  
(Rev. 1, 05-21-04)

A request for reconsideration is any written expression of dissatisfaction with the initial decision. The request may be in the form of a letter or statement that explains the issues, or the findings of fact with which the affected party disagrees, and the reasons for the disagreement. The reconsideration request may be signed by any responsible official of the provider or by an attorney on behalf of the provider. The SA officially dates or date-stamps any request the day of receipt in the SA.

3052C - Acknowledgment of Reconsideration Request  
(Rev. 1, 05-21-04)

The SA acknowledges the request promptly and forwards a copy of the request and acknowledgment letter to the RO immediately. The RO will advise if additional development is required. Also, the SA forwards any subsequent information received that would affect the reconsideration or hearing. If an attorney files the request, the SA sends a copy of the acknowledgment to the provider. Most cases require SA redevelopment, particularly if there are questions about the provider’s efforts and plans to correct previously cited deficiencies. If requesting additional evidence from the entity, stipulate in the acknowledgment a reasonable deadline for submittal.

3052D - Documentation of File  
(Rev. 1, 05-21-04)

A reconsideration review is not complete unless the file contains adequate documentation to fully explain every statutory deficiency and finding of noncompliance with program requirements. The SA sends the RO all reports of on-site visits and telephone contacts with the provider, as well as any pertinent information available from the licensing agency.

3054 - Reconsideration - RO Procedures – Excluding SNFs and NFs  
(Rev. 1, 05-21-04)

3054A - Review  
(Rev. 1, 05-21-04)

A reconsideration is a thorough, independent review of the prior decision and entire body of evidence, including any new information developed. If the provider/supplier has made corrections since the survey on which the original determination was based and the SA can verify this, the provider/supplier may no longer wish to pursue its recommendation request. For initial certifications only, the potential provider/supplier may withdraw its reconsideration request, ask for another initial certification (as long as the timeframe for the Form CMS-855 application has not expired) and accept certification based on the date of compliance with all Medicare conditions.
A reconsideration review is not complete unless and until every adverse finding (i.e., does not meet one or more statutory requirement, is not in substantial compliance with one or more CoPs) is adequately documented.

The reconsideration process for nursing homes was eliminated in 1995 when Informal Dispute Resolution (IDR) was formalized as part of the appeals process for SNFs and NFs. IDR duplicates in every pertinent way the reconsideration element of this process since it provides the opportunity for nursing home providers to conduct survey findings information (see §7212).

3054B - RO Receipt of Request  
(Rev. 1, 05-21-04)

A provider or supplier of services which has been denied participation may file a request for reconsideration in any manner through any CMS RO, SA, or intermediary. The request may be in the form of a letter or statement and may be signed by any responsible official of the provider/supplier, or by an attorney on behalf of the provider. The RO officially dates or date-stamps any request the day of receipt in the RO. The request for reconsideration must state the issues or the findings of fact with which the affected party disagrees, and the reasons for disagreement.

3054C - Acknowledgment or Reconsideration Request  
(Rev. 1, 05-21-04)

The RO acknowledges reconsideration requests within 3 working days of receipt. If an attorney filed the request, the RO sends a copy of the acknowledgment to the provider/supplier. If there is an offer to submit additional evidence or if the RO requests additional information, a reasonable deadline for its submittal is provided in the acknowledgment. The RO informs the provider/supplier that the SA may be in touch to obtain additional information. The RO informs the SA that the provider/supplier has requested reconsideration.

3054D - Reconsideration Determination  
(Rev. 1, 05-21-04)

The RO completes a Certification and Transmittal, Form CMS-1539, except for Items 17 and 18 if the original determination is reversed. The RO marks the top of all copies of the form “Reconsideration - (Affirmed) or (Reversed)” and distribute the copies as in an initial decision. The RO does not need to complete Form CMS-1539 if the SA has made a revisit and the certification is based on the revisit. The SA in this case will complete Form CMS-1539. If OCR clearance has not yet been received, the RO advises OCR that a previously denied provider/supplier has now been approved so that they may complete the clearance process. The cover letter should request immediate return of the forms and point out that a final determination has not been made on the request for reconsideration. If any title VI forms are needed, they are sent with the agreement forms.
3054E - RO Notice of Reconsidered Determination  
(Rev. 1, 05-21-04)

3054E1 - Denial Reversal (Approval)  
(Rev. 1, 05-21-04)

The RO confirms civil rights compliance before revising a denial of a provider on reconsideration. If the provider is not in compliance, see §2010.

After confirming that documentation is in order, the RO assigns a provider identification number and completes a Certification and Transmittal, Form CMS-1539, marking item 30, “Reconsideration Reversed.”

The RO issues a notice of acceptance as in a routine initial approval, enclosing the countersigned provider agreement. The notice of acceptance should reflect that the determination was reconsidered.

The RO notifies the SA of the revised decision and sends a Provider Tie-In Notice, Form CMS-2007, to the intermediary.

3054E2 - Denial Affirmed  
(Rev. 1, 05-21-04)

With the notice of decision, the RO includes a listing of each CoP, Condition for Coverage, or Requirement for SNFs with which the provider/supplier is not in compliance (for SNFs, substantial compliance). The RO includes a detailed explanation of why the deficiencies result in a determination and decision that the provider/supplier is not in compliance (substantial compliance for SNFs), the provider’s efforts and plans to correct deficiencies notwithstanding. The notice advises the provider/supplier of its rights to a hearing if it files a request within 60 calendar days of the date of receipt of the reconsideration decision of denial. The RO advises the provider/supplier to send the hearing request to it, Attn.: ARA, CMS. The RO forward all requests for hearings to the DAB for further action. The RO keeps a copy of the notice in the provider file and sends a copy to the SA. The RO acknowledges the receipt of the provider/supplier’s request for hearing in writing (see Exhibit 192).

3054E3 - Acting Official for Reconsideration Denial Notices  
(Rev. 1, 05-21-04)

If a reconsideration request is denied, the reviewer who was assigned the initial determination should not sign the denial notices at the reconsideration level.

- Initial Denial NOT Signed By ARA - If the ARA did not sign the initial denial notice he/she should sign the reconsideration denial notice.
• Initial Denial Signed By ARA - If the ARA signed the original notification of denial, forward the file and formal recommendation to the RA. The notice of reconsideration denial is released over the RA’s signature. The RO notifies all interested components and prepares and transmits the required Form CMS-1539.

• Problem Cases Needing CO Review - If the reconsideration action presents policy issues or problems that in the RO’s judgment need resolution, send the case to the CO with a statement of the problem and request guidance.

The RO prepares and transmits Form CMS-1539 and adjudicative notices to the SMA, SA, and intermediary.

3058 - Hearing on §1910(b) Cancellation of Medicaid Eligibility
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

If a hearing is requested on a termination of an ICF/IID’s Medicaid participation by CMS, send the provider’s request for a hearing to the DAB.

3060 - Appeals of Adverse Actions for Medicaid Non-State Operated NFs (Non-State Operated) and ICFs/IID (Not Applicable to Federal Terminations of Medicaid Facilities)
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

Denials, terminations, cancellations, and denials of payment for new admissions and other adverse actions to facilities participating in Medicaid-only are State administrative actions and decisions. State appeal procedures must be made available to facilities in cases of nonrenewal, denial, cancellation, or termination of the provider agreement. It is up to the State to designate the office or official having authority to hear and decide Medicaid appeals. Although the State retains considerable flexibility in developing its own appeal procedures, the procedures for an ICF/IID must at a minimum provide for an evidentiary hearing either before or within 120 calendar days after the effective date of the adverse action. The State must also provide an informal reconsideration prior to taking adverse action if it elects to provide a full evidentiary hearing after the effective date of the adverse action for an ICF/IID (42 CFR 431.150 through 431.154).

If a NF requests a hearing on a denial or termination of its provider agreement, the request does not delay the adverse action and the hearing need not be completed before the effective date of the action. However, a NF is entitled to a hearing before a CMP is collected (see §7526).

NOTE - In the procedures for denial of payment for new admissions for ICFs/IID (see §3060.C), a post-termination hearing is not a permitted option. The State must provide an informal hearing before the effective date of the denial of payments for new admissions. Consequently, reconsideration is not appropriate for these appeals.
3060A - Informal Reconsideration  
(Rev. 1, 05-21-04)

The State may develop and implement its own reconsideration proceedings. However, the process must include:

- Timely notice of the reason for the action;
- A reasonable opportunity for the provider/supplier to refute those reasons in writing; and
- A written decision prior to the effective date of the adverse action.

3060B - Evidentiary Hearing  
(Rev. 1, 05-21-04)

The evidentiary hearing must include:

- Timely written notice to the provider/supplier of the findings upon which the termination or denial is based, and disclosure of the evidence on which the decision is taken;
- An opportunity for the provider/supplier to appear before an impartial decision maker to refute the basis for the decision;
- An opportunity for the provider/supplier to be represented by counsel or another representative;
- An opportunity for the provider/supplier or its representatives to be heard in person, to call witnesses, and to present documentary evidence;
- An opportunity for the provider/supplier to cross-examine witnesses; and
- A written decision by an impartial decision-maker, setting forth the reasons for the decision and the evidence on which the decision is based.

3060C - Informal Reconsideration (Applies to ICFs/IID for Denial of Payment for New Admissions Only)  
(Rev. 92, Issued, 11-22-13, Effective: 11-22-13, Implementation: 11-22-13)

The informal hearing process must include:

- Timely notice of the reason for the action;
• A reasonable opportunity for the provider/supplier to present in writing or in person reasons for its disagreement;

• An opportunity for the provider/supplier or its representatives to be heard in person and to present documentation; and

• A written decision by an impartial decision maker, prior to the effective date of the denial of payment, setting forth the reasons for the determination. An evidentiary hearing does not follow the informal hearing.

3060D - Judicial Review
(Rev. 1, 05-21-04)

Federal regulations do not provide for judicial review of these appeals proceedings. State law governs judicial review.

3060E - Impartial Decision Maker (Hearing Officer)
(Rev. 1, 05-21-04)

The State has flexibility in selecting individuals to conduct the reconsideration and hearing proceedings. However, in both proceedings, certain individuals should be excluded from serving as decision-makers.

In reconsideration proceedings, the SA, as well as other persons directly involved in gathering and providing evidence upon which the adverse action is based, is ineligible to make decisions. (One person should not be both witness and judge.) However, the person who made the original determination based on the surveyors’ findings is not ineligible to decide the reconsideration. If the decision is originally made at the highest level, the appeal decision should also be made there. However, if a regional supervisor makes the original decision, have someone higher in authority review the appeal.

In administrative hearings, all persons directly involved in either the survey or the reconsideration process are ineligible for reasons of impartiality.
Prospective Payment System (PPS)

3100 - Hospitals and Hospital Units Excluded From the Inpatient Prospective Payment System (IPPS) - Introduction
(Rev. 198, Issued: 01-17-20, Effective: 01-17-20, Implementation: 01-17-20)

The IPPS determines Medicare payment for operating costs and capital-related costs of inpatient hospital services provided in short-term acute care hospitals. Certain hospitals and special hospital units may be excluded from this system and paid at a different Medicare reimbursement rate. These IPPS-excluded hospitals and units have their own specific reimbursement criteria. Title 42 CFR 412.20 through 412.29 describes the criteria under which these hospitals and units are excluded from IPPS.

3102 - General Information on IPPS Exclusion Deemed Providers and Suppliers
(Rev. 198, Issued: 01-17-20, Effective: 01-17-20, Implementation: 01-17-20)

The following providers and units are excluded from the Inpatient Prospective Payment System (IPPS):

- Psychiatric hospitals;
- Rehabilitation hospitals;
- Children’s hospitals;
- Long-term care hospitals;
- Psychiatric and rehabilitation units of hospitals;
- Cancer hospitals; and
- CAHs.

Certain kinds of Medicare-participating hospitals are paid under special provisions and are never subject to the IPPS. These hospitals need not be evaluated for compliance with the IPPS exclusion criteria:

- Hospitals paid under State cost control systems approved by CMS;
- Hospitals paid under demonstration projects approved by CMS;

As of the cost reporting period beginning on or after October 1, 2019, an IPPS-excluded hospital is no longer precluded from having an IPPS-excluded psychiatric and/or rehabilitation unit. For the purposes of payment, services furnished by a unit are
considered to be inpatient hospital services provided by the unit and not inpatient hospital services provided by the hospital operating the unit.

Although an IPPS-excluded hospital may have an IPPS-excluded unit, the excluded hospital may not have an IPPS-excluded unit of the same type (psychiatric or rehabilitation) as the hospital (for example, an Inpatient Rehabilitation Facility (IRF) may not have an IRF unit).

Note that a co-located, separately certified hospital or a separately certified hospital-within a hospital (HwH) is not a component of the hospital with which it shares a campus. For example, if a separately certified Medicare-participating short-term acute care IPPS hospital is located in the same building as a psychiatric hospital, it is not considered part of the psychiatric hospital and is not excluded from the IPPS based on the exclusion of the psychiatric hospital.

It is important to note that payment rules, such as the HwH (42 CFR 412.22(e)) or satellite facility (412.22(h)) rules, never waive or supersede the requirement that all hospitals must comply with the hospital conditions of participation (CoPs). All hospitals, regardless of payment status, must always demonstrate separate and independent compliance with the hospital CoPs, even when an entire hospital or a part of a hospital is located in a building also used by another hospital, or in one or more entire buildings located on the same campus as buildings used by another hospital.

An IPPS-excluded hospital operating an IPPS-excluded unit must continue to be in compliance with other Medicare regulations and CoPs applicable to the hospital and unit. An IPPS-excluded unit within a hospital is part of the hospital. Noncompliance with any of the hospital CoPs at 42 CFR 482.1 through 482.58 in any part of the certified hospital is noncompliance for the entire Medicare-certified hospital. Therefore, noncompliance with the hospital CoPs in an IPPS-excluded unit is noncompliance for the entire certified hospital (see 83 FR at 41514).

3104 - Criteria for PPS-Excluded Hospitals
(Rev. 1, 05-21-04)

3104A - Psychiatric Hospitals
(Rev. 1, 05-21-04)

A hospital certified as a psychiatric hospital under existing requirements is excluded.

3104B - Rehabilitation Hospitals
(Rev. 1, 05-21-04)

A hospital is an excluded rehabilitation hospital if:

- It has in effect a provider agreement to participate as a hospital;
During its most recent 12-month cost reporting period it treated an inpatient population of which at least 75 percent required intensive rehabilitative services for one or more of the following conditions:

- Stroke;
- Spinal cord injury;
- Congenital deformity;
- Amputation;
- Major multiple trauma;
- Fracture of femur (hip fracture);
- Brain injury;
- Poliarthritis, including rheumatoid arthritis;
- Neurological disorders, including multiple sclerosis, motor neuron diseases, polyneuropathy, muscular dystrophy, and Parkinson’s disease; or
- Burns.

It has in effect a preadmission screening procedure under which each prospective patient’s condition and medical history are reviewed to determine whether the patient is likely to benefit significantly from an intensive inpatient hospital rehabilitation program or assessment;

It ensures that patients receive close medical supervision, rehabilitation nursing, physical therapy, and occupational therapy plus, as needed, speech therapy, social services or psychological services, and orthotic and prosthetic services;

It has for each inpatient, a plan of treatment that is established, reviewed, and revised as needed by a physician in consultation with other professional personnel who provide services to the patient;

It uses a coordinated multidisciplinary team approach in the rehabilitation of each inpatient, as documented by periodic clinical entries to the patient’s medical record noting the patient’s status in relationship to goal attainment, and team conferences are held at least every two weeks to determine the appropriateness of treatment; and
• It has a director of rehabilitation who:
  
  o Provides services to the hospital and its inpatients on a full-time basis;
  
  o Is a Doctor of Medicine or Osteopathy licensed under State law to practice medicine or surgery; and
  
  o Has had, after completing a 1-year hospital internship, at least two years of training in the medical management of inpatients requiring rehabilitation services.

The inpatient population percent rule is applied to the 12-month cost reporting period immediately preceding the cost reporting period for which exclusion is sought. The cost reporting period need not be complete when this evaluation takes place. The RO requests the servicing intermediary to make the determination concerning the inpatient population percent rule.

Alternative: A hospital that seeks exclusion as a rehabilitation hospital for the first full 12-month cost reporting period that occurs after it becomes a Medicare participating hospital could not possibly meet the usual rule above, so a first-time exception to the rule is provided the hospital if it provides a written certification that the inpatient population it intends to serve will meet the inpatient population percent rule.

The written certification described above is effective for any cost reporting period of not less than one month and not more than 11 months occurring between the dates the hospital began participating in Medicare and the start of the hospital’s regular 12-month cost reporting period. This exception is available only once, when the facility first gains excluded status.

If a new rehabilitation hospital is excluded from PPS for a cost reporting period, but the inpatient population treated in the hospital during the period does not actually meet the inpatient population percent rule, a retroactive adjustment of payments to the hospital for the period is needed. Where this occurs, the FI advises the RO of the identity of the hospital and of the dates of the cost reporting period involved.

**3104C - Children’s Hospitals**

(Rev. 1, 05-21-04)

A hospital is an excluded children’s hospital if it has in effect an agreement to participate as a hospital, and more than 50 percent of its inpatients are individuals under the age of 18. The servicing intermediary verifies compliance.
A hospital is an IPPS-excluded long-term care hospital (LTCH) if it has in effect a provider agreement to participate as a hospital and the average inpatient length of stay is greater than 25 days. The average length of stay, for this purpose, is determined by dividing the total number of inpatient days for Medicare patients not paid at the site neutral rate or under a Medicare Advantage plan (excluding leave of absence or pass days) by the total number of Medicare discharges for the cost period not paid at the site neutral rate or under a Medicare Advantage plan. The servicing MAC verifies whether rehabilitation hospitals meet this length of stay criterion as LTCHs, and are, therefore, eligible for a LTCH exclusion and do not have to meet the special criteria otherwise established for these categories of facilities. The servicing MAC verifies length of stay for all LTCHs.

If an LTCH also has an IPPS-excluded psychiatric and/or rehabilitation unit, the days and discharges from those excluded units are not included in the calculation of an LTCH's average length of stay (83 FR 41515). Patients in IPPS-excluded units in an LTCH are not paid under the LTCH PPS.

Long-term care hospitals that occupy space in a building also used by another hospital, or in one or more buildings located on the same campus as buildings used by another hospital (i.e., the host facility), must meet additional “hospital-within-a-hospital” or satellite criteria.

NOTE: Section 15008(a) of the 21st Century Cures Act (Pub. L. 114-255) removed the LTCH category under section 1886(d)(1)(B)(iv)(II) of the Social Security Act (implemented in the regulations at 42 CFR 412.23(e)(2)(ii)) and created a new category of IPPS-excluded hospital at section 1886(d)(1)(B)(vi) of the Act (implemented in the regulations at 42 CFR 412.22(i)), which is referred to as extended neoplastic disease care hospitals, effective January 1, 2015. Although this category of hospitals have LTCH CCNs, they are not required to meet other LTCH specific requirements.

A IPPS-excluded hospital that occupies space in a building also used by another hospital which is not excluded from the IPPS, or in one or more entire buildings located on the same campus as buildings used by another hospital which is not excluded from the IPPS, must meet the criteria at 42 CFR 412.22(e) in order to maintain its IPPS-excluded status as follows:

- The hospital has a governing body that is separate from the governing body occupying space in the same building or campus, and the governing body is not controlled by the host facility or any third entity that controls both hospitals;
NOTE: For purposes of this section, “control” exists if an individual or an organization has the power, directly or indirectly, significantly to influence or direct the actions or policies of an organization or institution.

- The hospital has a chief medical officer who reports directly to the governing body and who is responsible for all medical staff activities of the hospital, and is not employed or under contract with the host facility or any third party that controls both hospitals;

- The hospital has a separate medical staff from the medical staff of the host facility, reports directly to the hospital’s governing body, and adopts and enforces bylaws governing medical care provided in the hospital and medical staff activities, including the granting of privileges to individual practitioners;

- The hospital has a single chief executive officer through whom all administrative authority flows and who exercises control and surveillance over all administrative activities at the hospital, and who is not employed by or under contract to the host facility or any third party who controls both hospitals; and

If a State hospital that is occupying space in the same building or on the same campus as another State hospital cannot meet the separate governing body criterion solely because its governing body is under the control of the State hospital with which it shares a building or a campus, or is under the control of a third entity that also controls the State hospital with which it shares a building or a campus, the State hospital can nevertheless qualify for an exclusion if it meets the other applicable criteria in §412.22(e) and the following:

- Both State hospitals occupy space in the same building or on the same campus and have been continuously owned and operated by the State since October 1, 1995;

- Is required by State law to be subject to the governing authority of the State hospital with which it shares space or the governing authority of a third entity that controls both hospitals; and

- Was excluded from the IPPS before October 1, 1995, and continues to be excluded from the inpatient prospective payment system through September 30, 2008.

If a hospital was excluded from the IPPS on or before September 30, 1995, and at that time occupied space in a building also used by another hospital, or in one or more buildings located on the same campus as buildings used by another hospital, it is not required to meet §412.22(e)(1)(i)-(iv) in order to maintain its IPPS-excluded status so long as it operates under the same terms and conditions in effect on September 30, 2003. For cost reporting periods beginning on or after October 1, 2006, a hospital may decrease its number of beds and remain excused from the separateness and control requirements as long as it does not increase its beds above the number it had on
September 30, 2003. Effective January 1, 2020 a grandfathered Children’s HwH may increase beds without losing its grandfathered status.

The SA notifies the CMS RO as soon as it becomes aware of any LTCH planning to operate as a HwH and notifies the facility immediately that it must demonstrate compliance with the special HwH criteria. The SA will review documentation for hospitals that intend to operate as HwH in order to make an initial recommendation to the ROs regarding a hospital’s compliance or noncompliance with the above criteria (See §3104.D). Final determinations will be made on a case-by-case basis by the RO using whatever procedure it deems appropriate. In some instances, it may be necessary to authorize a SA onsite inspection of the hospital by the State agency to collect additional information. The hospital must submit a completed Form CMS-855 to notify the MAC of its intent to be a HwH.

3106 - Criteria for Psychiatric and Rehabilitation Units
(Rev. 1, 05-21-04)

A PPS-excluded psychiatric unit must meet both the general criteria for units and the specific criteria for psychiatric units below. A PPS-excluded rehabilitation unit must also meet the general criteria for units and the specific criteria for rehabilitation units below.

3106A - General Criteria for Units
(Rev. 1, 05-21-04)

- The unit is a part of an institution that:
  - Has in effect a provider agreement to participate as a hospital;
  - Is not excluded in its entirety from PPS; and
  - Has enough beds not excluded from PPS to permit the provision of adequate cost information.

- The unit has written admission criteria applied uniformly to both Medicare and non-Medicare patients;

The unit has admission and discharge records that are identified separately from those of the hospital in which it is located, and that are readily retrievable. The medical records of the unit’s patients need not be physically separate from the records of patients in the acute care portion of the hospital. It is not necessary to create a second medical record when a patient is moved from the acute care portion of the hospital to the excluded unit or vice versa. The record, however, must indicate, for Medicare purposes, the dates of admission and discharge from the excluded unit. The unit’s policies provide that necessary clinical information accompany a patient upon transfer from the hospital to the unit;
• If State law provides special licensing requirements for psychiatric or rehabilitation units, the unit is licensed in accordance with the applicable requirements;

• The hospital’s UR plan includes specific standards for the type of care offered by the unit;

• The beds assigned to the unit are physically separate from (not commingled with) beds not included in the unit;

The unit must also meet the accounting requirements set forth in §§2803.B.1.g through j of the Provider Reimbursement Manual. These include requirements that:

• The unit is treated as a separate cost center for cost finding and apportionment purposes;

• The hospital’s accounting system properly allocates costs attributable to the unit and maintains statistical data that are adequate to support the basis of allocation of shared costs;

• The cost report for the hospital includes the costs of the unit in the same fiscal period and uses a single method of cost apportionment;

• As of the first day of the first reporting period for which all other exclusion requirements are met, the unit is fully equipped and staffed, and capable of providing inpatient psychiatric or rehabilitation care, regardless of whether there are any inpatients in the unit on that date; and

• Each hospital may have only one unit of each type (psychiatric or rehabilitation) excluded from PPS.

3106B - Specific Criteria for Psychiatric Units
(Rev. 121, Issued: 09-19-14, Effective: 09-19-14, Implementation; Upon Implementation of ICD-10)

An SA onsite verification or reverification survey for PPS exclusion of a psychiatric unit is required for a hospital filing a first-time request for PPS exclusion for its psychiatric unit, a psychiatric unit that has been selected as part of a sample for an annual validation survey, and/or a complaint against a psychiatric unit. For cost reporting periods following the first cost reporting period, the hospital is to self-attest that its psychiatric unit is in compliance with the requirements at 42 CFR 412.27.

3106B1 - Patient Criteria
The unit admits only patients requiring admission for active treatment, of an intensity that can be provided only in an inpatient hospital setting. The psychiatric principal diagnosis must be one contained in

- the Fourth Edition of the American Psychiatric Association Diagnostic and Statistical Manual;

- Chapter 5 (“Mental Disorders”) of the International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM); or

Chapter 5 ("Mental and Behavioral Disorders") of the International Classification of Diseases, Tenth Revision, Clinical Modification (ICD-10-CM.), upon implementation of ICD-10.

3106B2 - Services Provided

The unit furnishes, through the use of qualified personnel, psychological, social work, psychiatric nursing, occupational and recreational therapy services; and

3106B3 - Medical Records

The unit maintains medical records that permit determination of the degree and intensity of treatment provided to individuals who are furnished services in the unit, and that meet the following requirements:

a. Development of Assessment/Diagnostic Data

The medical record stresses psychiatric findings in the history, physical examination findings and treatment plan. It also includes the doctor’s orders for the psychiatric condition for which the patient is treated.

- The identification data includes the inpatient’s legal status (i.e., court commitment or voluntary admission);

- A provisional or admitting diagnosis is made at the time of admission. The record also includes the diagnoses of intercurrent diseases as well as psychiatric diagnoses;

- The reasons for admission are clearly documented as stated by the patient or others significantly involved, or both;

- The social service record, including reports of interviews with patients, family members, and others provides an assessment of home plans and family attitudes, community resource contacts and a social history; and
• When indicated, a complete neurological examination is recorded at the time of the admission physical examination.

b. Psychiatric Evaluation

Each patient receives a psychiatric evaluation that:

• Is completed within 60 hours of admission;
• Includes a medical history;
• Contains a record of mental status;
• Notes the onset of illness and the circumstances leading to admission;
• Describes attitudes and behavior;
• Estimates intellectual functioning, memory functioning, and orientation; and
• Includes an inventory of the patient’s assets in descriptive, not interpretive, fashion.

c. Treatment Plan

• Each patient has a comprehensive treatment plan based on an inventory of his/her strengths and disabilities; and

• The written plan includes:
  o A substantiated diagnosis;
  o Short-term and long-term goals;
  o The specific treatment modalities;
  o The responsibilities of each member of the treatment team; and
  o Documentation that justifies the diagnosis and treatment and all active therapeutic efforts.

d. Progress Notes

Physician progress notes must be documented by a Doctor of Medicine or Osteopathy responsible for the care of the patient, a nurse, a social worker and, when appropriate, others significantly involved in active treatment modalities. Progress notes’
frequency is based on the condition of the patient, but they must be recorded at least weekly for the first 2 months, and at least monthly thereafter. They should contain recommendations for revisions in the treatment plan as indicated, as well as a precise assessment of the inpatient’s progress in accordance with the original or revised treatment plan.

e. Discharge Planning and Discharge Summary

The record of each discharged patient has a discharge summary that must include a recapitulation of the patient’s hospitalization in the unit, recommendations from appropriate services concerning follow-up or aftercare, and a brief summary of the patient’s condition on discharge.

3106B4 - Staffing

The unit meets special staff requirements and has adequate numbers of qualified professional and supportive staff to evaluate inpatients, formulate written, individualized comprehensive treatment plans, provide active treatment measures, and engage in discharge planning, as follows:

a. Personnel

The unit employs or undertakes to provide adequate numbers of qualified professional, technical, and consultative personnel to:

- Evaluate patients;
- Formulate written individualized comprehensive treatment plans;
- Provide active treatment measures; and
- Engage in discharge planning.

b. Director of Inpatient Psychiatric Services and Medical Staff

The number and qualifications of Doctors of Medicine and Osteopathy are adequate to provide essential psychiatric services.

Inpatient psychiatric services are under the supervision of a clinical director, service chief, or equivalent who is qualified to provide the leadership required for an intensive treatment program and who:

- Meets the training and experience requirements for examination by the American Board of Psychiatry and Neurology or the American Osteopathic Board of Neurology and Psychiatry, and
• Monitors and evaluates the quality and appropriateness of services and treatment provided by the medical staff.

In verifying the training and experience requirements of the clinical director, the SA follows the Hospital Interpretive Guidelines and survey procedures specified in Appendix A. (See 42 CFR 482.62(b)(1).) A director is qualified to take the examination for board certification upon successful completion of a psychiatric residency program approved by either of the two boards.

c. Nursing Services

The unit has a qualified director of psychiatric nursing services. There are also adequate numbers of RNs, LPNs, and mental health workers to provide care necessary under each patient’s active treatment program and to maintain progress notes on each patient.

The director of psychiatric nursing services is an RN who has a master’s degree in psychiatric or mental health nursing, or its equivalent, from a school of nursing accredited by the National League for Nursing, or is qualified by education and experience in the care of the mentally ill. The director demonstrates competence to participate in interdisciplinary formulation of individual treatment plans; to give skilled nursing care and therapy; and to direct, monitor, and evaluate the nursing care furnished.

The staffing pattern ensures the availability of a registered nurse 24 hours each day. There are adequate numbers of registered nurses, licensed practical nurses, and mental health workers to provide the nursing care necessary under each inpatient’s active treatment program.

d. Psychological Services

The unit provides or has available psychological services to meet the needs of the patients. The services are furnished in accordance with accepted standards of practice and established policies and procedures.

e. Social Services

There is a director of social services who monitors and evaluates the quality and appropriateness of social services furnished. The services are furnished in accordance with accepted standards of practice and established policies and procedures.

Social service staff responsibilities include, but are not limited to, participating in discharge planning, arranging for follow-up care, and developing mechanisms for exchange of appropriate information with sources outside the hospital.

f. Therapeutic Activities
The unit provides a therapeutic activities program. The program is appropriate to the needs and interest of patients and is directed toward restoring and maintaining optimal levels of physical and psychosocial functioning. The number of qualified therapists, support personnel, and consultants are adequate to provide comprehensive therapeutic activities consistent with each patient’s active treatment program.

### 3106C - Specific Criteria for Rehabilitation Units

(Rev. 1, 05-21-04)

An SA onsite verification or reverification survey for PPS exclusion for a rehabilitation unit is to be performed for a hospital’s first-time request for PPS exclusion for its rehabilitation unit, a rehabilitation unit that is selected as part of a sample for an annual validation compliance survey, and/or a complaint against a rehabilitation unit. For cost reporting periods following the first cost reporting period, the hospital will self-attest that its rehabilitation unit is in compliance with the requirements in 42 CFR 412.29.

The unit meets the requirement in §3104.B, except as provided below, with respect to patients treated in the unit during the hospital’s most recent 12-month cost reporting period, i.e., the period immediately preceding the period for which the exclusion would be effective. This finding is based on the medical conditions of all (i.e., Medicare and non-Medicare) patients who occupy the beds assigned to the physically separate unit. The medical condition of all patients treated in the unit is considered.

If a hospital has not previously sought exclusion for any rehabilitation unit, and has both increased its bed capacity under Medicare certification and obtained approval for added bed capacity under State licensure, it may identify the new beds as a new rehabilitation unit for the first full 12-month cost-reporting period during which the unit is in service. For purposes of these provisions, “new beds” are defined as ones for which the hospital has obtained approval by increasing its bed capacity under both State licensure and Medicare certification. Note that there is no net increase if the hospital adds 20 new beds and deletes 20 beds previously licensed and certified for conversion. A unit that is comprised of some beds that were previously licensed and certified, and some new beds, will be recognized as a new rehabilitation unit only if over half of the beds are new. Beds are considered “new” only for the first full 12-month cost reporting period in which a hospital seeks exclusion of a new rehabilitation unit. The hospital may provide written certification that the inpatient population it intends the unit to serve meets the 75 percent rule instead of showing that it has treated such a population during its most recent 12-month cost reporting period.

The hospital that has an excluded rehabilitation unit must obtain approval for added bed capacity under State licensure requirements. If the hospital seeks to add the new beds to its existing excluded unit for the first full 12-month cost reporting period during which the new beds are used to furnish inpatient care, it must provide written certification that the new beds are intended to meet the inpatient population percent rule (see §3104.B).
instead of showing that those beds were used to treat such a population during the unit’s most recent 12-month cost reporting period.

The written certification described above is effective for any cost reporting period of not less than one month and not more than 11 months occurring between the dates the hospital began participating in Medicare and the start of the hospital’s regular 12-month cost reporting period. For purposes of this exclusion, a hospital that has undergone a change of ownership or leasing (see §3210) is not considered to have participated previously in the Medicare program.

If a hospital has a new rehabilitation unit excluded from PPS for a cost reporting period, or expands an existing PPS-excluded rehabilitation unit through the addition of new beds as defined above, but the inpatient population treated in the new unit of added beds during the period does not actually meet the inpatient population percent rule, a retroactive adjustment of payments to the hospital for the period is needed. Where this occurs, the FI advises the RO of the identity of the hospital and the dates of the cost reporting period involved.

If a hospital that has not previously participated in the Medicare program seeks exclusion of a rehabilitation unit, it may designate certain beds as a new rehabilitation unit for the first full 12-month cost reporting period that occurs after it becomes a Medicare participating hospital. The written certification described above also is effective for any cost reporting period of not less than one month and not more than 11 months occurring between the dates the hospital began participating in Medicare and the start of the hospital’s regular 12-month cost reporting period. For purposes of this exclusion, a hospital that has undergone a change of ownership or leasing (see §3210) is not considered to have participated previously in the Medicare program.

- The unit meets the requirements for a rehabilitation hospital in §3104. (The intermediary verifies the 75 percent rule.)

- The unit has a director of rehabilitation who:
  - Is a Doctor of Medicine or Osteopathy licensed under State law to practice medicine or surgery;
  - Has had, after completing a 1-year hospital internship, at least two years of training or experience in the medical management of inpatients requiring rehabilitation services; and
  - Provides services to the unit and its inpatients for at least 20 hours per week.

If the rehabilitation unit serves both inpatients and outpatients through a single, integrated unit, the time spent by the director in performing administrative duties for the entire unit counts toward the time requirement. The SA does not prorate this administrative time
between inpatients and outpatients. However, time devoted to performing direct patient care can count toward the time requirement only if furnished to inpatients of the unit.

**3108 - SA First-Time Verification Procedures for Hospitals and Units (Rev. 1, 05-21-04)**

At the time the SA is requested to verify PPS exclusion of a rehabilitation hospital, or a psychiatric or rehabilitation unit of a hospital for the first time, it completes the appropriate part of the Criteria Worksheet, Form CMS-437, 437A, or 437B (Exhibit 73). The SA verifies that the criteria specified below are met for exclusions of hospitals and units of hospitals. In addition, the RO may instruct the SA to verify some or all of the general criteria for units given in §3106.A.

**3108A - Rehabilitation Hospitals and Rehabilitation Units of Hospitals (Rev. 1, 05-21-04)**

1. Rehabilitation Hospitals - The SA verifies that the criteria in §3104.B are met. (The intermediary verifies the 75 percent rule.)

2. Rehabilitation Units - The SA verifies that the criteria in §3106.C are met. (The intermediary verifies the 75 percent rule.)

3. Criteria Presumed Met by Accredited Rehabilitation Hospitals and Units - A rehabilitation hospital or unit may be presumed to meet the criteria in §3104.B or §3106.C (excluding the 75 percent rule and the director requirement) if it is accredited by:

   - Commission on Accreditation of Rehabilitation Facilities (CARF), surveyed under the Comprehensive Inpatient Rehabilitation (CIRP) Program; or

   - Joint Commission on Accreditation of Healthcare Organizations (JCAHO). Hospitals/units surveyed under the Joint Commission Comprehensive Physical Rehabilitation Program or Unit includes the provisions of standard RH.3 in the Joint Commission Accreditation Manual for Hospitals.

Nevertheless, in either case, the SA verifies that the criterion for full-time director for rehabilitation hospitals or the 20-hour per week criterion for the rehabilitation unit is met. The SA reviews appropriate documentation submitted by the facility (e.g., payroll records, duty rosters, staff appointment notice specifying either full-time or part-time appointment of at least 20 hours per week).

**3108B - Psychiatric Units of Hospitals (Rev. 1, 05-21-04)**
The SA verifies that the criteria in §3106 are met.

The SA schedules an onsite verification visit at least 90 days prior to the end of the hospital’s cost reporting period. The SA records the survey findings using the applicable section of the Criteria Worksheet, Form CMS-437 and transmits the worksheet to the RO at least 60 days prior to the end of the hospital’s cost reporting period for inclusion with other information necessary for determining exclusion from PPS.

For a first-time verification survey, the SA verifies compliance with 42 CFR 412.27(a), (b), and (c) by reviewing at least one patient record. If there are no patients in the unit at the time of the survey, the SA reviews patient records (at least one) for patients treated in the unit within 6 months of the date of the survey. If the psychiatric unit has not treated any patients during the 6 month period prior to the survey (i.e., there are no closed or active patient records available for review), the unit cannot demonstrate compliance with 42 CFR 412.27(a), (b), and (c), and the SA survey should be rescheduled to a later date when records will be available.

Hospitals/units that do not meet all exclusion criteria at the time of the SA onsite verification may still be eligible for exclusion. They must provide strong evidence documenting compliance with exclusion requirements at least 15 days prior to the beginning of the cost reporting period. If necessary, the SA contacts the hospital/unit again to confirm their compliance. However, a second onsite visit generally is not required to confirm evidence submitted subsequent to onsite verification. For example, if a rehabilitation hospital has a part-time director and is able to furnish proof that a full-time director will be employed prior to the start of the cost reporting period, no revisit is necessary.

3110 - SA Reverification of PPS-Excluded Hospitals and Units
(Rev. 1, 05-21-04)

3110A - Annual Reverification Process for Nonaccredited, PPS-Excluded, Rehabilitation Hospitals and Units
(Rev. 1, 05-21-04)

- 120 days before the beginning of the next cost reporting period, the SA notifies the excluded hospital or unit (Exhibit 126) that it must self-attest to compliance with the appropriate requirements in 42 CFR 412.23(b), 412.25, and/or 412.29.

- The SA includes a copy of the attestation statement (Exhibit 127) and the appropriate hospital or unit criteria worksheet (Form CMS-437A or 437B, Exhibit 73);

- The hospital/unit is to return the completed/signed worksheet and signed attestation statement to the SA office no later than 90 days before the beginning of its next cost reporting period;
• The SA transmits the completed attestation statement and worksheet, along with its recommendation for reverification, to the RO at least 60 days prior to the end of the hospital’s cost reporting period for inclusion with other information necessary for determining exclusion from PPS.

3110B - Reverification Process for Rehabilitation Hospitals and/or Units Accredited by CARF Under CIRP or JCAHO
(Rev. 1, 05-21-04)

Accredited rehabilitation hospitals or units may be presumed to meet the criteria in §§3104.B or 3106.C, excluding the 75 percent rule (verified by the intermediary and the director requirement (42 CFR 412.23(b)(5) or 412.29(f)(1), as appropriate). Accredited rehabilitation hospitals/units self-attest to compliance with the director requirement on Form CMS-437A or Form CMS-437B using the same procedure and processing timeframes as used for nonaccredited hospitals/units.

3110C - Reverification Process for Psychiatric Units of Hospitals:
(Rev. 1, 05-21-04)

• The SA uses the same process and timeframes as those used for nonaccredited rehabilitation hospitals and units to determine if the unit meets the specific criteria for psychiatric units in §3106; and

• The SA provides the psychiatric unit with the criteria worksheet for psychiatric units, Form CMS-437 (Exhibit 73), along with Exhibit 126 and the attestation statement (Exhibit 127).

Hospitals/units that do not self-attest to meeting all exclusion criteria at the time of the annual self-attestation may still be eligible for exclusion. They must provide strong verifiable evidence documenting compliance with exclusion requirements at least 15 days prior to the beginning of the cost reporting period. If necessary, the SA contacts the hospital/unit again to confirm compliance. A SA onsite visit generally is not required to confirm evidence submitted subsequent to self-attestation.

3112 - RO Procedures for Exclusion from PPS for Hospitals and Units
(Rev. 1, 05-21-04)

For initial exclusion from PPS, hospitals (except those now certified as psychiatric hospitals) and hospital units that meet the criteria of this section have been instructed in the Provider Reimbursement Manual to notify the RO of their eligibility for exclusion. Notification should be in writing and include the following: name of hospital, type of hospital/units, address, current provider identification number, name of contact person, FI, and a statement that the hospital/units(s) meets the criteria for exclusion. Notification should be made, where possible, no later than 5 months before the date the hospital becomes subject to PPS.
Hospitals and units that have already been excluded from PPS need not reapply for exclusion. These facilities will be reevaluated based on the criteria described in §3112.2 to determine if they still meet the PPS exclusion criteria.

If the RO does not receive notification of a hospital/unit meeting the criteria of this section but has knowledge that it should be excluded, the RO identifies the hospital/unit for exclusion.

Except for currently certified psychiatric hospitals, the RO notifies the hospital and the FI, in writing, 45 days prior to the start of the cost reporting period, of the excluded status of the hospital or unit(s) and of any new provider number(s). The RO coordinates these activities very closely with the servicing intermediary of the hospital. Prior to making a decision on hospital/unit exclusion, the RO determines whether the hospital/unit meets all exclusion criteria including those criteria that are the responsibility of the servicing intermediary.

PPS excluded or nonexcluded status for a hospital or hospital unit remains in effect for the entire cost-reporting period for which the determination is made. If a change in meeting applicable criteria occurs during a cost reporting period or the hospital requests exclusion after the start of its cost reporting period, the status already determined for that period will remain for the duration of the period. However, any change in status resulting from the beginning or end of a hospital’s participation in an approved demonstration project or State reimbursement control program will be effective on the date the change occurs, whether or not the date coincides with the start of a cost reporting period. A hospital may increase or decrease the space (square footage or number of beds) assigned to an excluded unit only at the start of the hospital’s next cost reporting period.

3112.1 - RO Procedures for First-Time Exclusion of Hospitals and Units (Rev. 1, 05-21-04)

When considering a hospital or hospital unit for exclusion for the first time, the RO has the SA and or the intermediary verify the facility’s compliance with exclusion criteria, as follows:

- Psychiatric Hospitals - The RO verifies through a review of records that the hospital currently participates in Medicare as a psychiatric hospital, and that the hospital’s provider number identifies it as a psychiatric hospital. The hospital is not required to make a separate request for exclusion.

- Rehabilitation Hospitals - The RO has the SA verify that the exclusion criteria in §3104 are met. As noted in §3108, a rehabilitation hospital may be presumed to meet certain criteria based on accreditation by the Commission on Accreditation of Rehabilitation Facilities (CARF) or by the Joint Commission on Accreditation of Healthcare Organizations (JCAHO). However, the SA should verify compliance with the medical director requirement, and the intermediary should verify compliance with the 75 percent rule for hospitals other than new hospitals.
- Children’s Hospitals - The RO has the intermediary verify that the hospital has in effect an agreement to participate as a hospital and that a majority of the hospitals inpatients are individuals under the age of 18. The determination is to be based on the hospital’s most recently filed cost report, unless there is an indication that the age of the patient population has changed since the close of the period covered by the report. If the age of the patient population has changed since that period, the RO has the intermediary determine whether the age criterion is met by the patient population treated during the prior 6-month period. The intermediary may base the determination either on its knowledge of the provider or on a separate contact that results in an actual review of a sample of patient records.

- Long-term Care Hospitals - The RO has the intermediary verify that the hospital has in effect an agreement to participate as a hospital and that the average length of inpatient stay is greater than 25 days. The average length of inpatient stay is to be computed by dividing the number of total inpatient days (less leave or pass days) by the number of total discharges for the hospital’s most recent complete cost reporting period. However, if a change in the hospital’s average length of stay is indicated, the hospital’s average length of stay is to be computed by the same method for the immediately preceding 6-month period. Rehabilitation hospitals meeting the length-of-stay criterion for exclusion as a long-term hospital are to be excluded as long-term hospitals, and should not be evaluated for exclusion under the rehabilitation hospital criteria. (See §3104).

- Psychiatric Distinct-Part Units - The RO has the intermediary verify that the general criteria for exclusion for units in §3106 are met. The RO has the SA verify that the specific criteria for psychiatric units in §3106 are met.

- Rehabilitation Distinct-Part Units - The RO has the intermediary verify that the general criteria for units in §3106 are met. The RO has the SA verify that the exclusion criteria in §3104 are met. As noted in §3108, a rehabilitation hospital may be presumed to meet certain criteria based on accreditation by CARF or by JCAHO. However, the SA should verify compliance with the medical direction requirement and the intermediary should verify compliance with the 75 percent rule for rehabilitation distinct-part hospital units other than new units.

- Cancer Hospitals - The RO contacts the Center for Medicare Management to obtain a listing of the hospitals that have been designated as cancer hospitals.

3112.2 - RO Verifying Continued Compliance With Exclusion Criteria by Currently Excluded Hospitals or Units
(Rev. 1, 05-21-04)
3112.2A - Self-Attestation Procedures for PPS-Excluded Hospitals and Units
(Rev. 1, 05-21-04)

1. Rehabilitation Hospitals/Units and Psychiatric Units - Annual verification surveys for all previously excluded rehabilitation hospitals and units, and psychiatric units are no longer required. The new procedures is as follows:

   o At least 120 days prior to the beginning of the next cost reporting period, SAs provide (see Exhibit 126) excluded rehabilitation hospitals/units and psychiatric units with the attestation statement (Exhibit 127) and the appropriate Criteria Worksheet, Form CMS-437, 437A, or 437B;

   o Hospital/unit officials complete and sign the attestation statement and the appropriate Worksheet and return them to the SA no later than 90 days before the beginning of the next cost reporting period; and

   o After receiving the hospital/unit’s self-attestation materials from the SA, the RO notifies the hospital/unit (see Exhibit 193) that recertification has been approved.

   • Previously excluded hospitals/units are required to report any change in operations (e.g., expansion or downsizing) to the appropriate CMS RO and to provide the SA with a copy of the report within 10 days after the change occurs;

   • The SA conducts annual validation compliance surveys at excluded hospitals/units;

   • SAs continue to conduct complaint surveys at excluded hospitals/units;

   • SAs continue to conduct first-time verification surveys in connection with a hospital/unit’s first exclusion from PPS; and

   • FIs continue to verify, on an annual basis, compliance with the 75 percent rule (see 42 CFR 412.23 through 412.30) for rehabilitation hospitals and units.

3112.2B - RO Verifying Exclusion Eligibility of Other Facilities
(Rev. 1, 05-21-04)

1. Currently Certified Psychiatric Hospital - A hospital currently participating in Medicare and identified by its provider number as a psychiatric hospital is excluded from PPS and is not required to make any special requests for exclusion.

2. Children’s Hospital - A hospital is an excluded children’s hospital if it has in effect an agreement to participate as a hospital, and if the majority of its inpatients are individuals under the age of 18. The determination is based on the hospital’s
most recently filed cost report. If there is an indication that the age of the patient population has changed since the close of the period covered by the report, the RO uses data for the prior 6-month period, asking the servicing intermediary to verify whether the age criterion has been met (i.e., whether the majority of the hospital’s inpatients are individuals under the age of 18). This may be based on the intermediary’s knowledge of the provider, or based on a separate contact.

3. Long-term Hospitals - A hospital is an excluded long-term hospital if it has in effect an agreement to participate as a hospital and if the average inpatient length of stay is greater than 25 days. The RO bases its determination on the hospital’s most recently filed cost report. If there is an indication that the length of stay has changed since the close of the period covered by the report, use data for the prior 6-month period, asking the servicing intermediary to verify whether the length of stay criterion has been met (i.e., whether the average length of stay is in excess of 25 days). Rehabilitation hospitals meeting the length of stay criterion as a long-term hospital are eligible for a long-term hospital exclusion from PPS and do not have to meet the special criteria established for these categories of facilities.

4. The “hospital-within-hospital” criteria described in §3112.1 apply to all long-term hospitals with cost reporting periods beginning on or after October 1, 1995. If the RO becomes aware of any long-term hospital operated in a building or campus occupied by another hospital, the hospital must be in compliance with the criteria.

5. Cancer Hospitals - The RO verifies through contact with the Center for Medicare Management, that the hospital continues to be designated as a cancer hospital.

3112.3 - Role of FIs in Reverification of PPS Excluded Hospitals and Units
(Rev. 1, 05-21-04)

The FIs are to verify the following:

- Rehabilitation Hospitals and Units - 75 percent rule applied to diagnoses.
- Children’s Hospitals - Age criterion.
- Long-Term Hospitals - Length of stay criterion.
- All Distinct Part Units - Unit is a separate cost center for cost finding and apportionment, meeting requirements of Provider Reimbursement Manual §2803.

Changes in Provider Status or Services
3200 - Action Based on Changes in Provider Organization, Services, or Action of Other Approving Agencies
(Rev. 1, 05-21-04)

Notification that an entity has undergone organizational changes, added or relocated units, or received an accreditation may require a change in SA scheduling.

3202 - Change in Size or Location of Participating SNF and/or NF
(Rev. 1, 05-21-04)

Under §1866 of the Social Security Act (the Act), the Secretary has the authority to enter into an agreement with an institution or an institutional complex to provide covered services to our beneficiaries. The provider agreement requires compliance with the requirements the Secretary deems necessary for participation in the Medicare or Medicaid program. See §1866(b)(2) and §1902 (a)(27) of the Act. On the effective date of the provider agreement, the institution or institutional complex is deemed to have met the requirements for participation based upon a survey of the institution or institutional complex as it was configured (i.e., bed size/bed location configuration) on the date(s) of the survey. The CMS’ authority to regulate bed size changes in a SNF or a NF is based on the authority to ensure compliance with the provider agreement under §1866 of the Act and to further ensure that the configuration that has been approved for the institution or institutional complex does not so drastically change from that of the original certified configuration so as to endanger resident health and safety or otherwise change in a material fashion the identity of the entity that CMS originally certified for program participation.

An institution or institutional complex may choose to participate in the Medicare and/or Medicaid programs either in its entirety (i.e., fully participating), or a portion thereof (i.e., a distinct part), but not both. If only a portion of an institution or institutional complex actually participates in either program it is classified as a distinct part and must meet the criteria found in §2762. For example, an institution has 4 wings that consist of 25 beds each. Three contiguous wings that contain 75 beds are dually participating (i.e., participating in Medicare and Medicaid). The fourth wing is only certified to participate in Medicare. It consists of 25 beds. Therefore, in this instance the institution is fully participating for purposes of Medicare (i.e., 100 beds) and a distinct part for purposes of Medicaid (i.e., 75 beds). The policies on bed size changes and changes in designated bed locations that are included in this section apply, regardless of whether an institution is fully participating (i.e., all beds within the institution or institutional complex are certified to participate in the Medicare and/or Medicaid program) or participating as or with a distinct part.

A SNF or NF may be:

- An entire institution for skilled nursing or rehabilitative care, such as a nursing home; or
• A distinct part of an institution such as, a hospital, personal care home, assisted living facility, board and care home, domiciliary care facility, rest home, continuing care retirement community or nursing home.

An institution that is primarily for the care and treatment of mental diseases cannot be a SNF or NF.

3202A - Requirements for Distinct Part Certification
(Rev. 1, 05-21-04)

If the institution or institutional complex is participating as a distinct part SNF and/or NF, for a change to be approved, the requested change in bed size must conform to the requirements to be classified as a distinct part. The term “distinct part” refers to a portion of an institution or institutional complex (e.g., a nursing home or a hospital) that is certified to provide SNF and/or NF services. A distinct part must be physically distinguishable from the larger institution and fiscally separate for cost reporting purposes.

An institution or institutional complex can only be certified with one distinct part SNF and/or one distinct part NF. A hospital-based SNF is by definition a distinct part. Multiple certifications within the same institution or institutional complex are strictly prohibited.

The distinct part must consist of all beds within the designated area. The distinct part can be a wing, separate building, a floor, a hallway, or one side of a corridor. The beds in the certified distinct part area must be physically separate from (that is, not commingled with) the beds of the institution or institutional complex in which it is located. However, the distinct part need not be confined to a single location within the institution or institutional complex’s physical plant. It may, for example, consist of several floors or wards in a single building or floors or wards that are located throughout several different buildings within the institutional complex. In each case, however, all residents of the distinct part would have to be located in units that are physically separate from those units housing other patients of the institution or institutional complex.

Where an institution or institutional complex owns and operates a distinct part SNF and/or NF, that distinct part SNF and/or NF is a single distinct part even if it is operated at various locations throughout the institution or institutional complex. The aggregate of the SNF and/or NF locations represents a single distinct part subprovider, not multiple subproviders, and must be assigned a single provider number.
Illustration I, above, is an illustration of a floor plan of a nursing facility followed below by examples that meet the requirements for a distinct part, as well as examples that do not meet the requirements for a distinct part. The purpose of the Illustration is to assist the State and the RO in ensuring proper distinct part certification.

**3202A1 - Meet Distinct Part Certification**  
(Rev. 1, 05-21-04)

An institution or institutional complex can select any one of the following examples discussed in the context of Illustration I above that meets the requirements for distinct part certification.

- All rooms numbered 1 through 12 in wing 1 and all rooms numbered 1 through 12 in wing 2 constitute a distinct part. This option is approvable because it constitutes all beds in each wing.

- All rooms numbered 1 through 12 in wing 5. This option is approvable because it includes all beds in the wing.

- Room numbers 1 through 6 in wing 4 constitute a distinct part. This option is approvable because it includes all beds that constitute a single side of the corridor.
• Room numbers 7 through 12 in wing 2 and all rooms 1 through 12 in wing 1 constitute a distinct part. This option is approvable because it includes all beds in wing 1 and all beds that constitute a single side of the corridor in wing 2.

3202A2 - Do Not Meet Distinct Part Certification
(Rev. 1, 05-21-04)

Neither of the examples discussed below, in the context of Illustration I, meet the requirements for distinct part certification.

• Room numbers 1 through 12 in wing 1 and rooms 3, 4, and 5 in wing 6 do not constitute a distinct part. This option is not approvable because of the inclusion of the three rooms in wing 6.

• Room number 2 in wing 1, room numbers 5 and 7 in wing 6, and room numbers 4, 5, 6, 10, 11, and 12 in wing 4. This option is not approvable because the distinct part area must be physically separate from (that is, not commingled with) the beds of the institution or institutional complex in which it is located.

3202B - Changes in Bed Size of Participating SNF and/or NF
(Rev. 1, 05-21-04)

When an institution or institutional complex not previously certified as or with a SNF and/or NF establishes a SNF and/or NF, it must be initially certified and periodically recertified. If an institution or institutional complex has an existing SNF and/or NF agreement, it may elect to change the number of beds that are certified to participate in the Medicare or Medicaid program up to two times per cost reporting year in accordance with the requirements set out below. Where a change in the size of a SNF also impacts the size of a NF, or vice versa, this represents one change for the SNF and one change for the NF. An institution or institutional complex that is participating in the Medicare program can find these same requirements in §2337 of the Provider Reimbursement Manual. An institution or institutional complex may only change the bed size of its SNF and/or its NF once on the first day of the beginning of its cost reporting year and again on the first day of a single cost reporting quarter within that same cost reporting year in order to effect one of the following combinations:

• An increase in its bed size on the first day of the beginning of its cost reporting year and an increase in its bed size on the on the first day of a single cost reporting quarter that falls within the same cost reporting year; or

• An increase in its bed size on the first day of the beginning of its cost reporting year and a decrease in its bed size on the first day of a single cost reporting quarter that falls within the same cost reporting year; or
A decrease in its bed size on the first day of the beginning of its cost reporting year and an increase in its bed size on the first day of a single cost reporting quarter that falls within the same cost reporting year.

At no time can the RO or the SA approve two decreases in the bed size of an institution within the same cost-reporting year.

The institution or institutional complex may submit only ONE change in bed size at a time. Furthermore, an institution cannot request a change in its bed size just because it undergoes a change of ownership (CHOW) or because it has been approved to change its cost reporting year. In either of these circumstances, it is still bound by the filing requirements found in subsection C.

A request for a change in the number of certified beds cannot be approved on a retroactive basis. All changes are made on a prospective basis only in accordance with the effective date indicated above. The institution requesting a change in bed size must submit a written request to the RO or SA (as appropriate) in conformance with the requirements found in subsection C. An institution or institutional complex can not self-designate the effective date of a change in bed size.

### 3202C - General Request Filing Requirements
(Rev. 1, 05-21-04)

An institution or institutional complex seeking a change in the number of Medicare and/or Medicaid certified beds must:

- Submit a written request to the RO or SA (as appropriate) for the change 45 calendar days before:
  - The first day of its cost reporting year to effect a change on the first day of its cost reporting year; or
  - The first day of a single cost reporting quarter within the same cost reporting year at which time it seeks to change its bed size to effect a change on the first day of the designated cost reporting quarter.

- Submit floor plans identifying all areas of the institution or institutional complex with the current certified bed configuration and the proposed certified bed configuration in order for the RO or SA to determine that the proposed change is in fact, in conformance with the rules for full participation or distinct part certification, whichever applies.

- Include a reference to the cost-reporting year of the institution or institutional complex. If there has been a change in the cost-reporting year originally selected by the institution or institutional complex at the time of its initial certification,
submit a copy of the letter submitted to the fiscal intermediary and the fiscal intermediary’s response to the request.

3202D - Exceptions
(Rev. 1, 05-21-04)

There are certain situations (described below) that we believe warrant an exception to the above policy. Therefore, even if the institution or institutional complex has been approved for a change in bed size in accordance with the policies articulated above, the institution or institutional complex may be granted a change in bed size on the basis of one of these situations. To request a change in bed size based on one of these situations, the institution or institutional complex must file a written request with the RO or SA (as appropriate) 45 calendar days before the first day of its next cost reporting quarter, at which time the request will be effective if approved, along with floor plans identifying all areas of the institution or institutional complex with the current certified bed configuration and the proposed certified bed configuration. An exception may be granted based only on one of the following situations:

3202D1 - Life Safety Code (LSC) Requirements

An exception may be granted if the request is to reduce the size of the SNF or NF to avoid being out of compliance with LSC requirements (e.g., sprinkler installation). The proposed bed configuration must be separated from the rest of the institution or institutional complex by a 2-hour firewall, so that there is no danger of the fire spreading there from other parts not meeting safety requirements. In this case, the proposed reduction in the size of the SNF or NF may be established with an effective date that is requested by the institution or institutional complex, but not earlier than the date that the separation can be documented. A full survey by the fire authority must be performed if the reason for the request is to limit noncompliance with LSC requirements.

3202D2 - Elimination of Distinct Part

An exception may be granted if an institution or institutional complex concludes that it wants to become fully participating (i.e., all beds within the institution or institutional complex are certified to participate in the Medicare and/or Medicaid program). If the institution or institutional complex decides to become fully certified to participate in the Medicare and/or Medicaid program, it cannot return to distinct part certification until, at the earliest, the beginning of its next cost reporting year.

3202D3 - Enlargement through Construction, Purchase or Lease of Additional Space

An exception may be granted if the institution or institutional complex requests to increase the size of its SNF or NF to include space acquired through new construction, purchase or lease (e.g., constructing a new wing, purchasing an adjacent building or leasing a floor in a hospital).
3202E - Change in Designated Bed Location(s)
(Rev. 1, 05-21-04)

An institution or institutional complex may request to change its designated bed locations, as long as there is no change in the number of beds certified to participate in the Medicare and/or Medicaid program, by submitting a written request to the SA or the RO 30 calendar days in advance of such a change. In addition, the institution or institutional complex must submit floor plans identifying all areas of the institution or institutional complex with the current certified bed configuration and the proposed certified bed configuration in order for the RO or SA to determine that the proposed change is in fact, in conformance with the rules for full certification or distinct part certification, whichever applies. The institution or institutional complex must adhere to the notification requirements found in 42 CFR 483.10(b)(11)(ii)(A) and the residents’ rights requirements found in 42 CFR 483.10(o). The RO or SA must approve the request before the institution or institutional complex makes the change. No changes are made on a retroactive basis.

3202F - RO or SA (as appropriate) Actions Upon Receipt of Written Request for Change in Bed Size/Location
(Rev. 1, 05-21-04)

The RO or the SA must take the following actions when reviewing a request for a change in bed size:

- Date stamp the letter from the institution requesting a change in bed size with the date it was received by the RO or SA;

- Verify the cost-reporting year selected by the institution or institutional complex using the OSCAR/ASPEN system. The cost-reporting year of the provider must match what is contained in OSCAR/ASPEN. If the reported cost reporting year is different than that found in OSCAR/ASPEN it would be as a consequence of a change in cost reporting year (for Medicare) which must be approved by the fiscal intermediary in accordance with the requirements found in 42 CFR 413.34(f). Absent such a change, the institution or institutional complex must adhere to the cost reporting year selected at its initial certification;

- Document information as required under §2764;

- Complete the Form CMS-1539 reflecting the change in bed size/designated bed location(s) if the request is approved;

- Notify the institution or institutional complex in writing of the RO or SA decision to either approve or disapprove the request prior to the effective date of the change. If approved the letter must include the effective date of the change in bed
size and/or designated bed locations, the total number of beds certified and the designated bed locations. If disapproved the letter must explain the requirement(s) not met;

- Send a copy of the letter notifying the institution or institutional complex of the RO or SA decision to approve or disapprove the request to the appropriate fiscal intermediary;

- Update the OSCAR/ASPEN system.

Usually, advancing the scheduled SA standard survey to recertify the changed configuration is unnecessary. A telephone contact often resolves most questions, such as changing bylaws, staffing, or other issues regarding the capacity of the institution or institutional complex to furnish the level of care contemplated in the long term care requirements. The SA must advance the survey schedule and perform a survey if;

- There is reason to question whether the institution remains in compliance with the long term care requirements (e.g., the proposed relocation site is unsuitable);

- Information suggests that as a part of the change, a different governing body or managing personnel directs the distinct part. (See §3210.); or

- The area within the physical plant to be certified has not been subjected to a life safety code survey.

3202G - Evaluation
(Rev. 1, 05-21-04)

The SA bases its evaluation of the proposed certified area upon the following guidelines.

3202G1 - Shared Facilities and Services

Rarely is a distinct part SNF or NF so completely self-contained that it independently meets all of the long-term care requirements. Therefore, to the extent necessary, the SA evaluates services, facilities, and activities located outside the distinct part that are used by the distinct part’s residents. This evaluation is not an assessment of whether the distinct part meets the requirements to be considered provider-based for purposes of Medicare reimbursement.

Often, the distinct part will share central supporting services such as dietary, housekeeping, and plant maintenance with the rest of the institution or institutional complex. Depending on the size and type of the institution or institutional complex, the distinct part may also have shared administration and supervisory, medical, and therapeutic services.
The primary consideration in the evaluation of shared services is whether the sharing can be done without sacrificing the quality of care rendered to distinct part residents or endangering their health and safety. The distinct part must demonstrate a capacity to provide all of the services, facilities, and supervision required by the long term care requirements. For this reason, the SA may need to consider the total staff of an institution or institutional complex, particularly with respect to the amount of shared responsibilities.

3202G2 - Effect of Hospital Accreditation or Certification on SNF or NF

Make no assumption regarding a distinct part SNF or NF’s compliance with long term care requirements on the basis of the institutional or institutional complex’s accreditation by the Joint Commission on Accreditation of Healthcare Organizations or AOA or the institution or institutional complex’s Medicare participation. Survey and evaluate the institution or institutional complex to determine its compliance with all of the long-term care requirements.

3202G3 - SNF or NF as Distinct Part of a Psychiatric Hospital

The guidelines for the identification of a distinct part SNF or NF, regardless of the type of institution or institutional complex in which it is located, are generally applicable. However, there are special factors to consider when an institutional complex is certified to participate as a psychiatric hospital.

A SNF or NF cannot be certified if it is primarily for the care and treatment of mental diseases. In the context of a psychiatric hospital, for example, the presumption is that in most cases a SNF or NF distinct part of such a hospital is designed primarily for the care and treatment of patients with mental diseases. A distinct part SNF or NF cannot be established unless the psychiatric hospital either has a separate medical-surgical unit that is participating as a distinct part general hospital or has an arrangement with a community hospital for transfer to the hospital and back to the distinct part for post-hospital convalescence when a beneficiary requires medical-surgical services. In determining whether a distinct part SNF or NF is primarily for the care and treatment of mental diseases, the SA must look at the primary purposes for the unit’s existence, in combination with the requirements discussed above. A psychiatric hospital can have such a unit or section certified as a distinct part SNF or NF, only if the primary purpose of the unit is to provide medical services and the hospital meets one of the requirements discussed in the last sentence of the preceding paragraph.

In addition, a distinct part SNF or NF of a psychiatric hospital would also have to be licensed pursuant to the State or local law which provides for licensing of institutions of a type which qualify as SNFs, i.e., the distinct part would have to be licensed as a nursing home.
3202H - Survey Considerations  
(Rev. 1, 05-21-04)

Although an immediate survey is not mandatory, the SA must complete Form CMS-1539 promptly to report the change in size and location of the SNF or NF. Furthermore, the SA completes a spell of illness certification for any components of the institution or institutional complex that are being removed from inclusion in the SNF or NF. (See §2164.) If, in order to process this certification, the SA finds that survey is necessary, it may perform a full standard survey.

3206 - Existing ESRD Facility Relocation, Expansion, or Addition of New Service  
(Rev. 1, 05-21-04)

Enrollment with the FI and a new application is required when an ESRD facility relocates, expands, or adds a new service. An ESRD facility may relocate in order to expand because public transportation will make it more accessible to its patient population or because it wishes to add new services (see §2274).

3210 - CHOW of Providers and Suppliers  
(Rev. 1, 05-21-04)

Regulations covering CHOWs are at 42 CFR 489.18(ff).

The initial development of facts concerning a CHOW is made by the FI via the 855 process (see §2005.E). The FI sends the SA its recommendations and a final 855. After the SA concludes its fact-finding, it forwards the findings, with supporting documentation, to the RO with its recommendations for determination.

When a provider undergoes a CHOW, the provider agreement is automatically assigned to the new owner unless the new owner rejects assignment of the provider agreement. If the new owner rejects this assignment, the provider organization will not be able to participate in the Medicare program without going through the same process as any new provider, i.e., enrolling with the FI, applying for participation, undergoing Office of Civil Rights (OCR) clearance and an initial survey, having an effective date of participation assigned based upon regulation, etc. Automatic assignment of the existing provider agreement to the new owner means the new owner is subject to all the terms and conditions under which the existing agreement was issued. Terms and conditions include, but are not limited to:
3210A - Existing PoC  
(Rev. 1, 05-21-04)

The new owner must meet the timeframes for correcting deficiencies cited in the existing PoC. A CHOW is not a basis for extending the time given for correction. Documented evidence of effort and progress, and the absence of jeopardy to patient health and safety remain the only acceptable reasons for giving additional time for correction of deficiencies.

3210B - Compliance With Health and Safety Standards  
(Rev. 1, 05-21-04)

Assignment of an existing provider agreement assumes that a CHOW will have no adverse effect on patient health and safety. Consequently, a survey may not be required. If, however, there is any indication that patient care has deteriorated following a CHOW, the State must conduct a survey. If such a survey indicates noncompliance, the RO applies the enforcement action that is applicable to the provider/supplier type and appropriate to the level of noncompliance.

3210C - Compliance With Ownership and Financial Interest Disclosure Requirement  
(Rev. 1, 05-21-04)

Follow the guidance provided in §2005.F concerning enrollment and completion of the Form CMS-855.

3210D - Compliance With Civil Rights Requirements  
(Rev. 1, 05-21-04)

The RO notifies the OCR-RO of CHOWs of providers. Assignment of the existing provider agreement is not withheld pending civil rights clearance, and a new agreement can be issued before clearance by the OCR-RO is obtained. However, under these circumstances, a restricted provider agreement is issued with a contingency clause that states that if OCR clearance is not obtained, any payments made during the period will be recouped from the facility as of the effective date of the CHOW.

3210E - All Medicare Sanctions and Penalties  
(Rev. 1, 05-21-04)

Medicare sanctions and penalties are assigned to the new owner with the following exceptions:
1 - NATCEP

The restrictions preclude a State from approving (and requiring a State to withdraw from) Nurse Aide Training and Competency Evaluation Programs (NATCEPs/CEPs) offered by or in facilities that, within the previous two years, have been found to be out of compliance with certain CoPs. If there is a CHOW before such a 2-year restriction has run its course, whatever remains of the 2-year period will not be transferred to the new owner.

2 - Money Owed in Fraud Cases

The new owner is not responsible for money owed the Federal Government due to a determination that the previous owner is personally guilty of fraud. (However, if a determination of fraud is made against the corporation, and if the corporation is purchased and not incorporated as a new and separate corporation by the new owner, the new owner is subject to all Medicare penalties, sanctions, and liabilities.)

3210.1 - Determining Ownership
(Rev. 1, 05-21-04)

3210.1A - General
(Rev. 1, 05-21-04)

For certification and provider agreement purposes, the authorized official is an individual (such as independent practitioner or sole proprietor) or an appointed official (including, but not limited to, an officer, director, manager, general partner, limited partner, etc.) of a legal entity such as a corporation or general partnership who is directly responsible for the business enterprise and has been granted the legal authority to enroll it in Medicare, to make changes and/or updates to its status in the Medicare program, and to commit it to fully abide by the laws, regulations and applicable program memoranda and manual issuances of the Medicare program. This party is legally responsible for decisions and liabilities in a business management sense. The same party also bears the final responsibility for operational decisions made in the capacity of a “governing body” and for the consequences of those decisions.

Whether the owning party owns the provider enterprise premises or rents or leases them from a landlord or lessor is immaterial. Of course, if the owner enters into an agreement that allows the “landlord” to make or participate in decisions about the ongoing operation of the enterprise, this indicates that the owner has entered into either a partnership agreement or a management agency agreement instead of a property lease. A new partnership agreement constitutes a CHOW.

To determine ownership of any provider enterprise or organization, the SA determines which party (whether an individual or legal entity such as a partnership or corporation) has immediate authority for making final decisions regarding the operation of the
enterprise and bears the legal responsibility for the consequences of the enterprise’s operations.

CHOW processing is necessary for program participants that have Health Benefit Agreements or Provider Agreements in the Medicare program (hospital, SNF, HHA, hospice, CORF, OPT/SP providers and CMHC) because it must be determined who the responsible party is under the agreement. For the same reason, CHOW processing is necessary for supplier participants that have category-specific agreements with the Secretary (RHC, ASC, and FQHCs) or that must file cost reports (e.g., ESRD facilities). Somewhat less extensive CHOW processing is necessary for the remaining supplier types without agreements or cost report requirements (e.g., PXR) to ensure compliance with the statutory requirement for ownership disclosure and to ensure that the program has current, accurate records regarding participants.

3210.1B - SA Actions to be Taken Following CHOW
(Rev. 1, 05-21-04)

3210.1B1 - All Cases
(Rev. 1, 05-21-04)

The SA mails a set of initial certification forms to the new owner as soon as possible. (See Exhibit 63.) The SA sends Form CMS-1561 to the new owner for signature with a footnote that lists the original provider number, the name of the previous owner (the owner of record before the change of ownership), and his or her address, and it is placed in the empty space provided after the blocks furnished for the successor’s signature, title, and date. This serves to convey to each new owner at the outset that he or she is being assigned the previous owner’s provider agreement “subject to all the conditions specified in [the] agreement and 42 CFR Part 489, to include existing plans of correction....” This is important, because some providers have professed ignorance that they have been assigned the previous owners’ provider agreements, subject to the same terms and conditions that applied to the previous owners. This was a central point in the “U.S. v. Vernon Home Health, Inc.” case. Under 42 CFR 489.18(d) and early on any purchase of assets that involve the assignment of the provider agreement is subject to the relevant statutory and regulatory conditions. The new owners or prospective new owners must be clearly informed of their rights and responsibilities under the applicable Federal statutes and regulations, and it must be done as early in the process as possible to enable these individuals to make informed decisions.

Whenever an owner is contemplating or negotiating the sale of a provider, he or she notifies the SA, FI or the RO, as required in 42 CFR 489.18(b). The SA or the RO asks the prospective new owner if he or she intends to participate in the Medicare program, and if so, whether he or she intends to do so by accepting assignment of the previous owner’s provider agreement or by applying for a new provider agreement. This will prevent the confusion we have seen in the past and reduce the litigation. The new owner should be made aware that if the agreement is assigned to the new owner, the new owner is responsible for the former owner’s liabilities, including any Medicare payments. Also,
assignment of the agreement, in some cases, would result in the new owner receiving a Medicare underpayment. If the new owner states that assignment of the former owner’s provider agreement is not going to be accepted, but the new owner intends to continue the entity’s Medicare participation, inform the provider that there will be a break in the continuity of Medicare payment because CMS requires all new applicants to undergo a survey. If the new owner still does not wish to accept assignment, following a CHOW, the entity must enroll in the Medicare program as a new provider in accordance with the instructions found in §2005, and undergo the survey and certification process.

Sometimes the RO is unaware that a change of ownership has taken place until after the fact; sometimes months after the sales agreement has been consummated. In these cases, there is nothing that can be done except to ensure that the new owner understands the consequences of becoming a Medicare provider and accepting assignment of the previous owner’s provider agreement. Regardless of when the new owner is advised of the automatic assignment of the previous owner’s provider agreement (if the provider/supplier explicitly refuses to accept assignment, notify the RO immediately because payments may have already been made under the old provider agreement) the SA uses the footnote and has the new owner sign under the set of blocks on Form CMS-1561 labeled “Accepted for the Successor Provider of Services by.” This is an important step because it documents the fact that the new owner realizes that there is an assignment of the agreement. The SA informs the new owner of the requirement to submit the documents to its office no later than 2 working days after the consummation of the CHOW transaction, or, if the transaction occurred more than 2 calendar days ago, as soon as possible. The SA may accept the documents prior to the consummation of the CHOW transaction. However, because some CHOW transactions never are consummated, these documents should not be forwarded until the transaction has been completed. Similarly, the SA can only complete its processing after the CHOW date.

If the new owner indicates a desire not to participate in the program, the SA alerts the RO immediately by telephone and, if Medicaid is involved, the SMA. The SA obtains a written notice regarding the owner’s desire not to participate, and forwards it to the RO or to the SMA as appropriate.

A CHOW, per se, does not require a special survey. However, if new locations are added or different types of services will provided, we recommend that the SA may conduct a survey. If there is any reason to believe that the quality of services has deteriorated following the CHOW then the state must conduct a survey.

For portable x-ray suppliers, the SA obtains a copy of Form CMS-855B, and a statement from the new owner informing it of the CHOW effective date. Because there is no agreement to transfer in these cases and no cost report to be filed by the outgoing owner, it is not critical that the SA establish the CHOW date with the certainty required for providers and suppliers with Medicare agreements and/or cost reporting requirements.

For all providers, and suppliers with category-specific agreements (for example, RHC and ASC), and ESRD facilities, the SA obtains applicable Request to Establish Eligibility
form, an Expression of Fiscal Intermediary Preference, and documentation that proves a CHOW took place as well as exactly when it took place. The SA has all facilities complete an Expression of Fiscal Intermediary Preference form in every case to alert it to the instances in which the new owner is part of a CMS-recognized chain organization that uses a FI not commonly used in its State. Multi-regional chain operations due to their complexity are to be referred to the RO for adjudication in accordance with §3210.3. The SA includes on the Fiscal Intermediary Preference form a blank to be completed by the new owner indicating the ending date of the fiscal year that the new owner intends to use for purposes of Medicare and/or Medicaid reimbursement. The cost reporting year initially selected by the new owner must be used for cost reporting purposes and cannot be changed by the RO or SA. If the new owner decides that it wants to change its cost reporting year, it must do so in accordance with the requirements found in 42 CFR 413.24(f)(3). These requirements specify that the provider must submit a written request to their FI 120 calendar days before the close of the new reporting period requested by the provider and that a finding of good cause is made by the intermediary. Good cause would not be found if the FI determines the change would affect the initial date a hospital would be subject to the rate of increase ceiling or be paid under the prospective payment system.

For providers and suppliers with category-specific agreements, the SA obtains two signed originals of the applicable Medicare agreement form.

For providers, the SA obtains the applicable form required by OCR (HHS-441) and the appropriate attachments. See Exhibit 63 for a full listing of documents for the SA to submit to the RO.

**3210.1B 2 - CHOW during Termination Development**
(Rev. 1, 05-21-04)

The SA apprises the provider/supplier that termination actions already in process will not be postponed and the termination will only be avoided if compliance is attained. The SA notifies the RO by telephone of the CHOW and proceed to obtain and process the usual CHOW documents.

**3210.1B 3 - CHOW during “Reasonable Assurance Period”**
(Rev. 1, 05-21-04)

Following termination, a new owner may request approval for reentry, subject to operation of the facility for a certain period of time without recurrence of the deficiencies that were the basis for termination. The reentry may be through the SA survey process or through an accrediting organization recognized by CMS where appropriate under the regulations. The RO makes the determination as to whether the institution is eligible for readmission based on demonstrated compliance over a specified period of time. This “reasonable assurance period” will not be altered because a new owner takes over the provider organization or because the provider organization becomes accredited. If the
new owner wishes to proceed with reentry, the SA notifies the RO by telephone and obtains and processes the usual CHOW documents (see §2016).

**3210.1B 4 - Transfer Agreement Required of New Owner**  
(Rev. 1, 05-21-04)

For SNFs and NFs, a new owner will have to negotiate and submit a hospital transfer agreement relating to the new owning entity.

**3210.1B 5 – Relocation of Provider/Supplier Concurrent with CHOW**  
(Rev. 1, 05-21-04)

A new owner may propose to relocate the provider/supplier concurrent with the CHOW. If the relocation is to a site in a different geographic area serving different clients than previously served and employing different personnel to serve those clients, do not assign the agreement to the new owner. The provider/supplier must be treated as a new applicant to the Medicare program, rather than as an address change of an existing provider.

**3210.1C - Certification of Accredited Providers/Suppliers Which Change Ownership**  
(Rev. 1, 05-21-04)

While accreditation by a national accreditation body is not transferable to a new entity, accreditation does not automatically lapse when ownership changes. In the case of a provider or supplier that has been accredited by a national accreditation body, the provider/supplier must notify the accreditation body within 30 calendar days of the CHOW. Accreditation is continued until the accreditation body has determined whether a resurvey is necessary.

If an accredited provider/supplier is involved in a CHOW, the SA does not resurvey the provider/supplier. The SA secures the usual CHOW documents and forwards them to the RO.

If a participating provider/supplier that has been accredited by a national accreditation organization merges with a nonaccredited participating or nonparticipating provider/supplier, the accreditation organization may extend its accreditation to the nonaccredited provider/supplier as if it were deemed to meet the Medicare conditions. A CMS approved accreditation organization may extend its accreditation in a hospital it currently accredits. This could be a hospital that has undergone a change of ownership or a hospital that has expanded by acquiring another hospital or by establishing additional facilities. The JCAHO hospital accreditation program and all other CMS approved accreditation organizations have agreed to conduct surveys within 6 months the date of the extension agreement with the provider. The extension of the accreditation by the accreditation organization will serve in lieu of conducting a survey at this time.
3210.1D - CHOW Situations
(Rev. 1, 05-21-04)

Although 42 CFR 489.18 addresses only sole proprietorships, partnerships, corporations, and lease arrangements as CHOWs (as defined below) this does not preclude other transactions that constitute a CHOW (e.g. the creation of a Limited Liability Company). You should consult with your regional attorney when these other transactions are involved to determine if a CHOW has taken place. However, it is important for providers to understand that transactions that are considered a CHOW for purposes of certification may be treated differently for purposes of reimbursement under the program.

1 - Sole Proprietorship

If a provider of services is an entity owned by a single individual, a transfer of title to the enterprise to another person or firm, whether or not including transfer of title to the real estate, constitutes a CHOW. It is also a CHOW if the former owner becomes one of the members of a partnership or corporation succeeding him as the new owner.

2 - Partnership

In a partnership, the removal, addition, or substitution of an individual as a partner in the entity, in the absence of an express statement to the contrary (as permitted by State law) dissolves the old partnership and creates a new partnership and is a CHOW. State laws may vary regarding exceptions to this rule. If the surviving partners raise a question, the SA submits the facts without delay to the RO for a decision (or to the appropriate State authorities in a Medicaid-only case).

3 - Corporation

In an incorporated provider entity, the corporation is the owner. The governing body of the corporation is the group having direct legal responsibility under State law for operation of the corporation’s entity, whether that body is a board of trustees, a board of directors, the entire membership of the corporation, or is known by some other name. Even though one or more members of this governing body changes, and regardless of whether ownership of the corporation stock is transferred, there would not be a CHOW as long as the same corporation continues to be the legal entity responsible for operation of the provider organization.

A merger of one or more corporations with the Medicare-participating provider corporation surviving (i.e., a merger “into” the participating corporation) is not recognized as a CHOW of the surviving corporation. Also:

- If the corporation that survives is not the former owner of the provider entity, there is a CHOW; and
- Consolidation or merger of two or more corporations that results in the creation of a new corporate entity having ownership control over a provider organization constitutes a CHOW. The SA may need to refer to a corporation’s bylaws (on record with the State government) or board meeting minutes as it researches difficult cases.

4 - Leasing

When all or part of a participating provider facility is leased, it constitutes a CHOW. If only part of the provider is leased, the original provider agreement remains in effect only with respect to the unleased portion. The lease of part of the facility constitutes a change of ownership. The SA does a survey and prepares a certification covering the leased portion as a new provider.

When a new lease arrangement goes into effect at a participating facility, the SA obtains documents that indicate which individual or entity has first level authority over, and responsibility for, the provider located within the leased premises.

5 - Management Firm Operating Institution for Owners

A firm that contracts with the owners to manage an enterprise, subject to the owners’ general approval of operating decisions, is an agent of the owners rather than a partner or successor. If management in that sense is turned over to a management firm, this would not constitute a CHOW even though the management firm may appear to have wide latitude in making decisions, and even though its fee may be based on the net revenue or profit the facility receives from furnishing services.

The only case in which operation under a management agreement would constitute a CHOW is when the owner has relinquished all authority and responsibility for the provider organization. In questionable cases, the SA obtains and submits the management agreement to the RO along with its analysis and recommendation.

If a provider enterprise has been placed in the hands of a management firm and there is cause to suspect a problem in regard to patient care, the SA schedules and performs a survey.

6 – Franchise

If an entity states it is a franchisee of another entity which is the owner of the provider, do not enter into an agreement with either party or process a CHOW until you establish which entity is the provider which CMS will hold legally responsible for complying with the provisions of §1866 of the Act and applicable regulations. Please note that under 42 CFR 389.11 (see also §2781) if a provider wishes to participate in the Medicare program it must have an authorized official sign the provider agreement and submit a written statement indicating whether the provider has ever been adjudged insolvent or bankrupt or has such action pending.
A State licensing decision based upon a CHOW analysis conducted under the State’s criteria is not necessarily relevant to a Medicare CHOW determination. The Medicare determination must be made based exclusively on Medicare regulations and policies;

There can be no CHOW, i.e., transfer of Medicare participation, assignment of the provider agreement, and provider number, if there is no functioning provider enterprise in existence. If a provider ceases operations, it no longer meets the definition of any provider type and no longer has a right to a provider agreement or identification number;

As a rule, when a provider organization is sold, the Medicare provider number stays with it. A buyer is assigned the provider number and the provider agreement if the buyer purchases a participating provider organization. A provider number cannot be sold. A provider identification number is not the “property” of any individual or legal entity. The number is issued by the Medicare program and is under the control of the Secretary of DHHS, subject to law, regulation, and program policy;

To understand whether a CHOW took place in complex situations such as corporate reorganizations, it is often helpful to construct a simple “before and after” ownership diagram of the legal relationships among the owning entities and providers involved. The two-part diagram visually displays the ownership relationships as they appeared before and after the date of a possible CHOW;

In general, a CHOW recognized by the Medicare program is considered to have taken place at 12:01 a.m. on the date specified (i.e., in the first minute of the 24-hour day). Legal responsibility and the right to payment changes over when the clock moves past midnight into the CHOW effective date;

In general, the key date regarding a newly formed corporation is not the incorporation date (the date the corporation came into legal existence), but the date a provider was conveyed to the new corporation. Sometimes a new corporation becomes legally responsible for a provider the moment the corporation comes into existence, but there must be documented evidence that this is the case. It cannot be assumed;

It is not possible to know beforehand whether a CHOW will take place on a planned CHOW date. In every case, one must wait until after a proposed CHOW date to determine whether the planned CHOW event actually occurred. This means it is impossible to process a CHOW prior to the effective date; and
• The mere sale of any number of shares of an owning corporation does not constitute a Medicare CHOW because the responsible legal entity, the corporation, remains in place. For corporations that do not issue stock but are controlled by a “member” or “members” (which can be individuals, partnerships, or other corporations), the same principle holds true: a change in the individuals or entities controlling or owning the corporation is not relevant for CHOW purposes.

3210.2 - RO Role in CHOW Determination
(Rev. 1, 05-21-04)

The RO reviews the SA’s recommendation and, if it concurs that a CHOW has taken place, it annotates the certification file to reflect the change and:

• For providers and suppliers that require agreements, the RO issues a notice letter to the provider/supplier that recognizes the CHOW date. The RO sends copies of the letter to the SA, the SMA, the OCR-RO, and the FI(s). It is not necessary to acknowledge CHOWs that have taken place at the remaining supplier types;

• For providers and suppliers that require agreements, the RO signs the two original agreement forms previously signed by an authorized representative of the new owning entity and forwarded to the RO by the SA. The RO retains one original agreement form in its certification file and encloses the other with the notice letter to the provider/supplier;

• For providers and ESRD facilities, the RO prepares a Provider Tie-In Notice (Form CMS-2007) and forwards it to the FI. If there is a change in FI, the RO sends copies of Form CMS-2007 to the old and the new intermediaries. (See §3210.4);

• For all CHOWs, the RO should ensure that a kit of certification documents reflecting the change and all available current information is entered in the OSCAR/ASPEN system.

• As corporate structures have become increasingly complex, it has become more difficult for professionals in nonlegal entities to discern all of the management and control relationships in individual situations and to determine accurately whether or not a CHOW has occurred for purposes of the Medicare program. The RO should refer all CHOW determinations that are in any way complicated by the intricacies of the case or otherwise require professional legal expertise to the appropriate Regional Attorney.
3210.3 - CHOWS Involving Multi-Regional Chain Organizations
(Rev. 24, Issued: 01-26-07; Effective: 10-01-05; Implementation: 01-26-07)

When a CHOW involves a multi-regional chain organization, a lead or coordinating RO is designated. For certification purposes, this will typically be the RO serving the State in which the headquarters of the chain is located. The coordinating RO will notify all affected ROs of its lead role, make a CHOW determination that is uniform for all regions, and notify all affected ROs of the determination.

Exceptions to the lead RO procedure can occur. In an example involving 100 commonly-owned hospitals in eight regions, not all of the transactions met the definition of a CHOW under the regulations. Many of the transactions required individual, detailed analysis. In this case, no coordinating RO was designated because no uniform circumstances existed. In any situation in which the RO believes a coordinating RO may need to be designated, CO must be contacted for guidance.

New providers that belong to CMS-recognized chains have the option of being assigned to the local designated FI or to the FI that serves the chain home office.

The CORFs, CMHCs, OPT facilities, rehabilitation facilities, and ESRD facilities will be assigned to the local designated FI. This group of providers will also not be able to become chain organizations or receive single FI status. However, some exceptions have been made for ESRD facilities. Exceptions are made on a case-by-case basis where the ESRD provider makes a compelling argument that to become a chain organization or have single FI status would be in the best interests of the Medicare program.

3210.4 - Other Changes Related to CHOW - RO Procedures
(Rev. 24, Issued: 01-26-07; Effective: 10-01-05; Implementation: 01-26-07)

The new owner of a provider has the opportunity to change the provider's fiscal reporting period. The provider sends a written request to its fiscal intermediary. The fiscal intermediary reviews the request to determine if good cause exists, and will either approve or deny the request. See §3210.4B.

There are also some instances in which the RO must assign a different provider number. See §3210.4C.

3210.4A - New Owner Requests Different Intermediary
(Rev. 24, Issued: 01-26-07; Effective: 10-01-05; Implementation: 01-26-07)

All participating providers that change ownership and accept assignment of the existing provider agreement will continue with the same fiscal intermediary that served the previous owner. If the new owner does not accept assignment of the provider agreement,
the new provider will be treated as a new applicant to the Medicare program and once all Federal requirements have been met, a new provider agreement with a new provider number will be issued to the new owner. The new owner will then be assigned to the local designated FI.

For CORFs, CMHCs, OPT facilities, rehabilitation facilities, and ESRD facilities, please refer to §3210.3.

3210.4B - New Owner Sets Different Fiscal Reporting Period
(Rev. 24, Issued: 01-26-07; Effective: 10-01-05; Implementation: 01-26-07)

At the time of a CHOW, the new owner must select its cost reporting year for purposes of reimbursement under the program. The new owner may file its initial cost report covering a period of at least 1 month of provider operations under the program, but no more than 13 months of provider operations under the program.

The SA reports to the RO what cost reporting year the new owner has selected for use, and the RO indicates the selected cost reporting year on Form CMS-2007. Subsequent requests for changes in the fiscal reporting period must be made in writing by the provider to the FI. A change in the cost reporting period will be made only after the FI has established good cause. Good cause exists if there is a good reason or justifiable purpose in seeking a change in the cost reporting period. Neither the RO nor the SA can change the cost reporting year of a provider.

3210.4C - New Provider Number Must Be Issued
(Rev. 1, 05-21-04)

In most cases, the identification number previously assigned to the provider organization stays with the provider organization, regardless of which legal entity is the owner. A new number is not assigned based on a CHOW. There are exceptions, however. These exceptions involve cases in which the form of the provider number indicates a particular status of the provider or supplier to which it is assigned, and that status has changed as a result of the CHOW. For example, a hospital-based ESRD facility becomes freestanding or vice versa. The RO should note in the CHOW notice letter that one identification number is being retired effective with the CHOW date and a new one is being issued based on the change in provider/supplier status. The RO also includes this notice on Form CMS-2007.

Before issuing a CHOW acknowledgement letter, the RO should review §2779, “RO Assignment of Provider and Supplier Identification Numbers,” to be sure the previously-assigned identification number continues to be appropriate. For additional instructions regarding provider identification numbers in merger and CHOW situations, see §2779. Following are the most likely examples of the need to issue a new identification number as a result of a CHOW:
1 - ESRD

An ESRD facility can be classified as hospital-based and be located on the hospitals’ premises (having an identification number in the series 00-2300 through 00-2499), as a hospital satellite off the hospitals’ premises (having an identification number in the series 00-3500 through 00-3699), or as non-hospital/independent (having an identification number in the series 00-2500 through 00-2899). If the ESRD facility’s classification changes as a result of a CHOW, the RO retires the facility’s original number and issues a new number in the appropriate form, effective with the CHOW date. If the facility’s location changes as a result of a CHOW, the RO has the SA conduct a survey of the new location.

2- RHC

A RHC can be classified as freestanding (having an identification number in the series 00-3800 through 00-3974 or 8900-8999), or as provider-based (having an identification number in the series 00-3975 through 00-3999, 00-3400 through 00-3499, or 00-8500 through 00-8899). If the RHC’s classification changes as a result of a CHOW, the RO retires the original number and issues a new number in the appropriate form, effective with the CHOW date.

3210.5 - New Owner Refuses to Accept Assignment of the Provider Agreement
(Rev. 1, 05-21-04)

3210.5A - New Owner Refuses to Accept Assignment of Previous Owner’s Provider Agreement
(Rev. 1, 05-21-04)

A new owner may refuse to accept assignment of the previous owner’s provider agreement, which means that the existing provider agreement terminated effective with the CHOW date. The refusal to accept assignment must be put in writing by the new owner and forwarded to the RO 45 calendar days prior to the CHOW date to allow for the orderly transfer of any beneficiaries that are patients of the provider. The refusal can take the form of a letter initiated by the prospective owner or can be indicated in response to a letter sent to the new owner by the RO or the SA that is designed to document the new owner’s desire to continue program participation.

In all cases of refusal to accept assignment, all reasonable steps must be taken to ensure that beneficiaries under the care of the provider are aware of the prospective termination of the agreement. In this situation, there may be a period when the facility is not participating and beneficiaries must have sufficient time and opportunity to make other arrangements for care prior to the CHOW date.

After the CHOW has taken place, the RO acknowledges the refusal to accept assignment in a letter to the new owner, with copies to the SA and the FI. The RO completes a Form
CMS-2007 with the date the agreement is no longer in effect, noting that the termination is due to the new owner’s refusal to accept assignment of the provider agreement.

It is the responsibility of a prospective purchaser of a Medicare provider to know that it can refuse to accept assignment of the provider agreement and that it should formally indicate its choice in that regard. If, however, the CHOW goes into effect without a refusal or acceptance of assignment on record, the RO concludes that the agreement has been automatically assigned to the new owner and completes processing of the CHOW.

If the new owner refuses to accept assignment after the date the CHOW has taken place, the RO should contact its regional attorney for guidance.

If a new owner refuses to accept assignment and also wishes to participate in the Medicare program, the RO first processes the refusal as indicated above and then treat the new owner as it would any new applicant to the program: obtain and process application documents, have the SA perform an initial survey and, if all requirements for participation are met, assign an effective date of participation based upon the applicable regulation. (See 42 CFR 489.13.)

The earliest possible effective date for the applicant is the date the RO determines that all Federal requirements are met. The Federal requirements include, in addition to the CoP, enrollment as described in §2005, capitalization (HHAs), and any other special requirements such as the special provisions for psychiatric hospitals at 42 CFR 482.60. The aforementioned requirements are the same regardless of whether the new owner operates a non-accredited facility or is seeking Medicare compliance with the CoP via deemed status.

As mentioned above, these requirements include enrollment of the provider in accordance with the instructions in §2005. The Form CMS-855 must be submitted prior to the CHOW date. However, the subsequent survey of the new applicant must be performed (1) after the CHOW, because the provider agreement of the former owner terminates effective with the CHOW date and the new owner must be treated as a new Medicare applicant; and (2) after the FI makes a recommendation to CMS for approval in accordance with the current procedures. If for any reason the accrediting body of the entity seeking deemed status chooses not to conduct or to delay a survey of the new entity, CMS will inform the entity that is will be unable to participate in the Medicare program until a survey is conducted and CMS is assured that the new entity meets all applicable health and safety requirements. In such a circumstance the new applicant may choose to have the SA conduct its survey.

In addition to the policies articulated above and in §3210.1.C relating to accredited providers, the following policies apply.
Hospitals and Units Excluded from Medicare’s Prospective Payment System (PPS)

Accreditation by itself does not determine whether to exclude a hospital or unit from Medicare PPS (see Subpart B of 42 CFR 412). All PPS exclusion determinations must be made by the appropriate CMS RO based on the facts at the time the decision is made. PPS hospitals with such units must also be surveyed by the SA to determine if exclusion requirements are met by the new owner following the CHOW.

Rehabilitation Unit

If the rehabilitation unit is properly accredited as a rehabilitation program (§§3100-3112.3), the SA must verify only the requirements for the rehabilitation director at 42 CFR 412.23(b)(5). The FI must make its determination regarding the “75 percent rule” (See 42 CFR 412.23(b)(2)). If the rehabilitation program is not accredited, the rehabilitation unit must be surveyed onsite by the SA for compliance with the requirements at 42 CFR 412.25 and 412.29.

Psychiatric Unit

The SA must perform an onsite survey to determine if the psychiatric unit complies with the PPS exclusion criteria at 42 CFR 412.25 and 412.27 on or after the effective date of the CHOW. The FI must re-verify that other criteria at Part 412 are met.

CHOW of a Hospital within a Hospital

The non-assignment of a Medicare provider agreement involving a CHOW of a PPS excluded hospital within another hospital (HWH) may affect the PPS exclusion of the HWH if the contractual agreements between the HWH and the new host hospital have changed. If there has been a change, the HWH will lose its exclusion unless the agreements between the two hospitals are renegotiated with the new owner, and the HWH must submit updated evidence of compliance with the regulation to the RO via the SA. In addition, if the HWH is not in compliance with the CoP, it is subject to a loss of “deemed status” (if accredited) and placed under SA monitoring. It could subsequently be terminated if compliance is not achieved.

Also, when a host PPS hospital containing a PPS excluded HWH undergoes a CHOW, the HWH must be notified that it must show CMS that it complies with 42 CFR 412.22(e)(5) with the new provider within 30 calendar days of the CHOW date.

Offsite Location Based to Provider that Undergoes a CHOW and Non-Assignment of Provider Agreement

In the case of a provider with other providers or entities based to it that undergoes a CHOW with a new owner who chooses not to accept assignment of the current provider agreement, the provider-based status of the other providers or entities ends with the termination of the former owner’s provider agreement. The request for Medicare
approval of the new owner must include information related to other providers or entities if the new owner intends for these entities to meet the provider-based criteria found in §2004. In the case of hospitals with multiple components that operate as a single hospital, both §§2024 and 2004 apply. It is important to note that §2024 would be inapplicable to provider-based entities other than hospitals with multiple components that operate as a single hospital. The new owner must once again justify to the SA, FI and CMS that these provider-based entities meet CMS’ provider-based criteria. The accreditation body, and the SA as appropriate, must consider these entities in conducting the survey of the new provider for compliance with the CoP or accreditation standards. Specifically, the RO must ensure that all off-site entities that claim to be provider-based comply with the Medicare CoP, or in the case of accredited hospitals, comply with standards that are at least equivalent to the Medicare CoP.

CHOW Involves a Related Organization

In situations where the CHOW is an organization related to the former owner (e.g., CHOW from general corporate ownership to subsidiary corporation, limited partnership or other related entity) with assumption of the provider agreement the SA or the RO may wish to include the following paragraph in the notice to the provider if you believe there may be outstanding liabilities of the former owner. “The change of ownership does not release the former owner or successor owner from liabilities resulting from past provider operations. The former owner may be liable for overpayments, penalties and other payments arising from the period it owned the provider. In addition, successor owners have joint and several liability for these debts notwithstanding divestiture of assets by the former owner.”

In situations where the new owner fails to notify you of a CHOW timely, treat as an assigned agreement (§3210.5).

3210.5B - Withdrawal After CHOW - Provider
(Rev. 1, 05-21-04)

If, after a CHOW takes place, the RO receives notice that the new owner of a provider desires to withdraw from the program, the RO consults with the new owner to set a withdrawal date designed to protect the health and safety of program beneficiaries who may be patients of the provider. The RO sets a withdrawal date of up to 6 months beyond the provider’s notice of intent to withdraw. Under these circumstances, the RO processes a complete CHOW notice and a withdrawal.

3210.5C - CHOW and Withdrawal - Supplier
(Rev. 1, 05-21-04)

If the new owner of a supplier declines to participate, the RO negotiates a withdrawal date that does not disadvantage any program beneficiaries that the supplier may be serving. The RO processes the supplier withdrawal as usual.
Expansion of Services

3220 - Certifications of Additional Services
(Rev. 1, 05-21-04)

Several categories of providers/suppliers require specific approval prior to becoming eligible to receive Medicare and Medicaid payment for certain services beyond those for which they were initially approved. The specific provider/supplier types affected by this requirement are:

- HHAs;
- RHCs; and
- ESRD facilities.

During the initial survey and resurveys, the SA advises the provider/supplier to inform it and the appropriate FI or carrier promptly in writing when an additional service is contemplated, so that it can evaluate compliance with the pertinent CoPs or Conditions for Coverage. Do not accept oral requests. When the SA is notified by the provider or supplier or the FI or carrier of the addition by the provider/supplier or learns that a service has been added, it reviews applicable documentation and, as necessary, performs a survey of the new service promptly. The SA records the results on the appropriate survey report.

The only services of an HHA that when added would require a survey are OPT and speech language pathology services are when these services are provided by the HHA as a provider of outpatient physical therapy services (See 42 CFR 484.38 and §3222.) This usually does not require an immediate survey, but can be surveyed at the time of the next standard survey.

3220A - Services in Compliance
(Rev. 1, 05-21-04)

If a new survey is performed and the service meets the applicable Conditions, the SA recertifies the provider/supplier as continuing to meet the CoPs or Conditions for Coverage and complete Form CMS-1539. It includes an explanation in Item 17 concerning the added service and an evaluation of the new service.
3220B - Services Not In Compliance (HHAs, RHCs, and ESRD Facilities)
(Rev. 1, 05-21-04)

If a new service does not meet the applicable Condition(s), the SA informs the provider/supplier that it must either come into compliance or stop providing the service to avoid termination action.

If the provider/supplier agrees to stop providing a service, the SA notes this action in its files and follows up within 60 calendar days to ascertain whether the service has been discontinued. If the provider/supplier is unwilling to correct or stop providing the service, the SA initiates termination action.

3222 - Specific Requirements for Expansion of Services
(Rev. 1, 05-21-04)

3222A - HHA’s Request to Provide OPT Services on Its Premises
(Rev. 1, 05-21-04)

An HHA may provide OPT services on its premises as well as in patients’ homes. The HHA providing services on its premises must meet and should be surveyed for compliance with 42 CFR 485.723 and 485.727. These regulations do not apply to physical therapy services provided in patients’ homes.

3222B - RHC’s Request to Provide Visiting Nurse Services
(Rev. 1, 05-21-04)

After the clinic is approved as an RHC, it may also seek approval to provide covered visiting nurse services. An RN, LPN, or licensed vocational nurse must furnish these services. (See 42 CFR 405.2411, 405.2416, and 405.2417.)

When a request is received, the RO must determine whether there is a shortage of HHAs in the area. If there is an existing HHA furnishing services in the RHC area, the SA contacts the HHA for a statement of its ability or inability to adequately furnish nursing services in the area. In addition, the SA obtains information from the local or State health planning organization. The SA transmits the request and all pertinent documentation to the RO. The SA does not approve the visiting nurse services at this point.

If the RO determines that there is not a shortage of home health services for the area, authority to furnish visiting nursing services to homebound patients will be denied, and
the RHC will be expected to refer its homebound patients to the HHA serving the area. The SA will receive a copy of the RO determination notice.

For purposes of this development, a “homebound individual” is one permanently or temporarily confined to his/her place of residence because of a medical or health condition. The individual may leave the place of residence infrequently and still be considered homebound. However, an individual in a hospital or long term care facility is not “homebound” for purposes of visiting nurse services.

If the RO determines that there is a shortage of home health services, it will request that the SA evaluate the qualifications of RHC personnel who are responsible for delivery of nursing services.

The SA completes the applicable sections of the Rural Health Clinic Survey Report (Form CMS-30) for visiting nurse services and a written plan of care. The SA reviews records (plans of care and other appropriate records) to verify that such services are provided to homebound individuals and are furnished under written plans of care developed and signed by the supervising physician, nurse practitioner, physician assistant, or nurse midwife and reviewed by the supervising physician at least every 60 calendar days. If the service has recently been implemented, it may be necessary for the SA to follow-up later to determine that the 60-day requirement is being observed.

When the above development has been completed and evaluated, the SA completes a supplementary certification for the visiting nurse service.

Follow the procedure in §2274. (Also see Exhibit 27.)

3224 - Addition of Sites to an Existing Provider
(Rev. 1, 05-21-04)

It is inherent in the provider certification process that a provider give notification to CMS of its proposal to expand its service area by adding a branch, satellite or extension location. The Medicare statute and applicable regulations are implicit that the proposed expanded service area meet the Conditions of Participation the same as the primary location that has signed the provider agreement or that has been assigned a provider number or both. In the absence of notification, CMS has no way of determining whether the requirements critical to health and safety are met at the expanded location. For example, a hospice’s request for satellite location may be denied because it cannot demonstrate how the hospice will assume administrative and supervisory responsibility for the services provided at the expansion site. Moreover, there is no basis for a provider to bill Medicare for services provided by a site which has not been determined to meet applicable requirements of participation.
When an expansion request is received, before making a determination the RO considers the following:

- Whether the proposal meets Medicare statutory and regulatory requirements. For example, in the case of an HHA, does the proposed branch meet the definition of a branch office at 42 CFR 484.2. If it is possible to make a decision based on the provider’s description of how it intends to operate, an onsite survey may not be necessary.

- If the proposal complies with State and local laws related to the particular type of provider/supplier; and

- Whether Medicare reimbursement is affected by the proposal. For example, a hospital states that it has purchased a physicians’ clinic that is now a part of the hospital. In such a case, input from the Division of Medicare and the fiscal intermediary will likely be necessary. While CMS does not dictate to a provider how it should operate its business, the provider does have to comply with Medicare requirements. Whenever an entity can meet the requirements of two different categories; e.g., subunit and independent home health agency, it is generally CMS’ policy to designate the category for which there is the least potential to increase Medicare costs. If a proposed branch is in an area that would receive a different payment rate than the parent HHA, it could be found to be in a different geographic area and determined not to be a branch.

Although legal authority exists for conducting a survey, a survey may not be necessary because the provider furnishes the RO with sufficient information to make a determination about its proposed expansion either at the time of its initial request or subsequently. If the RO believes a survey is required, but the SA is unable to conduct a survey within a reasonable period of time, the RO may take one of the following actions:

- Make a determination based on the expansion information provided by the provider and inform the provider of the decision; and

- Inform the provider that a survey will be necessary and that it should not bill Medicare for services provided at the proposed expansion location until the survey is conducted and a determination is made.

In the absence of notification of an expansion, CMS has the authority to deny bills for services furnished at the expanded site. When notification is received of a proposed expansion, the RO should inform the provider of whether the expanded site meets applicable requirements. The fiscal intermediary should be notified of the RO’s decision.
Validation Surveys of Accredited Providers and Suppliers

3240 - Validation Surveys - General
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

A “deemed” provider or supplier means a provider or supplier which has voluntarily applied for and has been accredited by a CMS-approved Medicare accreditation program whose recommendation for deemed status for that provider or supplier has been accepted by the RO. A list of current CMS-approved Medicare accreditation programs may be found at: https://www.cms.gov/Medicare/Provider-Enrollment-and-Certification/SurveyCertificationGenInfo/Downloads/AOContactInformation.pdf

NOTE: For the purposes of this section, the term “Condition” refers to any Condition of Participation, Condition for Coverage, or Condition for Certification. (See Chapter VI for discussion of validation surveys for accredited laboratories.)

SAs may not conduct a Medicare survey of a deemed provider or supplier unless specifically authorized by the RO to do so. Sections 1864(c) and 1865 of the Act provide the basis for conducting validation surveys of deemed providers and suppliers. Regulations authorizing such surveys are found in 42 CFR Part 488. The CMS may require a survey of a deemed provider or supplier to validate the accreditation organization’s process. These surveys will be conducted on either a representative sample basis (i.e., representative sample validation survey), or in response to a substantial allegation of noncompliance (i.e., complaint survey).

A representative sample validation survey is usually a standard or full survey of all applicable conditions.

A substantial allegation/complaint survey is authorized by the RO in response to a credible allegation which, if substantiated, would result in a condition-level citation. The RO advises the SA which conditions are to be assessed for compliance, based on the nature of the complaint. See Chapter 5, Sections 5100 – 5110 for more information on substantial allegation survey policies, procedures and timeframes.

The Form CMS 2802 is transmitted from the RO to the SA via the ASPEN Central Office (ACO) system, for a representative sample validation survey, or via the ASPEN Complaint Tracking System (ACTS) for a substantial allegation validation survey. The RO uses this form to:

- authorize the SA to conduct a validation survey of a particular provider/supplier;
- identify the applicable AO(s);
- indicate the type of validation survey to be conducted;
- indicate, when applicable in the case of a representative sample validation survey, the AO’s survey end-date; and
- in the case of a substantial allegation validation survey, identify the specific conditions for which the SA must assess compliance.
If a provider or supplier selected for a validation survey (representative sample or substantial allegation) is found to have one or more condition-level deficiencies, it will no longer be deemed to meet the Medicare Conditions. The RO advises the provider or supplier that its deemed status is removed and that it is being placed under SA jurisdiction. However, no change is made to the provider’s or supplier’s deemed status in the Automated Survey Process Environment (AS Pen). Instead, the placement of the provider or supplier under SA jurisdiction is noted under the Deeming tab within the certification kit in ACO.

3241 - Objective of Validation Surveys
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Validation surveys are intended to develop a reasonable estimate of an accreditation organization’s performance. Validation surveys are to be conducted in accordance with the survey protocol for the provider/supplier type being surveyed to assure a fair basis for comparing the effectiveness of CMS-approved Medicare accreditation programs.

3242 - Representative Sample Validation Surveys of Deemed Providers/Suppliers
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

The deemed provider/supplier validation survey process is designed to evaluate the performance of a CMS-approved Medicare accreditation program by determining whether a provider/supplier that is deemed to be in compliance with the Medicare conditions on the basis of its accreditation and is, in fact, meeting Medicare health and safety requirements. Such validation surveys are authorized in accordance with §1864 of the Act. Moreover, CMS is required under §1875(b) of the Act to provide an annual report to Congress on the performance of all CMS-approved Medicare accreditation programs approved under Section 1865 of the Act, including a validation of the accreditation process.

The SA conducts validation surveys of deemed provider/supplier types in accordance with established survey protocols for the provider or supplier type.

CO selects a representative sample of deemed providers and suppliers for the SA to conduct a validation survey and forwards this listing each month to the ROs. Upon receipt of the monthly list, the RO electronically issues the applicable Form CMS 2802, depending on the type of provider or supplier, promptly to the SA via the ACO system deeming tab in the recertification/validation kit. The RO also provides CO via the designated Sharepoint site a copy of the Form CMS 2802 once it is issued, for tracking purposes.

The representative sample validation survey is a standard, i.e. full, survey of ALL conditions applicable to that provider or supplier type. Unless the survey is a mid-cycle survey (see below), representative sample validation surveys must be conducted within
60 calendar days following the scheduled end date of the accreditation organization survey. The CO advises the RO of the AO scheduled survey end date, and the RO shares this information with the SA via the electronic version of the applicable Form CMS 2802. The SA must not share this information with any outside parties. The SA must also not request a copy of the AO survey report from the AO, in order to avoid bias in the SA’s conduct of an independent survey of the provider or supplier.

In addition to the traditional representative sample validation surveys, the CO sample selection may include deemed facilities at various points in their accreditation cycle (e.g., mid-cycle). Generally the assignment of such surveys is not tied to the timing of a scheduled AO survey, and thus there is no AO survey end date that determines the timing of the SA’s validation survey. The CO will include any additional instructions (if applicable) for the conduct of such surveys, including the timeframe for completion of the survey by the SA.

3243 - Substantial Allegation Validation Surveys of Deemed Providers/Suppliers
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

A substantial allegation/complaint survey is authorized by the RO in response to a credible allegation which, if substantiated, would result in a condition-level citation. The RO advises the SA which conditions are to be assessed for compliance, based on the nature of the complaint. The SA conducts the complaint investigation in accordance with the established protocols for the provider or supplier type. It is not sufficient for the SA to review only the medical record(s) related to the specific complaint allegation; rather, the SA must assess the provider’s/supplier’s general, current compliance with each condition specified by the RO on the authorizing Form CMS 2802 transmitted in ACTS.

See Chapter 5, Sections 5100 – 5110 for more information on substantial allegation survey policies, procedures and timeframes.

3244 - SA Preparation for Validation Survey
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

The SA follows standard survey processes when conducting validation surveys. For example, all validation surveys are unannounced. All substantial allegation validation surveys must include an assessment of compliance with the condition(s) specified by the RO; it is not acceptable for the SA to review only the record and other information related to the specific complaint.

The SA assigns surveyors who normally conduct surveys of non-deemed providers or suppliers to conduct validation surveys for the same provider/supplier types. The size of the survey team and the time that the SA spends on-site must be comparable to that used by the SA for non-deemed providers/suppliers of similar size or complexity. Whenever possible, all team members should survey a provider or supplier concurrently. This applies to staff who conduct the LSC portion of a standard survey as well as to personnel...
who conduct the health portion of the survey. If situations occur where this is not feasible, the SA must contact the RO for further direction.

As with any other Federal survey, CMS personnel may be present during the survey to provide assistance and to help assure nationwide uniformity and validity.

3246 - Provider/Supplier Authorization for Validation Survey
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

During the entrance conference for either type of validation survey the SA presents to the provider or supplier a letter signed by the SA Director announcing the validation survey (Exhibit 37), as well as an “Authorization by Deemed Provider/Supplier Selected for Accreditation Organization Validation Survey,” (Exhibit 287). The SA requires the signature of the provider/supplier CEO or other authorized individual on the authorization document, acknowledging that the provider/supplier must permit the validation survey by the SA to take place, as well as SA monitoring of the correction of any substantial noncompliance found through the validation survey.

3248 – Provider/Supplier Refusal to Permit Validation Survey
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

If, after efforts have been made to explain the validation survey protocol procedure, the provider/supplier refuses to permit the validation survey to take place, the SA informs the provider/supplier that its “deemed” status will be removed and the provider/supplier may also be subject to termination from the Medicare program (and, where applicable, the Medicaid program. [see ref. 489.53(a)(4)] If the provider/supplier continues to refuse to permit the validation survey, the SA notifies the RO. The SA informs the RO of all efforts made to encourage compliance. The RO notifies the provider or supplier that its participation in Medicare will be terminated, following standard termination procedures.

3252 - SA Forwarding Validation Survey Records to RO
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Representative sample validation surveys: The SA submits the appropriate information as specified on the List of Documents in Certification Packet (see Exhibit 63) to the RO or through an update to the ASPEN database within 30 calendar days of completing the survey. In cases where immediate jeopardy exists, the SA submits all the appropriate information specified on the List of Documents in Certification Packet to the RO within two working days of completion of the survey, and enters survey data to the ASPEN database, in a timely manner. The SA also completes the Survey Team Composition and Workload Report (Form CMS-670).

Substantial allegation validation surveys: The SA follows the procedures in Chapter 5, Section 5110.
If the provider/supplier has been cited as a result of the validation survey for deficiencies at the standard level only, the provider/supplier remains deemed. It is not obligated to submit a plan of correction (PoC), although it may voluntarily choose to do. If the provider/supplier submits a PoC, the SA includes it in the survey file.

3254 - RO Actions Following Validation Survey
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

Upon receipt of survey materials, the RO analyzes and considers the SA findings and recommendations and determines whether it agrees with them. If the RO disagrees with the SA’s findings or recommendations, it may require the SA to revise the Form CMS 2567 before it is issued to the provider or supplier. The RO takes necessary action to make a compliance determination and notify the provider/supplier as soon as possible, in the case of an immediate jeopardy, and within 30 calendar days for all other cases. The RO inputs the information into ASPEN in a timely manner.

In the case of a representative sample validation survey only, once the survey report is finalized the RO forwards to CO via the designated Sharepoint site a copy of the Form CMS 2567 and the accompanying correspondence to the facility.

If the validation survey identifies substantial noncompliance, i.e., condition-level deficiencies, the RO removes the provider’s or supplier’s deemed status. The provider/supplier continues to be accredited by its accreditation organization and is still permitted to participate in the Medicare/Medicaid programs while correcting the deficiencies. However, until the provider/supplier either achieves substantial compliance or its Medicare agreement is terminated, it is subject to the same requirements, survey and enforcement procedures that are applied to non-accredited provider/suppliers found out of compliance following a survey. The SA monitors the provider/supplier until it reaches substantial compliance with all conditions or it is terminated from the Medicare program, and where applicable, the Medicaid program.

3254A – Providers/Suppliers Found in Compliance Following Validation Survey
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

If the provider/supplier is in compliance with all Medicare Conditions, the RO notifies the provider/supplier and sends a copy to the SA and the AO.

3254B - Providers/Suppliers Found Not In Compliance With One or More Conditions Following Validation Survey and Noncompliance Constitutes Immediate Jeopardy
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

If the SA finds deficiencies that pose an immediate jeopardy to patient’s health and safety, the SA immediately notifies the RO by phone and then forwards the survey packet
to the RO within two working days after the completion of the survey. If after the RO review of the survey packet, the RO determines that there are deficiencies that pose immediate jeopardy (IJ) to patient health and safety, the provider or supplier will be placed on the 23-day termination track. See Chapter 5, section 5110.2 for detailed procedures and timeframes for substantial allegation validation surveys.

In the case of a representative sample validation survey the RO removes the provider’s/supplier’s deemed status and places it under SA survey jurisdiction. The procedures and timelines in Chapter 5, Section 5110.4b apply.

### 3254C - Condition-level Deficiencies That Do Not Pose Immediate Jeopardy
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

If the SA determines, and the RO agrees, that the provider/supplier is out of compliance with one or more Conditions, but the deficiencies do not pose immediate jeopardy to patient health and safety, the SA and RO follow the procedures in Chapter 5, Section 5110.3 for substantial allegation validation surveys. For representative sample validation surveys, the RO removes the provider’s/supplier’s deemed status and places it under SA survey jurisdiction. The procedures and timelines in Chapter 5, Section 5110.4c apply.

### 3254E - Plans of Correction
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

See §2728B for discussion of the requirements for an acceptable PoC, as well as the options available to the provider/supplier.

See Chapter 5, Sections 5100 – 5110 for information on when the RO must review an SA recommendation on whether or not a PoC is acceptable and when the RO must send a notice to the provider/supplier with a copy to the applicable AO(s).

### 3254F - Termination
(Rev. 1, 05-21-04)

The CMS will terminate a provider/supplier if it does not submit an acceptable PoC, or if after a reasonable period of time, it does not correct the Conditions that have been determined to be noncompliant. The RO obtains copies from the SA of the latest survey material before proceeding with termination procedures.

### 3254G - Compliance with All Conditions After Correction of Deficiencies
(Rev. 1, 05-21-04)

When an accredited provider/supplier is determined to be in compliance with all Conditions, the RO notifies the provider/supplier accordingly (and where applicable, the
SMA). The RO informs the SA, in writing, to cease monitoring activities. Revisits by the SA are not authorized after an accredited provider/supplier has been notified that it is back in compliance with all Medicare Conditions, and its deemed status is reinstated.

3256 - RO Provision of Information to Accrediting Organizations
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

The RO must provide the AOs copies of every Form CMS 2567 issued to a deemed facility accredited by the AO(s) in a timely manner, i.e., as soon as possible after issuance to the provider/supplier. ROs have the option of sending this material to AOs electronically rather than mailing hard copies. Copies to be provided include all Form CMS 2567s resulting from a SA validation survey, including both representative sample and substantial allegation (complaint) surveys, as well as all Form CMS 2567s issued after the provider’s or supplier’s deemed status has been removed and the facility has been placed under SA jurisdiction until it can achieve substantial compliance. Copies of any correspondence sent to the provider or supplier related to a Form CMS 2567 must also be provided to the applicable AO(s), as well as correspondence concerning the status of the provider’s supplier’s Medicare provider agreement or supplier approval (e.g., issuance of an agreement/approval with CCN, notice of potential termination of the agreement/approval, notice of termination of the agreement/approval, etc.).

Note that a provider or supplier may be accredited under a CMS-approved Medicare accreditation program by more than one AO; in such cases all AOs must receive copies.

Forward copies to the applicable AO(s) using the contact information found at https://www.cms.gov/Medicare/Provider-Enrollment-and-Certification/SurveyCertificationGenInfo/Downloads/AOContactInformation.pdf

Where RO resources and workload resources permit, the RO should consider sending copies of this information electronically to the AO contact’s e-mail address.

3257 - Reinstatement to Accrediting Organization Jurisdiction
(Rev. 123, Issued: 10-03-14, Effective: 10-03-14, Implementation: 10-03-14)

A provider/supplier that has been under SA monitoring is returned to deemed status when it is determined that it is in substantial compliance with the applicable conditions. The provider or supplier is no longer monitored by the SA but is instead under the jurisdiction of the AO. The deeming tab in ASPEN is updated to reflect the end of the SA monitoring.

The SA may not conduct any further surveys of the provider/supplier without a new authorization from the RO via the applicable Form CMS 2802.

When the provider/supplier is returned to the accreditation organization’s jurisdiction, the RO notifies the provider or supplier in writing, with a copy to the applicable AO(s).
Termination

A CMS-approved AO must notify CMS, whenever it terminates the accreditation of a Medicare deemed provider or supplier. The notice must be submitted by e-mail simultaneously to:

- the CMS AO oversight program electronic mailbox; and
- the applicable CMS RO electronic mailbox:

CMS provides the specific electronic mailbox addresses to the AOs.

The AO’s notice to CMS notice must provide the effective date of the termination of accreditation, as well as the reason for the termination. The RO forwards the termination notice electronically to the applicable SA.

- Accreditation termination is concurrent with switch to another CMS-approved Medicare accreditation program, or provider/supplier was previously also deemed by another AO whose accreditation remains in effect:

  - Unless there was an involuntary termination for failure to comply with the AO’s accreditation standards, if the provider/supplier’s termination by one AO is concurrent with a new recommendation for accredited, deemed status by another CMS-approved AO, or if the provider/supplier was previously deemed based on multiple accreditations, each by a different AO, then the provider/supplier remains deemed and under the jurisdiction of the other AO. The recommendation for deeming is sent by the AO to the CMS AO oversight program and the applicable CMS RO, which forwards the AO’s recommendation letter electronically to the applicable SA. An update packet including the new recommendation for deemed status by another AO must be submitted by the SA to the RO. The SA also updates the information in the deemed status tab of the provider’s/supplier’s certification information in ASPEN to reflect both the termination of the first AO’s accreditation and, where there was a switch to another AO, the accreditation by the second AO.

- If the termination was involuntary due to failure to comply with the AO’s accreditation standards and if the provider or supplier’s deemed status has not already been removed due to a prior enforcement action, the RO must consider this a substantial allegation of noncompliance with Medicare standards and must authorize the SA to conduct a complaint investigation survey. The SA surveys the provider/supplier within 45 days (or, if the RO’s reason for termination suggests an immediate jeopardy, according to the immediate jeopardy timeline for complaints) in order to provide assurance that the facility is in substantial
compliance with the applicable health and safety standards. If the SA’s survey finds no condition-level deficiencies, the provider/supplier retains deemed status under the other/new AO. If the SA finds condition-level deficiencies, then deemed status is removed in the same manner as for any other survey of a deemed provider/supplier.

- Accreditation termination is not concurrent with switch to another AO, or provider/supplier was not previously deemed by multiple AOs:

  If there is no concurrent recommendation of deemed status for the provider/supplier from another AO or if the provider/supplier was not previously deemed based on multiple accreditations, each by a different AO, the provider’s/supplier’s deemed status is removed and it is placed under SA jurisdiction. The SA updates the information in the deemed status tab of the provider’s/supplier’s certification information in ASPEN to reflect the termination of the AO’s accreditation and removal of deemed status. The SA surveys the provider or supplier in order to provide assurance that the facility is in substantial compliance with the applicable health and safety standards. Timing of the SA survey is as follows:

  - When the AO advises CMS that the provider’s/supplier’s accreditation was involuntarily terminated due to failure to comply with the AO’s accreditation standards, the SA must conduct the compliance survey within 45 days or, if the RO’s reason for termination suggests an immediate jeopardy, according to the immediate jeopardy timeline for complaints.

  - In all other cases the SA prioritizes the provider’s/supplier’s survey on the basis of the current CMS policy concerning survey frequencies and SA workload priorities, using the date of the most recent accreditation survey to calculate the survey interval, unless:

    - The facility is a home health agency (HHA), then the SA must conduct the survey no later than 3 years after the last accreditation survey; or

    - The RO exercises its discretion to request the SA to conduct the survey by a specified date.

Adverse accreditation action other than termination

When an AO takes an adverse action that is not termination against the accreditation status of a provider/supplier, the AO is required to inform both the CMS CO and the appropriate RO of the adverse action. As long as provider’s/supplier’s accreditation is not terminated, the provider’s/supplier's participation in Medicare is not affected. Generally the RO will not authorize a validation survey by the SA, but it has the discretion to do so in rare circumstances.
Note that none of the above scenarios concerning termination or other adverse
accreditation actions apply to the situation where a provider or supplier is acquired by a
new owner who rejects assignment of the prior Medicare agreement. In such a situation
CMS terminates the provider agreement of the seller as a voluntary cessation of business.
It does not matter whether the terminated provider or supplier was deemed, whether
under one or multiple CMS-approved Medicare accreditation programs, nor are any
further actions taken by CMS or the SA in response to a notification by an AO of an
accreditation termination or other adverse accreditation action related to the provider or
supplier covered by that prior Medicare agreement. (See Section 2003B.)
3300 - Confidentiality and Disclosure of Records - Citations and Applicability
(Rev. 1, 05-21-04)

Section 1106 of the Social Security Act (the Act) prohibits disclosure of any file, record, report, or other writing, or any information obtained at any time by or from the Secretary or an office or employee of DHHS in the course of discharging his duties under the Act, except as prescribed by regulations. The applicable regulations are found in 42 CFR 401, Subpart B (Confidentiality and Disclosure).

The regulations set out what records are available, how they may be obtained, and, where applicable, when a fee is paid to offset the cost of administrative activity involved in furnishing the information.

3302 - Federal Freedom of Information Act (FOIA)
(Rev. 1, 05-21-04)

Coexisting with the confidentiality provision of §1106 of the Act are the provisions of the 1967 “Freedom of Information” amendment to the Administrative Procedures Act. This amendment establishes the right of the public to access numerous types of Federal records and information. Exempted from mandatory disclosure under this amendment, however, are records and information that other Federal confidentiality statutes prohibit being disclosed. See 42 CFR 401.118 regarding the deletion of identifying details.

3304 - Multi-Program Information in SA Files
(Rev. 1, 05-21-04)

The CMS’ rules governing disclosure of Medicare/Medicaid/CLIA records and information to the public may be more or less restrictive than SA rules or those of other Federal programs. The SA should carefully distinguish between:

A. Records and information the SA acquires as an agent of a CMS program, and

B. Other records and information which:

1. The SA independently acquires through a State program; or

2. Are known to other parties who are not subject to a restriction or disclosure. The information known to these parties is considered as having entered the public domain.

Only the information the SA acquired in its role as an agent of CMS, and which has not otherwise entered the public domain, is subject to CMS’ disclosure rules.
When the SA obtains a record or an item of information that is not in the public domain and is held for joint use by the Medicare/Medicaid programs and other State or Federal programs, the SA applies the most restrictive confidentiality policies of all the programs to which the information relates.

Once any record or item of information has been forwarded to CMS, it is treated according to whatever CMS rule is applicable. Consequently, the SA is free to disclose the information listed in §3308 below, either on the basis that the State is an agent of the Medicare/Medicaid/CLIA program or on the basis that such information has entered the public domain through CMS.

3305 - Sharing State Licensure Information With Medicare Contractors
(Rev. 1, 05-21-04)

To promote the fiscal integrity of Federal health programs it is essential that all health care professional licensure revocations, non-renewals and denials as well as other changes in the legal ability to practice a profession or operate a health care business are reported to any appropriate Medicare fiscal intermediary, carrier or other Medicare contractor within two weeks after the State takes action as the result of such a change. Notice of a change in licensure status or the legal ability to practice should be reported in writing (hard copy or electronically) and should contain the reason for change in the licensure status, as well as all information necessary to adequately identify the health professionals or entities in question. This would include entities and individuals such as physicians who, although they are not surveyed for compliance with Conditions, nevertheless play an important role in the Medicare program by approving plans of care or treatment and ordering items and services for Medicare beneficiaries.

3308 - Information That May Be Disclosed to Public
(Rev. 1, 05-21-04)

(See Chapter 7 for disclosure requirements for SNFs and NFs)

3308A - Information Disclosable to Public Under CMS Rules That May Be Disclosed Directly by the SA
(Rev. 1, 05-21-04)

1. A facility does or does not participate in the Medicare/Medicaid/CLIA program;

2. The official Medicare/Medicaid/CLIA report of a survey, except to the extent that it contains:
   - The name of any patient;
   - Medical information about any identifiable patient;
   - The identity of a complainant;
• The address of anyone other than an owner of the facility; or

• Information which could be defamatory toward any identifiable person.

NOTE - The SA reviews the report of survey, and if it contains any of the above elements, it deletes the information from the report by blocking it out fully prior to release of the report. (See 42 CFR 401.118)

3. Citations of deficiencies that have been conveyed to the provider following a survey, except to the extent the report contains any of the identifiable information listed above. The SA blocks this information out prior to release of the statement of deficiencies;

4. PoC and pertinent comments submitted by the provider relating to Medicare/Medicaid/CLIA deficiencies cited following a survey, except to the extent the PoC or comments contain any of the identifiable information listed above. The SA blocks this information out prior to release of the PoC;

5. Official notices of involuntary provider termination;

6. Reports and information about a laboratory's performance in proficiency testing programs;

7. The CMS manuals distributed to the SAs, intermediaries, carriers, providers, or suppliers; and

8. Statistical data on provider characteristics that do not identify any specific provider or individual.

3310 - Requests for Information About Nonparticipating Institutions
(Rev. 1, 05-21-04)

Information disclosable about an institution, agency, organization, or supplier that has never been surveyed for participation in the Medicare/Medicaid/CLIA program is limited to the fact that it has never participated.

3312 - Charges for Information
(Rev. 1, 05-21-04)

If a member of the public requests from CMS copies of the records and information described in §3308 of this manual, there will generally be a charge. The SA does not provide free copies of more than a few pages of disclosable information if the cost of reproduction is to be charged to the Medicare/Medicaid/CLIA program. Charges should be in accordance with 42 CFR 401.136 and 401.140 for Medicare, or with any applicable State fee schedules for reproduction.
3314 - Time Periods for Disclosure Other Than Nursing Homes
(Rev. 1, 05-21-04)

(See Chapter 7 for Disclosure requirements for SNFs and NFs)

For Medicare, §1864(a) of the Act specifies that the Secretary shall disclose survey information within 90 days following the completion of each survey. Implementing regulations in 42 CFR 401.133(a) and (b) further provide that survey-related information prepared by the State and responded to by the facility must be disclosed within 90 days following completion of the survey by the State but may not exceed 30 days following CMS’ receipt of such information, or if prepared by CMS, within 30 days following the final preparation.

For Medicaid, §1902(a)(36) of the Act and 42 CFR 431.115 provide that survey-related information must be disclosed either upon determining that a provider is eligible to be certified or recertified, or within 90 days after completion of the survey, whichever occurs first. Any applicable State law(s) would also be considered.

Sections 1819(g)(5) and 1919(g)(5) of the Act provide disclosure requirements specific to SNFs and NFs within 14 calendar days after the information is made available to the facility. (Also see 42 CFR 488.325.)

For both Medicare and Medicaid, the date the survey is completed is the last day of the survey.

The provisions of FOIA require that when a request to disclose information is received, the information be released within 20 working days, or if this is not possible, the requester be notified within 20 working days when the information will be released. Follow this policy unless the State has other disclosure rules in which case those rules apply to the time allowed for disclosure. (See PL 104-321.)

When the SA receives a request for information, disclosure of which is not clearly permissible under the preceding sections, the SA declines to disclose the information requested on the basis of the provisions of §1106 of the Act and refers the request to the RO. The SA notifies the requester of this action within 20 working days and directs the requester to contact the RO for additional information concerning the denial. (See PL 104-321.)

3316 - Information Furnished to Original Source
(Rev. 1, 05-21-04)

Any party is entitled to information which that party supplied initially. Accordingly, one copy of the current survey report (subject to the restrictions in §§3308 and 3320) or other official form completed by or for the provider/supplier, correspondence from the
provider/supplier, and documentation submitted by the provider/supplier may be furnished to the original source without charge.

When a request is made by a provider/supplier to examine or secure copies of documents it has submitted, but the item requested has been sent to the RO, the SA acknowledges receipt of the request and refer the request to the RO, which will arrange for the original documents to be examined or copied, as appropriate.

**3318 - Disclosure of Information To and From Operating Components**  
(Rev. 1, 05-21-04)

According the 42 CFR 401.134, the SA may disclose confidential certification information about a provider/supplier without the provider/supplier’s authorization when such disclosure is necessary for proper performance of the duties of:

- An officer or employee of DHHS;
- An officer or employee of a SA or an intermediary participating in administration of title XVIII and/or title XIX by contract, agreement, or State plan for purposes of carrying out such contract, agreement, or State plan; and
- An officer or employee of a SA carrying out duties under State law in licensing or approving facilities.

Generally, confidential certification information is not to be released outside of the State survey and certification unit. Release by a SA to another State component or to a county or other local entity that performs survey functions for the SA is predicated on first obtaining agreement by the component or county to use the information only for certification or licensure purposes. The receiving SA component or other State or local entity may not release any certification information, subject to the penalty provisions of §1106 of the Act.

**3319 - Monthly Quality Indicator Comparison Reports Policy**  
(Rev. 1, 05-21-04)

**3319A - Purpose**  
(Rev. 1, 05-21-04)

The Monthly Quality Indicator Comparison Report was created by CMS in order to give state personnel and facilities a report that can be selectively released to certain requestors without violating the confidentiality of resident information. The Facility Quality Indicator Profile Reports, which provide data on the MDS-based quality indicators at a facility level have been available for some time. Because of issues of resident privacy, it has not been permissible to release these reports to requestors such as consumers. The Monthly Quality Indicator Comparison Report contains facility-level information while conforming to privacy rules; thus, States and facilities may release these reports.
3319B - Contents
(Rev. 1, 05-21-04)

The Monthly Quality Indicator Comparison Report is based on the same data used to produce the Facility Quality Indicator Profile Report, but the data are presented differently. (A sample of the report is attached to Survey and Certification Policy Letter 01-17.) The Monthly Quality Indicator Comparison Report contains both state percent and national percent comparison columns. This allows the reviewer to compare the performance on each quality indicator measure to that of the state and of the nation. The state percent column contains the “percent triggered” for the quality indicator averaged over all facilities within the state. This is an unweighted average, meaning that each facility has an equal impact on the average, regardless of number of residents. The national percent is also an unweighted average. It represents the “percent triggered” for the quality indicator averaged over all the facilities in the nation.

3319C - Differences From Facility Quality Indicator Profile Report
(Rev. 1, 05-21-04)

Because of concerns with protection of the privacy of residents, some of the data presented on the Facility Quality Indicator Profile Report are not included on the Monthly Quality Indicator Comparison Report. The numerator and denominator used to calculate the percents are not presented on the Monthly Quality Indicator Comparison Report. Additionally, totals are not displayed in instances when the number of residents in a facility is 10 or fewer. Also, if the percent triggered is 90 percent or greater for any given quality indicator, the exact percent triggered will not be displayed. Instead, a notation indicating that the percent triggered is “greater than or equal to 90 percent” will be displayed.

3319D - Delivery of Report
(Rev. 1, 05-21-04)

At the end of every month, the Monthly Quality Indicator Comparison Report is generated for each facility within a state, and is placed in that facility’s directory. The report does not allow an option for user-defined dates. Due to a built-in lag in the data system, the data that appear on the report generated in a given month are based on quality indicators from four months prior. The name of the report text file in the directory reflects this. The file name is MSRmmyyyy.txt, where mmyyyy is the month and year of the data from which the report was created. This date also appears at the top of the report itself.

3319E - Release of Report
(Rev. 1, 05-21-04)

The Monthly Quality Indicator Comparison Reports were designed for use by:

1. Family members of a facility resident;
2. Potential facility residents or their family members; and

3. The State ombudsman.

Release of the reports for a large number of facilities or to individuals other than those named above is at the State’s discretion. Although CMS has suppressed certain data in order to protect individual privacy, the state and the facilities must be cautious in evaluating requests for these reports and in releasing them:

- The reports were designed to enable individuals with a personal or professional interest in a facility, or a few facilities to have access to quality indicator information at a facility level. A State ombudsman, discharge planner, or other appropriate person could help the consumer understand and utilize the information in a small number of reports.

- Since release of these reports was not designed for large batches, the data system does not include a direct method to print reports for a large number of facilities at once.

- Explanatory language appropriate for consumers is not included with the reports; therefore the reports are not designed to be published or to be posted on the Internet.

- State or facility staff should remind recipients of the reports that the information contained in the report is not a measure of a facility’s quality, but is rather an indicator of the potential for quality problems. Recipients should not use these reports alone to draw conclusions about a facility’s quality. They should also visit the facilities that they are evaluating.

- Each recipient of a Monthly Quality Indicator Comparison Report should also receive a copy of the sheet “The Monthly Quality Indicator Comparison Report: Guidelines” (see F below). The sheet will aid recipients in the appropriate use of the report.

3319F - Monthly Quality Indicator Comparison Report Guidelines
(Rev. 1, 05-21-04)

The Monthly Quality Indicator Comparison Report was created by the CMS to give individuals with either a personal or a professional interest in a specific facility access to facility-level data on quality indicators without violating the confidentiality of the nursing home residents. The report allows comparisons between a specific nursing home and the state, as well as the nation about the proportion of residents with a particular condition.

- The data in these reports are only a starting point in reaching a decision about the quality of a nursing facility. The data should be viewed as measures of the
potential for quality problems. There may be a legitimate reason why a facility has higher than average percentages on a particular indicator.

- In making decisions about quality of care, it is important to actually visit the nursing home being considered. Information in this report can help decide what type of questions an individual may want to ask on a visit to the nursing home.

- Additional information about individual facilities can be found in the Nursing Home Compare section of the CMS Web site http://www.medicare.gov/. This includes the State inspection results, found in the “About the Nursing Home Inspections Results” section of this web site.

3320 - Necessary Preclearance With RO Before Releasing Confidential Information
(Rev. 1, 05-21-04)

The SA obtains advice from the RO when a request is received for confidential Medicare and/or Medicaid information under circumstances other than as permitted by §3308. To the extent known, indicate the purpose to be served by the release, the specific information to be released, and, if appropriate, the availability of the information elsewhere.
Additional State Agency Responsibilities

3330 - HHA Toll-Free Hotline and Investigative Unit
(Rev. 1, 05-21-04)

Section 1864(a) of the Act requires that the State establish a toll-free hotline to collect, maintain, and continually update information on Medicare participating HHAs including certification-related deficiencies found regarding patient care, corrective actions, and sanction activity during its most recent survey. Complaints and questions will also be received over the hotline concerning HHAs in the State. The State is to inform Medicare beneficiaries of the availability of the hotline, its purpose, the hotline telephone number, and hours during which it is in service. The SA (under the §1864 agreement) may subcontract with the appropriate State or local agency for operation of the toll-free hotline. However, the SA is required to investigate certification-related complaints received by the hotline.

The instructions are minimum requirements for establishing and maintaining the hotline. Additional operating procedures may be used to complement the requirements. This activity will be monitored by the RO and enforced through the §1864 agreement process.

3330A - HHA Hotline Function
(Rev. 1, 05-21-04)

• Operate the toll-free hotline 6 hours per day (between 7:00 a.m. - 6:00 p.m.) Monday - Friday, except on State or Federal holidays. Notify the public in advance if there are changes in hours of operation;

• Provide HHA hotline information to HHAs and the public by telephone or in writing;

• Record most recent certification and enforcement findings regarding patient care within 30 calendar days of the determination of certification or recertification, approved PoC, correction of deficiencies, and any sanctions imposed, including termination of a Medicare participating HHA;

• Record complaints received on the HHA hotline;

• Refer HHA hotline certification-related complaints to the survey unit for possible investigation; and

• Refer non-certification related complaints to the appropriate components in the State agencies.
3330B - HHA Hotline Information
(Rev. 1, 05-21-04)

These items must be maintained and readily retrievable:

- Name, address, and Medicare provider number of Medicare HHAs in the State;
- Date of most recent Medicare certification or recertification survey of individual HHAs;
- Record of any Condition level deficiencies found regarding patient care in the most recent survey conducted by the SA;
- Date(s) of planned corrective action(s) and completed corrective action(s) for Condition level deficiencies; and
- Date(s) and type of sanction(s), if any imposed, including termination.

3330C - Disclosure of Information
(Rev. 1, 05-21-04)

When responding to calls requesting information, the SA releases the information verbally and as necessary in writing in accordance with CMS public disclosure rules. (See §§3300-3320.)

3330D - Record Keeping Requirements
(Rev. 1, 05-21-04)

The SA records each call as either a complaint or a general inquiry. The record keeping system may be manual or automated and must indicate the general nature of the inquiry or complaint and the resolution, i.e., question answered, material sent, or complaint referred for investigation.

3330E - Public Awareness
(Rev. 1, 05-21-04)

The State is to notify Medicare beneficiaries and HHA’s of the availability of the hotline. In addition, the State is to notify each Medicare HHA’s about the HHA hotline, the telephone number, purpose, and hours of operation.

3330F - Hotline Investigative Unit
(Rev. 1, 05-21-04)

The SA investigates the certification-related complaints received by the hotline following procedures in SOM Chapter 5. In instances where a complaint is not certification-related,
the SA refers that complaint to the proper State agency (e.g., the State medical society, the State licensure agency, or the State social services department), or the RO.

**Response to Subpoenas Served On and Suits Against the State Agency**

**3350 - Subpoena for Program Records**
(Rev. 1, 05-21-04)

When one of the officers or employees of an SA is served with a subpoena or other legal proceeding to produce title XVIII and/or title XIX records or CLIA records, whether or not the information is partly or wholly disclosable, accept the subpoena. The SA should immediately notify its legal advisor or counsel and the RO of receipt of the subpoena, and provide copies of the subpoena and other pertinent documents to both. The SA determines whether the subpoenaed records and information are routinely disclosable to the public or contain confidential information that is normally withheld. The SA includes this information with the notification.

The SA should place the subpoenaed records and information in a secure area to prevent unauthorized disclosure and assure the availability to counsel for review.

If not included in the subpoena but known to the SA, it informs its legal advisor and the RO of the names, addresses, and telephone numbers of the presiding judge and attorneys, and the purpose of the subpoena.

After the RO receives the materials, it coordinates with the SA legal advisor and takes other actions necessary to assist the SA. After consultation with the SA, the RO and the SA legal advisor determine whether the SA must produce the subpoenaed records and information and if so, notify the SA director to comply. If it is determined that all or part of the subpoenaed records and information are to be withheld, the RO and/or SA legal advisor represents the SA in dealing with the court of jurisdiction and enter the motions to quash the subpoena, based on the provisions of §1106 of the Act which prohibits disclosure of confidential records and information and other pertinent statutes and regulations.

**3352 - Forthwith Subpoena**
(Rev. 1, 05-21-04)

On rare occasions the SA may be served with a Forthwith Subpoena. Unlike a standard subpoena, which requires that records and information be produced on a specified date and delivered to the court, a Forthwith Subpoena requires that the SA immediately provide the subpoenaed records and information. The SA accepts the subpoena and requests the individual to wait.

The SA immediately notifies its SA legal advisor and requests that the legal advisor call the RO.
The RO and the SA legal advisor will determine by telephone the appropriate action to be taken. The SA legal advisor will deal with the individual waiting for the records and information. The SA should not take unilateral action of any type.

3354 - Subpoena for SA Licensure Records
(Rev. 1, 05-21-04)

If the SA has integrated certification and licensure files and it receives a subpoena requiring it to produce licensure information, it should consult with the RO before complying with the subpoena to determine if the integrated file contains information prohibited from disclosure by §1106 of the Act. (See §3300.) If the files contain such information, the RO assists the SA legal advisor in determining the appropriate procedures and action to be taken in responding to the subpoena.

3356 - Suit Against SA
(Rev. 1, 05-21-04)

Where a suit is brought against the SA as a result of its title XVIII and/or title XIX or CLIA functions, the SA forwards the notice of the suit to the RO. The RO coordinates with the SA and its legal advisor.
### Transmittals Issued for this Chapter

<table>
<thead>
<tr>
<th>Rev #</th>
<th>Issue Date</th>
<th>Subject</th>
<th>Impl Date</th>
<th>CR#</th>
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</thead>
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<tr>
<td>R198SOMA</td>
<td>01/17/2020</td>
<td>Revisions to the State Operations Manual (SOM) Chapter 2 and Chapter 3</td>
<td>01/17/2020</td>
<td>N/A</td>
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<td>R190SOMA</td>
<td>06/14/2019</td>
<td>Updates to the State Operations Manual (SOM) Chapters 2, 3 and 9 to add instructions for Organ Transplant Programs.</td>
<td>06/14/2019</td>
<td>N/A</td>
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<td>R188SOMA</td>
<td>04/26/2019</td>
<td>Revisions to the State Operations Manual (SOM 100-07) Chapter 2, The Certification Process, Chapter 3, Additional Program Activities, and Chapter 4, Program Administration and Fiscal Management</td>
<td>04/26/2019</td>
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<td>123SOM</td>
<td>10/03/2014</td>
<td>Revisions to State Operations Manual (SOM) Chapters 1, 2 and 3</td>
<td>10/03/2014</td>
<td>N/A</td>
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<td>R121SOM</td>
<td>09/19/2014</td>
<td>Update to State Operations Manual (SOM), Publication 100-07, Chapter 3, to Provide Language-Only Changes for Updating ICD-10</td>
<td>Upon Implementation of ICD-10</td>
<td>N/A</td>
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<td>R112SOM</td>
<td>04/11/2014</td>
<td>State Operations Manual (SOM) Chapter 3 Policy Revisions For Organ Procurement Organizations (OPOs)</td>
<td>04/11/2014</td>
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<td>R92SOM</td>
<td>11/20/2013</td>
<td>State Operations Manual (SOM) Chapter 3 Policy and Nomenclature revisions for Intermediate Care Facilities for Individuals with Intellectual Disabilities (ICF/IID)</td>
<td>11/22/2013</td>
<td>N/A</td>
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<td>R24SOM</td>
<td>01/26/2007</td>
<td>Sunset of the Policies for Provider Nominations for an Intermediary and the Provider Requests for a Change of Intermediary</td>
<td>01/26/2007</td>
<td>N/A</td>
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<td>R01SOM</td>
<td>05/21/2004</td>
<td>Initial Issuance of Pub 100-07</td>
<td>N/A</td>
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