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Message from URAC

Dear interested party or applicant:

Quality-based operations should be the centerpiece of any company doing business in today’s health care system. Quality improvement activities promote a wide range of benefits such as increasing operational efficiencies, reducing business risks and improving patient health outcomes. However, health care professionals must identify and implement a quality improvement methodology that really works for their particular business model and health care setting.

Through its modular approach to accreditation, URAC works with the industry and other key stakeholders to benchmark URAC standards against key organizational structures and business functions. Now in its 18th year of operation, URAC offers 20 different accreditation and certification programs. URAC has issued more than 10,000 accreditation certificates to companies operating in all 50 states and internationally. URAC also is recognized as part of the regulatory process in three-fourths of the states and by four federal agencies.

URAC, as a nonprofit, independent accreditation agency, brings to the table a nationally-recognized accreditation process and seal of approval. URAC’s success is tied in large part to the broad-based, consensus-driven process by which hundreds of volunteers help URAC draft and update its standards and also oversee the accreditation system. These volunteers represent the interests of a wide variety of stakeholders including purchasers, regulators, consumers, providers and industry representatives.

All companies that apply for URAC accreditation make improvements to their operations as a result of the review process. The desktop review of the application identifies issues early on in the process and helps focus the onsite review, which is designed to confirm compliance with the standards. During the onsite visit, accreditation reviewers exchange information with applicants in what often becomes a mutual learning experience. URAC’s goal is to identify and promote best practices for each market segment that it accredits.

Receiving the accreditation certificate signifies a job well done and distinguishes the organization as having met a standard of excellence. As a result, URAC accredited organizations join the ranks of a select community who have documented and verified their commitment to quality health care.

Please contact URAC if you would like to find out more about the accreditation process or to become involved with one of our committees, educational programs, research initiatives, or other projects. We look forward to hearing from you.
Introduction to URAC Accreditation Guides and Standards

URAC offers two references addressing standards. The Accreditation Standards provides a copy of the standards produced by URAC and defined terms, which are italicized within the standards. It is a resource for government agencies and private entities wanting to examine the standards for their own purposes. For organizations contemplating accreditation, URAC’s Accreditation Guide provides, in addition to the standards, information about the documentation to submit as evidence for meeting the intent of the standards as well as the types of materials and activities URAC’s accreditation reviewers will be examining during an onsite visit. Both the Accreditation Standards and Guide are available through URAC’s Business Development Department at (202)216-9010 or send an e-mail to BusinessDevelopment@urac.org. For detailed information about how to prepare an application for accreditation, please go to: http://www.urac.org/healthcare/accreditation/accreditnet.aspx for a copy of the AccreditNet Instruction Booklet, designed to complement the Accreditation Guide for applicant organizations.

The Accreditation Standards and Accreditation Guide are intended to provide guidance only. The URAC Accreditation Committee and Executive Committee hold the final authority to make determinations regarding interpretation and application of standards, and an applicant’s compliance with standards.

The Accreditation Guide is provided to assist applicants understand the meaning or intent of the standards. That being said, it cannot cover all possible situations and subsequent interpretations that may apply. Therefore, applicants should be aware that the standards are subject to ongoing interpretation and as such, changes can be made to the Accreditation Guide.

Each company applying for accreditation should carefully review URAC’s accreditation standards and the defined terms italicized within the standards, then use the Accreditation Guide and AccreditNet Instruction Booklet to prepare an application for submittal to URAC.

Modular Concept

URAC uses a “modular accreditation system” that is adaptable to the continuing evolution of the health care system. A module is a set of standards established for a particular health care function. The collection of standards contained within modules are unique to that health care service or function. The Core Standards incorporate the basic elements necessary to promote quality for any type of health care organization and were designed for two purposes: 1) to act as a “foundation” for function-specific
accreditation programs, and 2) to act as a “stand-alone” accreditation program for companies not delivering services under one of the specific functions or modules.

Each accreditation will include Core and the module(s) covering the functions.

\[
\text{Core Standards + Module(s) = Specific Accreditation}
\]

Core “stand-alone” accreditation will only consist of Core Standards.

\[
\text{Core Standards Only = Core “Stand-Alone” Accreditation}
\]

Eligibility to apply for Core “stand-alone” accreditation is determined on the basis of how a company markets itself. If a health care organization markets any of the services addressed under one of URAC’s modules, Core “stand-alone” accreditation is not an option. An example of an organization that would be eligible for Core “stand-alone” accreditation is an organization that provides health care educational services.

For applicants, the modular system provides the flexibility to choose from a variety of accreditation programs. For example, an applicant may choose to apply for Utilization Management (UM) accreditation initially, and when up for reaccreditation, add the Case Management (CM) module.

\[
\text{Core with Single-Module Application (Example: Core & UM only)}
\]

\[
\text{Core with Multi-Module Application (Example: Core & UM + CM)}
\]

With several choices available, an applicant can tailor the accreditation to its current needs and business goals. If you are not sure what modules would best fit your organization, URAC’s Business Development Department can be reached at BusinessDevelopment@urac.org or at (202) 216-9010 to answer questions, provide pricing information and help organizations decide the best course of action.

**Compliance with State and Federal Law**

The Accreditation Guide provides information on URAC’s expectations regarding compliance with each standard. Some standards require applicants to attest to compliance with specific state regulations regarding operational policy and procedure. Prior to submitting an application the applicant should conduct a review of its legal obligations, including those addressed in the standards. Although it is not indicated for each standard, URAC expects that the applicant will be in compliance with all applicable state and federal laws that pertain to relevant operations. State and federal laws supersede URAC Standards if
the laws or regulations are more rigorous than URAC Standards. Conversely, an applicant must comply with URAC Standards if the standards are more stringent. If an applicant is required by law to carry out its business in a manner not consistent with URAC Standards, then the applicant may request a variance from a URAC Standard. A copy of the relevant statute or regulation must accompany the request submitted for that standard in the application.

**Standards and Interpretation**

The standards are grouped together into modules, with each module representing various health care functions. Individually, the standards address the structures and processes that need to be in place for performing the function to be accredited according to national standards. For the most part, an applicant is expected to be in compliance with all applicable standards at the time of application for accreditation.

In the **Accreditation Standards**, you will find:

- **Definitions.** All italicized terms found in the standards are defined in this section.
- **Standards with assigned Weights.** Standard elements include assigned weights for scoring. If an element in and of itself does not contain enough information to evaluate compliance without the following sub-element, then it is noted as "Not Weighted."

In the **Accreditation Guide**, you will find:

- **Definitions.** All italicized terms found in the standards are defined in this section.
- **Standards with assigned Weights and Interpretive Information for each Standard.**
- **Points to Remember and Scope of Standards.** These bullet points identify important issues to consider when documenting your organization’s compliance with the standard. In some cases, additional details are provided that will help your compliance efforts and in other cases, these details will alert you to potential pitfalls.
- **Evidence for Meeting the Standards: Desktop Materials and Onsite Review Materials and Activities.**
- **Bright Ideas.** This section is not used for every standard and contains common industry practices that may be helpful to the applicant organization. (Note: adoption of a “bright idea” is not required for compliance with a standard, nor does adoption of the “bright idea” guarantee compliance with that standard.)
- **Related Standards.** This section is not used for every standard, but helps to identify relationships between standards that are not always obvious and helpful to know about.
Scoring Methodology

Policy Regarding “Not Applicable” Elements and Standards

If a weighted standard element or an entire standard is determined to be *not applicable*, then they are not included in the scoring calculations (i.e., deducted from the denominator). This includes the rare instance when a “variance” is granted by the requisite URAC committee. As a result, applicants are not penalized when a standard element or standard is not applicable.

If a mandatory standard element is determined to be *not applicable*, then it does not count against the applicant when determining an accreditation category; however, applicant organizations must have a policy that meets the intent of a mandatory standard element even if it is not currently being implemented.

An applicant may choose not to meet any or all leading indicator standard elements (i.e., leading indicators are *optional*); therefore, a leading indicator standard element, which is not weighted, cannot be made *not applicable*. Not all accreditation standard sets have leading indicators.

Standard Element Weights

URAC’s Scoring System has six (6) distinct categories of standard elements:

- **Weight = 1**: Emerging Practice
- **Weight = 2**: Basic Infrastructure
- **Weight = 3**: Promotes Quality
- **Weight = 4**: Key Stakeholder Right / Empowers Consumers
- **Mandatory = M**: Non-weighted, mandatory element with a direct or significant impact on consumer safety and welfare
  - All mandatory elements must be met at 100% compliance in order to achieve a Full accreditation
  - If determined to be not applicable, applicant must have a policy and procedure in place that meets the intent of the element should the organization need to implement it in the future
Leading Indicator = L: Non-weighted, optional element highlighting effective practices not yet widely adopted in health care

- Potential forecast of where the health care industry may be heading
- Provides a way for an organization to distinguish itself from other accredited companies
- Leading indicators are not reported to URAC's Accreditation or Executive Committees and do not influence an applicant's final accreditation score or category
- Cannot be designated “not applicable” given that they are optional
- Before URAC will acknowledge that an applicant has met a leading indicator:
  - Full accreditation must be achieved, and
  - Element must be met at 100% compliance
  - Initially URAC will list leading indicators in the Accreditation Summary Report (ASR)
- Other types of marketing exposure may be considered in the future (e.g., Website, conferences, etc.)

Definitions for the standard element categories are listed on the following pages. As you analyze the standard elements to assign a weight, keep in mind the following:

- Standards are no longer weighted, but standard elements are. Elements are the components of a standard that are evaluated through the accreditation review process.
- Standard elements are no longer designated as “primary” or “secondary.”

Computing an Accreditation Score

- Scoring an Element
  - Element weight x Compliance
- Scoring a Standard
Total points achieved ÷ Total points possible

Scoring a Module

Total points achieved ÷ Total number of standards

Scoring a multi-Module accreditation

(Core score x .30) + (Module score x .70)

Scoring a multi-Site accreditation

Lowest site score determines the application score

Determining an Accreditation Category

- If one Mandatory standard element is not met Conditional
- If two Mandatory standard elements are not met Corrective Action
- If three Mandatory standard elements are not met Denial
- If all Mandatory standard elements are met:

≥ 94 points/100 and complies 100% on at least one “Leading Indicator” standard à Include compliance with Leading Indicator(s) on the Accreditation Summary Report (ASR)

≥ 94 points/100 Full Accreditation

≥ 90, but < 94 points/100 Conditional Accreditation

≥ 85, but < 90 points/100 Corrective Action

< 85 points/100 Denial

Rating Compliance with a Standard Element

Standard elements are individually rated at 100% (full compliance), 50% (partial compliance) or 0% (no compliance) as follows:

For elements that require a file/record audit, the audit must reveal:
% Compliance with Element

100% Compliance (Full Compliance) = Audit score ≥ 80%;

Note: A mandatory element must be met at 100% compliance with an audit score of ≥ 90%; if not, then the mandatory element is considered not met. Credentials verification must be met at 100%. Applicants must meet all applicable mandatory elements in order to achieve Full accreditation.

Note: A leading indicator element must be met at 100% compliance; if not, then the leading indicator is considered not met. Applicants do not have to meet leading indicators since these types of elements are optional, acting as “extra credit.”

Note: A minimum of 30 files will be pulled for file review, but if the applicant does not have 30 files, then all files will be reviewed. For monitoring reviews, files will be pulled from the first 12 months of the accreditation cycle; for reaccreditation reviews, files will be pulled from the prior 24 months.

- 50% Compliance = Audit score ≥ 65%, but < 80%, or for contracts the audit score is < 80%, but the applicant has an internally approved, compliant contract template.
- 0% Compliance = Audit score < 65%, or for contracts the applicant does not have an internally approved, standards-compliant contract in place.

For elements that do not require a file/record audit:

% Compliance with Element

100% Compliance (Full Compliance) = Element documented pursuant to the standard element and upon verification is found to be fully implemented.

Note: A mandatory element must be met at 100% compliance; if not, then mandatory element is considered not met. Applicants must meet all applicable mandatory elements in order to achieve Full accreditation.
Note: A leading indicator element must be met at 100% compliance; if not, then the leading indicator is considered not met. Applicants do not have to meet leading indicators since these types of elements are optional, acting as “extra credit.”

- A standard element is implemented, where at least one of the following onsite activities is verified:
  - Staff is observed conducting the procedure correctly; or
  - Staff verbalizes the procedure correctly; or
  - Documented examples of implementation are surveyed; or
  - Documentation of oversight is reviewed; or
  - Management attests to its implementation and provides supporting documentation (i.e., sign-in sheets showing staff training session occurred, CV of newly hired medical director, sample of revised and distributed documentation such as a provider directory, notification letters, etc.)

- 50% Compliance (Partial Compliance) = Element documented pursuant to the standard element, but not consistently or completely implemented.

- One or more incidences of non-compliance in implementation will lower the compliance rating to 50%. This would include:
  - Errors implementing work processes during on-site observation by the URAC Reviewer.
  - Mistakes during interviews. If staff catches the error – misspoke – and corrects it, then this will not count as evidence of non-compliance.
  - Reports with data or analysis demonstrating non-compliance (e.g., not meeting timelines, wrong staff conducted the procedure, provider listed in a directory prior to credentialing, etc.)
  - Meeting minutes revealing decisions contrary to meeting the intent of the standards or lacking documentation indicating that a key activity did not take place (e.g., vote to eliminate provider appeal mechanism, minutes do not reflect review and update of the quality management program, etc.)
0% Compliance (No Compliance) = No evidence or incomplete evidence of compliance with the standard element in documentation or, regardless of documentation, applicant has not implemented the structures or processes needed to comply with the standard element. No compliance is exemplified when any one of the following statements is true.

The standard element is:

- Not addressed in documentation,
- Only partially addressed in documentation,
- Addressed in documentation, but does not meet the intent of the standard element,
- Not implemented, which does not include situations where:
  - The organization did not have the opportunity to implement. An example of this would be where an organization has an appeal process in place, but is either not contracted to do appeals or simply has not had an appeal of the type addressed by the standard.
  - The organization has been in business < six (6) months and is therefore eligible for a provisional accreditation.
- Implemented in a non-compliant manner, or
- Implemented, but one or more staff shows a pattern (≥ 4 occurrences) of non-compliance with an element over a period of time (within six (6) consecutive months), regardless of any warnings, corrective action taken (including training or procedural changes), or relative improvement over time.

**Scoring a Weighted Standard Element**

To score a weighted standard element ("element").

- Multiply the element weight by the compliance factor achieved (e.g., 0 for no compliance, .50 for partial compliance or 1.0 for full compliance).

  Calculation for Standard Element Score

(Compliance factor) × (Weight of standard element) = Score for an Element
Scoring a Standard

To score a standard with weighted elements,

- Sum the score achieved for each element.
- Divide by the number of points possible (sum of the element weights) for the entire standard.

- Weights for the elements determined to be not applicable are not included in the denominator and as such do not count against the applicant.

- Multiply by 100; this provides a percentage score for the standard.

Calculation for Standard Score

\[
\frac{\text{(Sum of all applicable element scores)}}{\text{(Total possible points for the standard)}} \times 100 = \text{Standard Score}
\]

Calculating a Final Total Accreditation Score for Core-only Applications and Accreditations that do not include Core

To calculate a final score for Core-only and non-Core Module Applications,

- Sum the scores for each standard.
- Divide by the total number of applicable standards.

- Standards determined to be not applicable are not included in the denominator and as such do not count against the applicant.

Calculation for Core and non-Core Module Score

\[
\frac{\text{(Sum of all applicable standard scores)}}{\text{(Total number of applicable standards)}} = \text{Module Score} = \text{Final Total Score (round to nearest tenth)}
\]

Note: For purposes of calculating an accreditation score, the number of standards possible is the total count of standards that have at least one weighted element; standards that do not have any weighted elements (i.e., only mandatory and/or leading indicator elements) are not included in the count since they do not contribute a standard score towards the module score. It is not logical to include standards in the denominator when it has no points to contribute to the score and to include such a standard in the denominator would unfairly penalize the applicant.
Note: If all weighted elements in a standard are determined to be not applicable, then the standard does not count towards the total number of applicable standards for purposes of scoring a module. To count them would unfairly penalize the applicant and is contrary to the URAC policy on standards determined to be not applicable.

Calculating a Final Total Accreditation Score for Multi-Module Accreditations

To calculate a final total accreditation score for multi-module accreditations, which includes:

· Core + a single Module (e.g., Health UM, IRO, CES, etc.)
· Core + multiple Modules (e.g., Health Plan, Health Network, etc.)

For Core:

- Sum all of the Core standard scores.
- Divide by the total number of applicable standards.

- Standards determined to be not applicable are removed from the denominator and as such do not count against the applicant.

- Multiply the Core score by .30 since Core is 30% of the final score for multi-module accreditations that include Core.

For non-Core module score:

- Sum all of the standard scores from all of the non-Core modules.
- Divide by the total number of applicable standards.

- Standards determined to be not applicable are removed from the denominator and as such do not count against the applicant.

- Multiply the non-Core score by .70 since the non-Core modules collectively contribute to 70% of the final score.
- Sum the percentage score for Core and the modules; a perfect score would be 100%.

Final Total Score
(Core score × .30) + (non-Core score × .70) = Final Total Score for One or Multiple Sites (round to nearest tenth)

**Final Total Score if all Sites achieve a Full (≥ 94)**

\[
\text{Final Full Accreditation Score} = \frac{\text{Sum of all Full onsite scores}}{\text{Number of sites that had an onsite review}}
\]

(round to nearest tenth)

**Note:**

- Each site that has an onsite review receives its own score; however, the lowest of these scores determines the score for the overall application.
- If all sites receiving an onsite review achieve a Full, the average score for these sites is the final score for the application. (See calculation above.)
- If there is a trend of three (3) or more sites that achieve less than a Full accreditation, then the URAC Reviewer has the discretion to visit all of the sites in the application.

**Determining How Many Mandatory Elements are Met and Not Met**

Mandatory elements are non-weighted elements and all applicable mandatory elements must be met at 100% compliance in order for an applicant to achieve Full accreditation.

- Mandatory elements that are determined to be not applicable are subtracted from the total count of mandatory elements and do not count against an applicant.
- Mandatory elements with a compliance level less than 100% (e.g., partial compliance [.5] and no compliance [0]) are considered *not met* and count against an applicant when determining an accreditation level. (See “Determining an Accreditation Level” below.)
Definitions for Health Accreditation Standards

(NOTE: Defined terms appear in italics throughout the standards).

Being familiar with these definitions is critically important to accurate understanding of URAC Standards. In the Standards, defined terms are italicized. Readers are encouraged to refer to the definitions section each time they encounter an italicized term until they feel they have committed the meaning of that term to memory.

Abandonment Rate: The percentage of calls offered into a communications network or telephone system -- i.e., automatic call distribution (ACD) system of a call center -- that are terminated by the persons originating the call before answer by a staff person.

Access: The consumer’s or client’s ability to obtain services in a timely manner.

Interpretive Note: The measures of access for consumers are determined by components such as the availability of services, their acceptability to the consumer, consumer wait time, and the hours of operation.

The measures of access for clients are determined by components such as turn-around time and other metrics as they may be defined in written business agreements, etc.

Adverse Benefit Determination: A decision by the Organization to deny or reduce a benefit for some or all of the lines of a claim (other than the application of a deductible or other cost-sharing).

Adverse Event: An occurrence that is inconsistent with or contrary to the expected outcomes of the Organization’s functions.

Advisory Board of Osteopathic Specialists (ABOS): American Osteopathic Association (AOA) certification agent organized in 1939 for the purpose of establishing and maintaining standards of
osteopathic specialization and pattern of training.

**American Board of Medical Specialties (ABMS):** Organized originally in 1933 as the Advisory Board of Medical Specialties, the ABMS (1970), in collaboration with the American Medical Association (AMA), is the recognized certifying agent for establishing and maintaining standards of medical specialization and pattern of training.

**Appeal:** A written or verbal request by a consumer, ordering provider or prescriber to contest an organizational determination (e.g., services have been denied, reduced, etc.).

**Interpretive Note for term “Appeal”:** Specific terms used to describe appeals vary, and are often determined by law or regulation.

**Appeals Consideration:** Clinical review conducted by appropriate clinical peers, who were not involved in peer clinical review, when a decision not to certify a requested admission, procedure, or service has been appealed. Sometimes referred to as “third level review.”

**Assessment:** A process for evaluating individual consumers that have been identified as eligible for the disease management program, to identify specific needs relating to their clinical condition and associated co-morbidities.

**Attending Physician:** The doctor of medicine or doctor of osteopathic medicine with primary responsibility for the care provided to a patient in a hospital or other health care facility.

**Attending Provider:** The physician or other health care practitioner with primary responsibility for the care provided to a consumer.
**Average Speed of Answer:** The average delay in seconds that inbound telephone calls encounter waiting in the telephone queue of a call center before answer by a staff person.

**Benefit Calculation:** An adjustment or calculation by the Organization of the financial reimbursement for a claim under the terms of the applicable benefit plan, provisions, criteria, provider contracts, or state rules.

**Benefits Program:** An arrangement to pay for health care services provided to a consumer. “Benefits program” includes, but is not limited to, health and medical benefits provided through the following organization types:

- Health maintenance organizations (HMOs);
- Preferred provider organizations (PPOs);
- Indemnity health insurance programs;
- Self-insured plans;
- Public programs, such as Medicare and Medicaid; and
- Workers’ compensation insurance programs.

**Blockage Rate:** The percentage of incoming telephone calls “blocked” or not completed because switching or transmission capacity is not available as compared to the total number of calls encountered. Blocked calls usually occur during peak call volume periods and result in callers receiving a busy signal.

**Board-certified:** A certification – approved by the American Board of Medical Specialties, the American Osteopathic Association, or another organization as accepted by URAC – that a physician has expertise in a particular specialty or field. To the extent that future URAC standards include other certifications, URAC will specify further approved boards.

**Interpretive Note for term “Board-certified”:** URAC recognizes that ABMS- and AOA-approved board certifications may not be the only certification programs that may be acceptable for health professionals in URAC-certified organizations. For example, non-physician professionals will have appropriate certifications that are not ABMS- of AOA-approved. Any applicant wishing to have URAC recognize another board certification program should notify URAC early in the certification process. URAC will then
take this recommendation to URAC’s Accreditation Committee.

The Accreditation Committee will review all requests, and will decide to approve or reject the certification. The Accreditation Committee will consider the following criteria in judging whether a certification is acceptable:

• Is the certification accepted within its target community of health professionals?
• Was the certification developed through an open, collaborative process?
• Does the certification reflect accepted standards of practice?
• Is the certification administered through an objective process open to all qualified individuals?

**Caller:** The consumer inquiring to obtain health care information. This may also be a representative inquiring on behalf of the consumer.

**Case:** A specific request for medical or clinical services referred to an organization for a determination regarding the medical necessity and medical appropriateness of a health care service or whether a medical service is experimental/investigational or not. It is a non-approval regarding medical necessity and medical appropriateness decisions for services covered under a health benefit plan’s terms and conditions or for coverage decisions regarding experimental or investigational therapies that is at issue during the independent review process.

**Case Involving Urgent Care:** Any request for a utilization management determination with respect to which the application of the time periods for making non-urgent care determinations a) could seriously jeopardize the life or health of the consumer or the ability of the consumer to regain maximum function, or b) in the opinion of a physician with knowledge of the consumer’s medical condition, would subject the consumer to severe pain that cannot be adequately managed without the care or treatment that is the subject of the case. (Note: This definition is derived from the Department of Labor’s definition of “claim involving urgent care.”)

**Interpretive Note for term “Case Involving Urgent Care”:**

While the URAC standards are silent on the methods by which a claim is determined to be a “case involving urgent care,” the Department of Labor claims regulation (29 C.F.R. § 2560.503-1(m)(1)) specifies that whether a claim is a “claim involving urgent care” is to be determined by an individual acting on behalf of the health benefits plan applying the judgment of a prudent layperson who possesses an average knowledge of health and medicine. Any claim that a physician with knowledge of the claimant’s medical
condition determines is a “claim involving urgent care” shall be treated as a “claim involving urgent care.

**Case Management:** A collaborative process which assesses, plans, implements, coordinates, monitors, and evaluates options and services to meet an individual’s health needs using communication and available resources to promote quality cost-effective outcomes.

**Certification:**

1) **(UM Specific Definition)** A determination by an organization that an admission, extension of stay, or other health care service has been reviewed and, based on the information provided, meets the clinical requirements for medical necessity, appropriateness, level of care, or effectiveness under the auspices of the applicable health benefit plan.

Interpretive Note for term “Certification”: “Determination” may vary depending on context.

2) **(General Definition)** A professional credential, granted by a national organization, signifying that an individual has met the qualifications established by that organization. To qualify under these standards, the certification program must:

   - Establish standards through a recognized, validated program;
   - Be research-based; and
   - Be based (at least partially) on passing an examination

**Claim:** Any bill, claim, or proof of loss made by or on behalf of a consumer or health care provider to an Organization (or its intermediary, administrator, or representative) for which the consumer or health care provider has a contract for payment of health care services. (Note: definition based on Code of Virginia § 38.2-3407.15.)

**Claimant:** A person or entity who submits a claim, or on whose behalf a claim is submitted. (Includes “consumer” for URAC’s Core Standards.)

**Claims Administrator:** Any entity that recommends or determines to pay claims to enrollees, physicians, hospitals, or others on behalf of the health benefit plan. Such payment determinations are made on the basis of contract provisions. Claims administrators may be insurance companies, self-insured employers,
third party administrators, or other private contractors.

**Claims Processing Organization:** An organization that seeks accreditation under these standards. Examples of organizations that process claims include but are not limited to:

- Health insurance companies;
- Health maintenance organizations (HMOs);
- Preferred provider organizations (PPOs);
- Third-party administrators (TPAs);
- Disability insurance carriers; and
- Workers’ compensation insurance carriers.

**Interpretive Note for term “Claims Processing Organization”:** Throughout this document the term “organization” refers to claims processing organization.

**Clean Claim:** A claim that has no material defect, impropriety, lack of any required substantiating documentation, or special circumstance(s) – such as, but not limited to, coordination of benefits, pre-existing conditions, subrogation, or suspected fraud – that prevents timely adjudication of the claim.

**Client:** A business or individual that purchases services from the Organization.

**Interpretive Note for term “Client”:** Here are some examples of client relationships:

- If a health plan provides health coverage to an employer, the employer is the client.
- If a health plan contracts for utilization management services from a utilization management organization, the health plan is the client.
- If a PPO contracts for credentialing services with a CVO, the PPO is the client.

**Clinical Activities:** Operational processes related to the delivery of clinical triage and health information services performed by clinical staff.

**Clinical Decision Support Tools:** Protocols, guidelines, or algorithms that assist in the clinical decision-making process.
Clinical Director: A health professional who: (1) is duly licensed or certified; (2) is an employee of, or party to a contract with, an organization; and (3) who is responsible for clinical oversight of the utilization management program, including the credentialing of professional staff and quality assessment and improvement functions.

Clinical Peer: A physician or other health professional who holds an unrestricted license and is in the same or similar specialty as typically manages the medical condition, procedures, or treatment under review. Generally, as a peer in a similar specialty, the individual must be in the same profession, i.e., the same licensure category as the ordering provider.

Clinical Practice Guidelines: Systematically developed statements to assist decision-making about appropriate health care for specific clinical circumstances.

Clinical Rationale: A statement that provides additional clarification of the clinical basis for a non-certification determination. The clinical rationale should relate the non-certification determination to the patient’s condition or treatment plan, and should supply a sufficient basis for a decision to pursue an appeal.

Clinical Review Criteria: The written screens, decision rules, medical protocols, or guidelines used by the organization as an element in the evaluation of medical necessity and appropriateness of requested admissions, procedures, and services under the auspices of the applicable health benefit plan.

Clinical Staff: Employees or contracted consultants of the health care organization who are clinically qualified to perform clinical triage and provide health information services.

Clinical Triage: Classifying consumers in order of clinical urgency and directing them to appropriate health care resources according to clinical decision support tools.

Comparable: Data about performance is compared to an historical baseline (which may be internal) and ongoing progress is recorded in regular intervals (e.g., monthly, quarterly, or annually). External benchmarks also may be used for purposes of comparison.
Complaint: An expression of dissatisfaction by a consumer expressed verbally or in writing regarding an organization’s products or services that is elevated to a complaint resolution system.

Interpretive Note for term “Complaint”:

- This term is sometimes referred to as “grievance.”
- This definition does not include appeals.

Concurrent Review: Utilization management conducted during a patient’s hospital stay or course of treatment (including outpatient procedures and services). Sometimes called “continued stay review”.

Condition: A diagnosis, clinical problem or set of indicators such as signs and symptoms a consumer may have that define him/her as eligible and appropriate to participate in a disease management program.

Conflict of Interest: Any relationship or affiliation on the part of the organization or a reviewer that could compromise the independence or objectivity of the independent review process. Conflict of interest includes, but is not limited to:

- An ownership interest of greater than 5% between any affected parties;
- A material professional or business relationship;
- A direct or indirect financial incentive for a particular determination;
- Incentives to promote the use of a certain product or service;
- A known familial relationship;
- Any prior involvement in the specific case under review.

Consumer: An individual person who is the direct or indirect recipient of the services of the Organization. Depending on the context, consumers may be identified by different names, such as “member,” enrollee,” “beneficiary,” “patient,” “injured worker,” “claimant,” etc. A consumer relationship may exist even in cases where there is not a direct relationship between the consumer and the Organization. For example, if an individual is a member of a health plan that relies on the services of a utilization management organization, then the individual is a consumer of the utilization management organization.

Interpretive Note for term “Consumer”:

In the case of a consumer who is unable to participate in the decision-making process, a family member or other individual legally authorized to make health care decisions on the consumer behalf may be a consumer for the purposes of these standards.
**Contractor:** A business entity that performs delegated functions on behalf of the Organization.

**Interpretive Note for term “Contractor”:**

For the purposes of these standards, the term “contractor” includes only those contractors that perform functions related to the key processes of the Organization. It is not URAC’s intent to include contractors that provide services unrelated to key processes. For example, a contractor that provides catering services would not fall within the definition of “contractor” in these standards. Conversely, a company that provides specialty physician reviewers to a UM organization would clearly fall within the definition of “contractor.”

**Covered Benefits:** The specific health services provided under a health benefits program, including: cost-sharing and other financial features; claims submission and reimbursement processes; requirements and processes (if any) for prior authorization or other approval of health services.

**Covered Service:** A health care service for which reimbursement or other remuneration is provided to a consumer or on behalf of a consumer under the terms of the consumer’s benefits program.

**Credentials Verification:** A process of reviewing and verifying specific credentialing criteria of a practitioner.

**Credentials Verification Organization (CVO):** An organization that gathers data and verifies the credentials of health care practitioners.

**Criteria:** A broadly applicable set of standards, guidelines, or protocols used by the organization to guide the clinical processes. Criteria should be:

- Written;
- Based on professional practice;
- Literature-based;
- Applied consistently; and
- Reviewed, at a minimum, annually.
**Data Integrity:** The quality or condition of being accurate, complete and valid, and not altered or destroyed in an unauthorized manner.

**Date of Receipt:** The date on which a claim arrives at an Organization (or, for claims that arrive on a non-business day, the date of the first business day thereafter).

**Delegation:** The process by which an organization contracts with or otherwise arranges for another entity to perform functions and to assume responsibilities covered under these standards on behalf of the organization, while the organization retains final authority to provide oversight to the delegate.

**Discharge Planning:** The process that assesses a patient’s needs in order to help arrange for the necessary services and resources to effect an appropriate and timely discharge or transfer from current services or level of care.

**Disease Management (DM):** According to the Disease Management Association of America, “Disease management is a system of coordinated healthcare interventions and communications for populations with conditions in which patient self-care efforts are significant. Disease management: supports the physician or practitioner/patient relationship and plan of care, emphasizes prevention of exacerbations and complications utilizing evidence-based practice guidelines and patient empowerment strategies, and evaluates clinical, humanistic, and economic outcomes on an ongoing basis with the goal of improving overall health. Disease management components include: population identification processes; evidence-based practice guidelines; collaborative practice models to include physician and support-service providers; patient self-management education (may include primary prevention, behavior modification programs, and compliance/surveillance); process and outcomes measurement, evaluation, and management; routine reporting/feedback loop (may include communication with patient, physician, health plan and ancillary providers, and practice profiling).”

**Disease Management Program:** A program or entity that provides the scope of functions and activities necessary to provide disease management.

**Electronic:** Mode of electronic transmission including the Internet (wide-open), Extranet (using Internet technology to link a business with information only accessible to collaborating parties), leased lines, dial-up lines, private networks, and those transmissions that are physically moved from one location to another using magnetic tape, disk, or compact disk media. (Final Rule, Department of Health and Human

**Electronic Health Record:** An electronic record of health-related information on an individual that conforms to nationally recognized interoperability standards and that can be created, managed, and consulted by authorized clinicians and staff across more than one health care organization.

**Electronic Medical Record:** An electronic record of health-related information on an individual that can be created, gathered, managed, and consulted by authorized clinicians and staff within one health care organization.

**Engagement:** Proactive outbound contact with consumers, by phone or mail, within some specified time frame of identification of eligible consumers, with tracking of interactions.

**Evidence-based:** Recommendations based on valid scientific outcomes research, preferably research that has been published in peer reviewed scientific journals. Evidence-based information can be used to develop protocols, pathways, standards of care or clinical practice guidelines and related educational materials.


**Expedited Appeal:** An appeal of a non-certification involving imminent or ongoing services.

**Facility:** An institution that provides health care services.

**Facility Rendering Service:** The institution or organization in or by which the requested admission, procedure, or service is provided. Such facilities may include, but are not limited to: hospitals; outpatient
surgical facilities; individual practitioner offices; rehabilitation centers; residential treatment centers; skilled
nursing facilities; laboratories; imaging centers; and other organizational providers of direct services to
patients.

**Family:** Individuals whom the consumer chooses to involve in the decision-making process regarding the
consumer’s health care. In the case of a consumer who is unable to participate in the decision-making
process, “family” shall include any individual legally authorized to make health care decisions on the
consumer’s behalf.

**Health Assessment Tool:** A data collection instrument that allows a consumer and the organization to
ascertain the consumer’s health status and health risks. Such tools are often administered online,
although other formats are permissible. Other common terms used to describe health assessment tools
are “health risk assessment” and “health survey.”

**Health Benefits Plan:** An arrangement to pay for medical services provided to a consumer. “Health
benefits plan” includes (but is not limited to):

- HMOs;
- PPOs;
- Indemnity health insurance programs;
- Self-insured plans;
- Public programs, such as Medicare and Medicaid; and
- Workers’ Compensation insurance programs.

**Health Care Team:** The attending physician and other health care providers with primary responsibility for
the care provided to a consumer.

**Health Content Reviewer:** An individual who holds a license or certificate as required by the appropriate
jurisdiction in a health care field (where applicable), has professional experience in providing relevant
direct patient care or has completed formal training in a health-related field.
**Health Information:** Educational resources designed to enhance the knowledge and understanding of health topics to promote wellness and self-care.

**Health Information Exchange:** The electronic movement of health-related information among organizations according to nationally recognized standards.

**Health Information Organization:** An organization that oversees and governs the exchange of health-related information among organizations according to nationally recognized standards.

**Health Literacy:** The degree to which individuals have the capacity to obtain, process, and understand basic health information and services needed to make appropriate decisions regarding their health.

**Health Professional:** An individual who: (1) has undergone formal training in a health care field; (2) holds an associate or higher degree in a health care field, or holds a state license or state certificate in a health care field; and (3) has professional experience in providing direct patient care.

**Health-Related Field:** A professional discipline that promotes the physical, psychosocial, or vocational well being of individual persons.

**Individually Identifiable Information:** Any information that can be tied to an individual consumer, as defined by applicable laws.

**Independent Review:** A process, independent of all affected parties, to determine if a health care service is medically necessary and medically appropriate or experimental/investigational. Independent review typically (but not always) occurs after all appeals mechanisms available within the health benefits plan have been exhausted. Independent review can be voluntary or mandated by law.
**Interpretive Note for term “Independent Review”:** The IRO standards focus on cases regarding questions of medical necessity/medical appropriateness and whether a treatment is experimental/investigational. Some independent review programs may also address administrative issues and disputes. However, these issues are outside the scope of these standards. Therefore, accreditation under these standards should not be interpreted (or represented) as an endorsement of an organization’s process for conducting reviews of administrative disputes.

**Independent Reviewer:** The individual (or individuals) selected by the organization to consider a case. Selection of the reviewer(s) for a case must be conducted in accordance with standards IR 1 and IR 6. All reviewer(s) who are health care practitioners must have the following qualifications:

- Active licensure;
- Recent experience or familiarity with current body of knowledge and medical practice;
- At least 5 years experience providing health care;
- If the reviewer is an M.D. or D.O., board certification by a medical specialty board approved by the American Board of Medical Specialties or the American Osteopathic Association.
- If the reviewer is a D.P.M., board certification by the American Board of Podiatric Surgery.

**Initial Clinical Review:** Clinical review conducted by appropriate licensed or certified health professionals. Initial clinical review staff may approve requests for admissions, procedures, and services that meet clinical review criteria, but must refer requests that do not meet clinical review criteria to peer clinical review for certification or non-certification. Sometimes referred to as “first level review.”

**Interoperability:** Ability of two or more systems or components to exchange information and to use the information that has been exchanged.
Knowledge Domains – Areas of specific expertise.

License: A license or permit (or equivalent) to practice medicine or a health profession that is 1) issued by any state or jurisdiction in the United States; and 2) required for the performance of job functions.

Interpretive Note for term "License":

In this definition, the word “equivalent” includes certifications, registrations, permits, etc. Specific terms will vary by state and health profession.

Medical Director: A doctor of medicine or doctor of osteopathic medicine who is duly licensed to practice medicine and who is an employee of, or party to a contract with, an organization, and who has responsibility for clinical oversight of the organization’s utilization management, credentialing, quality management, and other clinical functions.

Medical Management – A general term encompassing activities such as utilization management, case management, and the clinical aspects of quality management.

Non-Certification: A determination by an organization that an admission, extension of stay, or other health care or pharmacy service has been reviewed and, based on the information provided does not meet the clinical requirements for medical necessity, appropriateness, or effectiveness under the applicable health benefit plan.

Non-Clinical Administrative Staff: Staff who do not meet the definition of health professional (including intake personnel).

Non-Clinical Staff: Employees or contracted consultants of a health care organization who do not perform clinical assessments or provide callers with clinical advice. They may be responsible for obtaining demographic information, providing benefit information, and re-directing callers.
Off-shoring: The relocation of an organizational function to a foreign country under the same organizational control (ownership).

Note related to off-shoring: In health care management, outsourcing distinct functions to a foreign subcontractor is the more common trend. See the definition for "outsourcing."

Opt-in: Affirmative consent actively provided by a consumer to participate in an activity or function of the program, provided after the program has fully disclosed the terms and conditions of participation to the consumer.

Note for term "Opt-in": Auto enrollees are not considered "opt-in" enrollees of the program.

Opt-out: A process by which an enrolled consumer declines to participate in an activity or function of the program.

Ordering Provider: The physician or other provider who specifically prescribes the health care service being reviewed.

Organization: A business entity that seeks accreditation under these standards.

Interpretive Note for term “Organization”: This can include a program or department and can be geographically defined.

Organizational Conflict of Interest: A conflict that affects objectivity between the organization's financial interests and the organization's obligations to the client.

Outcome: A consumer's health status following services.

Outsourcing: The delegation of services or functions from internal production to an external entity outside of the United States.

Oversight: Monitoring and evaluation of the integrity of relevant program processes and decisions affecting consumers.
Participant (participating): An eligible consumer or treating provider that has had one or more inbound or outbound contacts with the disease management program, and if a consumer, has not opted out of the program.

Participating Provider – A provider that has entered into an agreement with the organization to be part of a provider network.

Patient: The enrollee or covered consumer for whom a request for certification may or may not have been filed.

Interpretive Note for term “Patient”: In the case of a patient who is unable to participate in the decision-making process, a family member or other individual legally authorized to make health care decisions on the patient’s behalf may be a patient for the purposes of these standards.

Peer Clinical Review: Clinical review conducted by appropriate health professionals when a request for an admission, procedure, or service was not approved during initial clinical review. Sometimes referred to as “second level review.”

Peer-to-Peer Conversation: A request by telephone for additional review of a utilization management determination not to certify, performed by the peer reviewer who reviewed the original decision, based on submission of additional information or a peer-to-peer discussion.

Personal Health Record: An electronic record of health-related information on an individual that conforms to nationally recognized interoperability standards and that can be drawn from multiple sources while being managed, shared, and controlled by the individual.

Personally-identifiable Information: Any information that can be tied to an individual identifier.

**Plain Language:** Communication that uses short words and sentences, common terms instead of (medical) jargon, and focuses on the essential information recipients need to understand.

**Population:** Depending on the model of the disease management program, the population for which it is responsible may be all of the consumers identified with the disease condition, or the population may be only those consumers identified to the disease management program by client referral or another mechanism. In some instances the disease management program may be responsible for identification of the population, and in other instances the client may conduct identification (and stratification) activities.

**Potential Enrollees:** Employees and eligible dependents of employer/purchasers who are offering enrollment in the organization’s products as part of the employee benefits package. In the case of organizations that offer products in the individual market, potential enrollees include individuals from the general public in the geographic area where the organization offers the products.

**Practitioner** – An individual person who is licensed to deliver health care services without supervision.

**Pre-Review Screening:** Automated or semi-automated screening of requests for authorization that may include: (1) collection of structured clinical data (including diagnosis, diagnosis codes, procedures, procedure codes); (2) asking scripted clinical questions; (3) accepting responses to scripted clinical questions; and (4) taking specific action (certification and assignment of length of stay explicitly linked to each of the possible responses). It excludes: (1) applying clinical judgment or interpretation; (2) accepting unstructured clinical information; (3) deviating from script; (4) engaging in unscripted clinical dialogue; (5) asking clinical follow-up questions; and (6) issuing non-certifications.

**Primary Physician:** The physician who is primarily responsible for the medical treatment and services of a consumer.

**Primary Source Verification or Primary Source:** Verification of a practitioner’s credentials based upon evidence obtained from the issuing source of the credential.
**Principal Reason(s):** A clinical or non-clinical statement describing the general reason(s) for the non-certification determination (“lack of medical necessity” is not sufficient to meet this).

**Professional Competency:** The ability to perform assigned professional responsibilities.

**Prospective Review:** Utilization management conducted prior to a patient’s admission, stay, or other service or course of treatment (including outpatient procedures and services). Sometimes called “precertification review” or “prior authorization.”

**Protected Health Information:** Individually identifiable health information: (1) Except as provided in paragraph (2) of this definition, that is: (i) Transmitted by electronic media; (ii) Maintained in any medium described in the definition of electronic media at Sec. 162.103 of this subchapter; or (iii) Transmitted or maintained in any other form or medium. (2) Protected health information excludes individually identifiable health information in: (i) Education records covered by the Family Educational Rights and Privacy Act, as amended, 20 U.S.C. 1232g; (ii) Records described at 20 U.S.C. 1232g(a)(4)(B)(iv); and (iii) Employment records held by a covered entity in its role as employer. (67 Fed. Reg. at 53,267 (Aug. 14, 2002); 65 Fed. Reg. at 82,805 (Dec. 28, 2000) (to be codified at 45 C.F.R. pt. 164.501)).

**Provider:** A licensed health care facility, program, agency, or health professional that delivers health care services.

**Provider Network** – A group of providers with which the organization contracts to provide health services to consumers.

**Provider-Specific Information:** Information that is sufficient to allow identification of the individual provider.

**Quality Improvement Project:** An organization-wide initiative to measure and improve the service and/or care provided by the organization.

**Quality Management (QM) program:** A systematic data-driven effort to measure and improve consumer and client services and/or health care services.

**Quality Review Study:** A scientific and systematic measurement of the effects or results of treatment modalities or practices for a particular disease or condition. The goal of quality measurement is to improve
health care services by monitoring and analyzing the data and modifying practices in response to this data.

**Rationale:** The reason(s) or justification(s) – commonly based on criteria – for a specific action or recommendation.

**Re-assessment:** Re-evaluation of an individual participating in the disease management program on a specified frequency, using the same or similar tools that were used in the initial assessment. Re-assessment may also include re-stratification.

**Referring Entity:** The organization or individual that refers a case to an organization. Referring entities may include insurance regulators, health benefits plans, consumers, and attending providers. Some states may limit by law which individuals or organizations may be a referring entity.

**Regional Health Information Organization:** A health information organization that brings together health care stakeholders within a defined geographic area and governs health information exchange among them for the purpose of improving health and care in that community.

**Retrospective Claim:** A claim presented after services have been provided (i.e., a post-service claim) and presented for consideration under a contract or policy.

**Retrospective Review:** Review conducted after services (including outpatient procedures and services) have been provided to the patient.

**Interpretive Note for term “Retrospective Review”:** Retrospective medical necessity determinations are considered utilization management (and subject to these standards).

**Review of Service Request:** Review of information submitted to the organization for health care services that do not need medical necessity certification nor result in a non-certification decision.

**Reviewer(s):** The individual (or individuals) selected by the organization to consider a case. Selection of the reviewer(s) for a case must be conducted in accordance with standards IR 1 and IR 6. All reviewer(s) who are health care practitioners must have the following qualifications:

- Active licensure;
- Recent experience or familiarity with current body of knowledge and medical practice;
- At least 5 years experience providing health care;
- If the reviewer is an M.D. or D.O., board certification by a medical specialty board approved by the American Board of Medical Specialties or the American Osteopathic Association.
• If the reviewer is a D.P.M., board certification by the American Board of Podiatric Surgery

Second Opinion: Requirement of some health plans to obtain an opinion about the medical necessity and appropriateness of specified proposed services by a practitioner other than the one originally making the recommendation.

Secondary Source Verification or Secondary Source: Verification of a practitioner’s credentials based upon evidence obtained by means other than direct contact with the issuing source of the credential (e.g., copies of licenses and certifications and data base queries).

Service Requests: Screening callers to determine the services that are necessary at the time of the call. This is usually performed by a non-clinical staff person to determine if the call is clinical and requires transfer to a clinical staff person.

Staff: The Organization’s employees, including full-time employees, part-time employees, and consultants.

Standard Appeal: An appeal of a non-certification that is not an expedited appeal. In most cases, standard appeals will not relate to cases involving urgent care. However, standard appeals may also include secondary appeals of expedited appeals.

Statistically valid: Based on accepted statistical principles and techniques.

Stratification: A process for sorting a population of eligible consumers into groups relating to the need for disease management interventions. Stratification and assessment are inter-related, and both provide data used to assign interventions. Stratification may use a variety of data sources, including but not limited to claims, pharmacy, laboratory, or consumer-reported data.

Structured Clinical Data: Clinical information that is precise and permits exact matching against explicit medical terms, diagnoses or procedure codes, or other explicit choices, without the need for interpretation.

Therapeutic: Of or relating to the treatment of disease or disorders by remedial agents or methods.

Treating Provider: The treating provider is the individual or provider group who is primarily managing the treatment for a consumer participant in the disease management program. The treating provider is not necessarily the consumers’ primary care physician. The consumer may have a different treating provider for different conditions.
Utilization Management (UM): Evaluation of the medical necessity, appropriateness, and efficiency of use of health care services, procedures, and facilities. UM encompasses prospective, concurrent and retrospective review in which clinical criteria are applied to a request. UM is sometimes called “utilization review”.

Worker: An ill or injured individual (or representative acting on behalf of the worker) who is eligible for workers’ compensation benefits and who files for, or for whom a workers’ compensation claim has been filed.

Written Agreements: A document – including an electronic document – that specifies the terms of a relationship between the Organization and a client, consumer, or contractor. This term may include a contract and any attachments or addenda.

Written Notification: Correspondence transmitted by mail, facsimile, or electronic medium. Individually identifiable health information is information that is a subset of health information, including demographic information collected from an individual, and: (1) Is created or received by a health care provider, health plan, employer, or health care clearinghouse; and (2) Relates to the past, present, or future physical or mental health or condition of an individual; the provision of health care to an individual; or the past, present, or future payment for the provision of health care to an individual; and (i) That identifies the individual; or (ii) With respect to which there is a reasonable basis to believe the information can be used to identify the individual. (65 Fed. Reg. at 82,804 (to be codified at 45 C.F.R. pt. 164.501))

(Note: This definition is derived from the federal Health Insurance Portability and Accountability Act (HIPAA)).
## Crosswalk between Core v2.1 & v3.0

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**Consumer Satisfaction**
## Core v3.0 Consumer-Facing Accreditations

### Consumer-Facing Accreditations

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Core standards apply to all accreditations unless determined to be N/A.

**ALL HEALTH ACCREDITATIONS (that include Core)**

Core standards apply if accreditation requires an organization to have credentialed clinical staff that **MUST HAVE LICENSED/CERTIFIED CLINICAL STAFF**

Core standards apply if accreditation requires an organization to have clinical staff that interfaces with consumers on an ongoing basis to provide health care management support services directly to the consumer.

**CONSUMER-FACING Health Accreditation Products that Include Core**

<table>
<thead>
<tr>
<th>Health Accreditation Products that Include Core</th>
<th>Core 1-23 &amp; 25-29</th>
<th>Core 30-35</th>
<th>Core 24 (safety QIP) &amp; Core 36-40</th>
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<td>Core (stand-alone)</td>
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1For Core stand-alone accreditation applicants, Core 30-35, 24 & 36-40 may or may not apply depending upon the type of business covered by the accreditation. URAC staff will work with these applicants to determine which standards are applicable and which ones are not.
Applicability of Core Organizational Quality Standards, Version 3.0

For the URAC Core Organizational Quality Standards version 3.0 (Core), the standards have been reorganized to ease the application process. Standards listed first apply to all health care organizations. These standards are followed by those where the applicant organization must have credentialed licensed or certified clinical staff in order to conduct the function covered by the accreditation. The last sections of Core apply to those organizations that interact directly with health care consumers on an ongoing basis in order to provide health care management support services directly to the consumer. This was done to address the fact that many of the organizations which URAC accredits under the Core standards do not interact directly with consumers, but instead are business partners or vendors to organizations that do.

The following sections of standards address infrastructure and human resources and are applicable to all organizations applying for accreditation or certification including stand-alone Core applicants, though some standards may be determined to be not applicable. For example, the delegation standards would not apply if an organization does not delegate.

- Organizational Structure
- Policies and Procedures
- Regulatory Compliance
- Inter-Departmental Coordination
- Oversight of Delegated Functions
- Marketing and Sales Communications
- Business Relationships
- Information Management
- Quality Management
- Staff Qualifications
- Staff Management
The Clinical Staff Credentialing and Oversight Role section covers clinical oversight and are applicable to organizations applying for accreditations that require licensed or certified clinical staff in order to perform the function covered by the accreditation. These standards may not apply to certain stand-alone Core applicants.

Lastly, the Core sections on Health Care System Coordination and Consumer Protection and Empowerment are applicable to organizations that are involved with care coordination and interact with consumers. These accreditation products cover functions that require an organization to interact with consumers on an ongoing basis in order to provide health care management services directly to the consumer.

The table below is color coded to show which standards are applicable under the various types of URAC accreditation products.

Blue: Standards apply regardless of accreditation, though it may be determined that some standards are not applicable (“N/A”) for a given organization.

Purple: Standards are applicable if the accreditation requires an organization to have licensed or certified clinical staff. (Note that these standards may or may not apply to an organization applying only for Core accreditation.)

Black: Standards are applicable for accreditations that require an organization to have licensed or certified clinical staff that interfaces with consumers on an ongoing basis in order to provide health care management support services directly to the consumer. (Note that these standards may or may not apply to an organization applying only for Core accreditation.)

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<td>Consumer Protection and Empowerment</td>
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### Organizational Structure

**CORE - 1 - Organizational Structure**

The *organization* has a clearly defined organizational structure outlining direct and indirect oversight responsibility throughout the *organization*. (2)

**CORE - 2 - Organization Documents**

*Organization’s* documents address: (No Weight)

- (a) Mission statement; (2)
- (b) Organizational framework for program; (2)
- (c) The population served; and (2)
- (d) Organizational oversight and reporting requirements of the program. (2)

### Policies and Procedures

**CORE - 3 - Policy and Procedure Maintenance, Review, and Approval**

The *organization*: (No Weight)

- (a) Maintains and complies with written policies and documented procedures that govern core business processes of its operations related to the scope of the accreditation; (Mandatory)
- (b) Maintains the ability to produce a master list of all such policies and procedures; (2)
- (c) Reviews written policies and documented procedures no less than annually and revises as necessary; (3)
(d) Includes the following on the master list or on all written policies and documented procedures: (No Weight)

   (i) Effective dates, review dates, including the date of the most recent revision; and (2)

   (ii) Identification of approval authority. (2)

---

### Regulatory Compliance

**CORE - 4 - Regulatory Compliance**

The *organization* implements a regulatory compliance program that: (No Weight)

(a) Tracks applicable laws and regulations in the jurisdictions where the *organization* conducts business; (Mandatory)

(b) Ensures the *organization's* compliance with applicable laws and regulations; and (Mandatory)

(c) Responds promptly to detected problems and takes corrective action as needed. (4)

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### Inter-Departmental Coordination

**CORE - 5 - Inter-Departmental Coordination**

The *organization* establishes and implements mechanisms to promote collaboration, coordination and communication across disciplines and departments within the *organization*, with emphasis on integrating administrative activities, quality improvement, and where present, clinical operations. (3)

---

### Oversight of Delegated Functions
**CORE - 6 - Delegation Review Criteria**

The organization establishes and implements criteria and processes for an assessment prior to the delegation of functions. (3)

**CORE - 7 - Delegation Review**

Prior to delegating functions to another entity, the organization: (No Weight)

(a) Establishes and implements a process to conduct a review of the potential contractor’s written policies and documented procedures and capacity to perform delegated functions; and (3)

(b) Outlines and follows criteria and processes for approving contractors. (3)

**CORE - 8 - Delegation Contracts**

The organization enters into written agreements with contractors that: (No Weight)

(a) Specify those responsibilities delegated to the contractor and those retained by the organization; (2)

(b) Require that services be performed in accordance with the organization's requirements and URAC standards; (Mandatory)

(c) Require notification to the organization of any material change in the contractor’s ability to perform delegated functions; (4)

(d) Specify that the organization may conduct surveys of the contractor, as needed; (2)

(e) Require that the contractor submit periodic reports to the organization regarding the performance of its delegated responsibilities; (3)

(f) Specify recourse and/or sanctions if the contractor does not make corrections to identified problems within a specified period; (2)
(g) Specify the circumstances under which activities may be further delegated by the contractor, including any requirements for obtaining permission from the organization before any further delegation; and (4)

(h) Specify that, if the contractor further delegates organizational functions, those functions shall be subject to the terms of the written agreement between the contractor and the organization and in accordance with URAC standards. (Mandatory)

**CORE - 9 - Delegation Oversight**

The organization establishes and implements an oversight mechanism for delegated functions within the scope of accreditation that includes: (No Weight)

(a) A periodic review (no less than annually) of the contractor's written policies and documented procedures and documentation of quality activities for related delegated functions; (2)

(b) A process to verify (no less than annually) the contractor's compliance with contractual requirements and written policies and documented procedures; and (Mandatory)

(c) A mechanism to monitor financial incentives to ensure that quality of care or service is not compromised. (3)

**Marketing and Sales Communications**

**CORE - 10 - Review of Marketing and Sales Materials**

The organization follows marketing and sales practices that include: (No Weight)

(a) Mechanisms to clearly and accurately communicate information about services inclusive of delegated activities; (3)

(b) A formal process of inter-departmental review of marketing and sales materials before dissemination to safeguard against misrepresentations about the organization's services; (3)

(c) Monitoring of existing materials for accuracy; and (3)
(d) Responds promptly to detected problems and corrective action as needed. (4)

Business Relationships

CORE - 11 - Written Business Agreements

The organization maintains signed written agreements with all clients describing the scope of the business arrangement. (2)

CORE - 12 - Client Satisfaction

The organization implements a mechanism to collect or obtain information about client satisfaction with services provided by the organization. (3)

Information Management

CORE - 13 - Information Management

The organization implements information system(s) (electronic and paper) to collect, maintain and analyze information necessary for organizational management that: (No Weight)

(a) Provides for data integrity; (Mandatory)

(b) Includes a plan for storage, maintenance and destruction; and (2)

(c) Includes a plan for interoperability: (No Weight)

   (i) Between internal information systems; and (Leading Indicator)

   (ii) With external entity information systems. (Leading Indicator)
CORE - 14 - Business Continuity

The organization implements a business continuity plan for program operations, including information system(s) (electronic and paper) that: (No Weight)

(a) Identifies which systems and processes must be maintained and the effect an outage would have on the organization’s program; (3)

(b) Identifies how business continuity is maintained given various lengths of time information systems are not functioning or accessible; (3)

(c) Is tested at least every two years; and (3)

(d) Responds promptly to detected problems and takes corrective action as needed. (3)

CORE - 15 - Information Confidentiality and Security

The organization provides for data confidentiality and security of its information system(s) (electronic and paper) by implementing written policies and/or documented procedures that address: (No Weight)

(a) Assessment of the potential risks and vulnerabilities to the confidentiality, integrity and availability of information systems; (3)

(b) Prevention of confidentiality and security breaches; and (Mandatory)

(c) Detection, containment and correction of confidentiality and security violations. (Mandatory)

CORE - 16 - Confidentiality of Individually-Identifiable Health Information

The organization implements written policies and/or documented procedures to protect the confidentiality of individually-identifiable health information that:

(No Weight)

(a) Identifies how individually-identifiable health information will be used; (Mandatory)
(b) Specifies that individually-identifiable health information is used only for purposes necessary for conducting the business of the organization, including evaluation activities; (Mandatory)

(c) Addresses who will have access to individually-identifiable health information collected by the organization; (Mandatory)

(d) Addresses oral, written or electronic communication and records that are transmitted or stored; (Mandatory)

(e) Address the responsibility of organization employees, committee members and board members to preserve the confidentiality of individually-identifiable health information; and (Mandatory)

(f) Requires employees, committee members and board members of the organization to sign a statement that they understand their responsibility to preserve confidentiality. (Mandatory)

Quality Management

CORE - 17 - Quality Management Program

The organization maintains a quality management program that promotes objective and systematic measurement, monitoring and evaluation of services and implements quality improvement activities based upon the findings. (Mandatory)

CORE - 18 - Quality Management Program Resources

The organization employs staff and/or provides the resources necessary to support the day-to-day operations of the quality management program. (3)

CORE - 19 - Quality Management Program Requirements

The organization has a written description for its quality management program that: (No Weight)

(a) Is approved by the organization’s appropriate oversight authority; (2)

(b) Defines the scope, objectives, activities, and structure of the quality management program; (2)
(c) Is reviewed and updated by the Quality Management Committee at least annually; (2)

(d) Defines the roles and responsibilities of the Quality Management Committee; and (2)

(e) Designates a member of senior management with the authority and responsibility for the overall operation of the quality management program and who serves on the Quality Management Committee. (3)

**CORE - 20 - Quality Management Committee**

The organization has a quality management committee that: (No Weight)

(a) Is granted authority for quality management by the organization's oversight authority; (3)

(b) Provides ongoing reporting to the organization's oversight authority; (3)

(c) Meets at least quarterly; (3)

(d) Maintains approved records of all committee meetings; (2)

(e) If applicable, includes at least one participating provider or receives input from participating providers; (4)

(f) Provides guidance to staff on quality management priorities and projects; (3)

(g) Approves the quality improvement projects to undertake; (3)

(h) Monitors progress in meeting quality improvement goals; and (3)

(i) Evaluates the effectiveness of the quality management program at least annually. (3)

**CORE - 21 - Quality Management Documentation**

The organization, as part of its quality management program, provides written documentation of: (No Weight)

(a) Objectives and approaches utilized in the quality management activities; (3)
(b) Identification and tracking and trending of performance measures relevant to the scope of the accreditation including, but not limited to: (Mandatory)

   (i) Access to services; (3)

   (ii) Complaints; and (3)

   (iii) Satisfaction; (3)

(c) Measures that are quantifiable and used to establish acceptable levels of performance; (Mandatory)

(d) Measuring baseline level of performance; (Mandatory)

(e) Re-measuring level of performance at least annually; (Mandatory)

(f) The implementation of action plans to improve or correct identified problems or meet acceptable levels of performance on measures; (Mandatory)

(g) The mechanisms to communicate the results of such activities to relevant staff; and (3)

(h) The mechanism to communicate the results of such activities to the quality management committee. (3)

CORE - 22 - Quality Improvement Projects

At any given time, the organization maintains no less than two quality improvement projects that address opportunities for error reduction or performance improvement related to the services covered by the accreditation. (Mandatory)

CORE - 23 - Quality Improvement Project Requirements

For each quality improvement project, the organization will:
(No Weight)

   (a) Establish measurable goals for quality improvement; (3)
(b) Design and implement strategies to improve performance; (3)

c) Establish projected time frames for meeting goals for quality improvement; (3)

d) Re-measure level of performance at least annually; (3)

(e) Document changes or improvements relative to the baseline measurement; and (3)

(f) Conduct an analysis if the performance goals are not met. (3)

**CORE - 24 - Consumer Organizations: Quality Improvement Projects**

For an organization that interacts with consumers: (No Weight)

(a) At least one of the two quality improvement projects must address consumer safety for the population served; and (Mandatory)

(b) If the quality improvement project is clinical in nature, then the organization demonstrates the involvement of a senior clinical staff person in judgments about the use of clinical quality measures and clinical aspects of performance. (Mandatory)

**Staff Qualifications**

**CORE - 25 - Job Descriptions**

The organization has written job descriptions for staff that address requirements pertinent to the scope of the positions' roles and responsibilities: (No Weight)

(a) Required education, training, and/or professional experience; (2)

(b) Expected professional competencies; (2)

(c) Appropriate licensure/certification requirements; and (2)

(d) Current scope of roles and responsibilities. (2)
CORE - 26 - Staff Qualifications

Staff meets qualifications as required in written job descriptions. (3)

Staff Management

CORE - 27 - Staff Training Program

The organization has an ongoing training program that includes: (No Weight)

(a) Initial orientation and/or training for all staff before assuming assigned roles and responsibilities; (2)

(b) Training in current URAC standards as appropriate to job functions; (2)

(c) Conflict of interest; (4)

(d) Confidentiality; (Mandatory)

(e) Documentation of all training provided for staff; and (2)

(f) Ongoing training, at a minimum annually, to maintain professional competency. (2)

CORE - 28 - Staff Operational Tools and Support

The organization provides staff with: (No Weight)

(a) Written policies and/or documented procedures appropriate to their jobs; (2)

(b) Clinical decision support tools as appropriate; and (2)

(c) Regulatory requirements as related to their job function. (2)
CORE - 29 - Staff Assessment Program

The organization maintains a formal assessment program for individual staff members, which includes: (No Weight)

(a) An annual performance appraisal; and (2)

(b) A review of relevant documentation produced by that individual staff member. (3)

Clinical Staff Credentialing and Oversight Role

CORE - 30 - Clinical Staff Credentialing

The organization implements a written policy and/or documented procedure to: (No Weight)

(a) Primary source verify the current licensure or certification of staff whose job description requires licensure or certification upon hire, and thereafter no less than every three (3) years; (Mandatory)

(b) Require staff to notify the organization in a timely manner of an adverse change in licensure or certification status; (Mandatory)

(c) Implement corrective action in response to adverse changes in licensure or certification status; and (Mandatory)

(d) Primary source verify current licensure and certification upon hire, and thereafter no later than scheduled expiration. (Leading Indicator)

CORE - 31 - Senior Clinical Staff Requirements

The organization designates at least one senior clinical staff person who has: (No Weight)

(a) Current, unrestricted clinical license(s) (or if the license is restricted, the organization has a process to ensure job functions do not violate the restrictions imposed by the state licensure board); (Mandatory)
(b) Qualifications to perform clinical oversight for the services provided; (Mandatory)

(c) Post-graduate experience in direct patient care; and (Mandatory)

(d) Board certification (if the senior clinical staff person is an M.D. or D.O.). (3)

**CORE - 32 - Senior Clinical Staff Responsibilities**

A senior clinical staff person's program responsibilities include: (No Weight)

(a) Provides guidance for clinical operational aspects of the program; (3)

(b) Is responsible for oversight of clinical decision-making aspects of the program; (Mandatory)

(c) Has periodic consultation with practitioners in the field; and (3)

(d) Ensures the organizational objective to have qualified clinicians accountable to the organization for decisions affecting consumers. (Mandatory)

**CORE - 33 - Financial Incentive Policy**

If the organization has a system for reimbursement, bonuses or incentives to staff or health care providers based directly on consumer utilization of health care services, then the organization implements mechanisms addressing how the organization will ensure that consumer health care is not compromised. (Mandatory)

**CORE - 34 - Access to Services**

The organization implements written policies and/or documented procedures to ensure access to services covered by the accreditation. (Mandatory)

**CORE - 35 - Consumer Complaint Process**

The organization maintains a formal process to address consumer complaints that includes: (No Weight)

(a) A process to receive and respond in a timely manner to complaints; (Mandatory)
(b) Notice (written or verbal) of final result with an explanation; (4)

(c) Informs consumers of the avenues to seek further review if an additional complaint review process is available; (4)

(d) Evidence of meeting the organization’s specified time frame for resolution and response; and (4)

(e) Reporting analysis of the complaints to the quality management committee. (3)

Health Care System Coordination

CORE - 36 - Coordination with External Entities

The organization establishes and implements mechanisms to promote collaboration and communication with applicable external entities to coordinate health services for consumers. (1)

Consumer Protection and Empowerment

CORE - 37 - Consumer Rights and Responsibilities

The organization implements a mechanism for informing consumers of their rights and responsibilities. (4)

CORE - 38 - Consumer Safety Mechanism

The organization has a mechanism to respond on an urgent basis to situations that pose an immediate threat to the health and safety of consumers. (Mandatory)

CORE - 39 - Consumer Satisfaction

The organization implements a mechanism to collect or obtain information about consumer satisfaction with services provided by the organization. (3)
CORE - 40 - Health Literacy

The organization will implement written policies and/or documented procedures addressing health literacy that: (No Weight)

(a) Require consumer materials to be in plain language; (Leading Indicator)

(b) Assess the use of plain language in consumer documents; and (Leading Indicator)

(c) Provide relevant information and guidance to staff that interfaces directly with, or writes content for, consumers. (Leading Indicator)
Health Utilization Management, Version 6.0

Review Criteria

HUM - 1 - Review Criteria Requirements

The organization utilizes explicit clinical review criteria or scripts that are: (No Weight)

(a) Developed with involvement from appropriate providers with current knowledge relevant to the criteria or scripts under review; (3)

(b) Based on current clinical principles and processes; (3)

(c) Evaluated at least annually and updated if necessary by: (3)

  (i) The organization itself; and (3)

  (ii) Appropriate, actively practicing physicians and other providers with current knowledge relevant to the criteria or scripts under review; and (Mandatory)

  (d) Approved by the medical director (or equivalent designate) or clinical director (or equivalent designate). (Mandatory)

Accessibility of Review Services

HUM - 2 - Access to Review Staff

The organization provides access to its review staff by a toll free or collect telephone line at a minimum from 9:00 a.m. to 4:00 p.m. of each normal business day in each time zone where the organization conducts at least two percent of its review activities. (4)

HUM - 3 - Review Service Communication and Timeframes

The organization maintains processes to: (No Weight)
(a) Receive communications from providers and patients during the business day and after business hours; (4)

(b) Respond to communications within one business day; and (4)

(c) Conduct its outgoing communications related to utilization management during providers' reasonable and normal business hours, unless otherwise mutually agreed. (4)

**HUM - 4 - Review Service Disclosures**

The organization: (No Weight)

(a) Requires utilization management staff to identify themselves by name, title, and organization name; and (2)

(b) Upon request, verbally informs patients; facility personnel; the attending physician and other ordering providers; and health professionals of specific utilization management requirements and procedures. (4)

**On-Site Review Services**

**HUM - 5 - Onsite Review Requirements**

For on-site review services, the organization: (No Weight)

(a) Requires on-site reviewers to carry a picture ID with full name and the name of the organization; (2)

(b) Schedules reviews at least one business day in advance, unless otherwise agreed; and (4)

(c) Requires the on-site reviewers to follow reasonable hospital or facility procedures, including checking in with designated hospital or facility personnel. (4)
HUM - 6 - N/A
This standard number is reserved to synchronize with URAC's Workers' Compensation Utilization Management Standards. There is no current Standard UM 6. (No Weight)

**Initial Screening**

**HUM - 7 - Limitations in Use of Non-Clinical Staff**
For initial screening, the organization limits use of non-clinical administrative staff to: (No Weight)

(a) Performance of review of service request for completeness of information; (2)
(b) Collection and transfer of non-clinical data; (2)
(c) Acquisition of structured clinical data; and (2)
(d) Activities that do not require evaluation or interpretation of clinical information. (Mandatory)

**HUM - 8 - Pre-Review Screening Staff Oversight**
The organization ensures that licensed health professionals are available to non-clinical administrative staff while performing initial screening. (Mandatory)

**HUM - 9 - Pre-Review Screening Non-Certifications**
The organization does not issue non-certifications based on initial screening. (Mandatory)

**Initial Clinical Review**

**HUM - 10 - Initial Clinical Reviewer Qualifications**
Individuals who conduct initial clinical review: (No Weight)

(a) Are appropriate health professionals; and (Mandatory)
(b) Possess an active professional relevant *license*. (Mandatory)

**HUM - 11 - Initial Clinical Reviewer Resources**

Individuals who conduct *initial clinical review* have *access* to consultation with a: (3)

(a) *Licensed* doctor of medicine or doctor of osteopathic medicine; or (No Weight)

(b) *Licensed health professional* in the same licensure category as the *ordering provider*; or (No Weight)

(c) *Health professional* with the same clinical education as the *ordering provider* in clinical specialties where *licensure* is not issued. (No Weight)

**HUM - 12 - Initial Clinical Reviewer Non-Certifications**

The *organization* does not issue *non-certifications* based on *initial clinical review*. (Mandatory)

**Peer Clinical Review**

**HUM - 13 - Peer Clinical Review Cases**

The *organization* conducts *peer clinical reviews* for all *cases* where a *certification* is not issued through *initial clinical review* or *initial screening*. (Mandatory)

**HUM - 14 - Peer Clinical Reviewer Qualifications**

Individuals who conduct *peer clinical review*: (No Weight)

(a) Are appropriate *health professionals*; (Mandatory)

(b) Are qualified, as determined by the *medical director* or *clinical director*, to render a clinical opinion about the medical *condition*, procedures, and treatment under review; and (Mandatory)

(c) Hold a current and valid *license*; (Mandatory)
(i) In the same licensure category as the ordering provider; or (No Weight)

(ii) As a doctor of medicine or doctor of osteopathic medicine. (No Weight)

Peer-to-Peer Conversation

HUM - 15 - Peer-to-Peer Conversation Availability

Health professionals that conduct peer clinical review are available to discuss review determinations with attending physicians or other ordering providers. (4)

HUM - 16 - Peer-to-Peer Conversation Alternate

When a determination is made to issue a non-certification and no peer-to-peer conversation has occurred: (No Weight)

(a) The organization provides, within one business day of a request by the attending physician or ordering provider, the opportunity to discuss the non-certification decision: (No Weight)

(i) With the clinical peer reviewer making the initial determination; or (4)

(ii) With a different clinical peer, if the original clinical peer reviewer cannot be available within one business day); and (4)

(b) If a peer-to-peer conversation or review of additional information does not result in a certification, the organization informs the provider and consumer of the right to initiate an appeal and the procedure to do so. (4)

Time Frames for Initial UM Decision

HUM - 17 - Prospective Review Timeframes

For prospective review, the organization issues a determination: (No Weight)

(a) As soon as possible based on the clinical situation, but in no case later than 72 hours of the
receipt of request for a *utilization management* determination, if it is a *case involving urgent care*; or (4)

(b) Within 15 calendar days of the receipt of request for a *utilization management* determination, if it is a non-urgent *case*. (4)

(c) For non-urgent cases this period may be extended one time by the *organization* for up to 15 calendar days: (No Weight)

(i) Provided that the *organization* determines that an extension is necessary because of matters beyond the control of the *organization*; and (4)

(ii) Notifies the *patient*, prior to the expiration of the initial 15 calendar day period of the circumstances requiring the extension and the date when the plan expects to make a decision; and (4)

(iii) If a *patient* fails to submit necessary information to decide the case, the notice of extension must specifically describe the required information, and the *patient* must be given at least 45 calendar days from receipt of notice to respond to the plan request for more information. (4)

**HUM - 18 - Retrospective Review Timeframes**

For *retrospective review*, the *organization* issues a determination: (No Weight)

(a) Within 30 calendar days of the receipt of request for a *utilization management* determination; and (3)

(b) This period may be extended one time by the *organization* for up to 15 calendar days: (No Weight)

(i) Provided that the *organization* determines that an extension is necessary because of matters beyond the control of the *organization*; and (4)

(ii) Notifies the *patient*, prior to the expiration of the initial 30 calendar day period of the circumstances requiring the extension and the date when the plan expects to make a decision; and (4)
(iii) If a patient fails to submit necessary information to decide the case, the notice of extension must specifically describe the required information, and the patient must be given at least 45 calendar days from receipt of notice to respond to the plan request for more information. (4)

**HUM - 19 - Concurrent Review Timeframes**

For concurrent review, the organization adheres to the following time frames: (No Weight)

(a) For reductions or terminations in a previously approved course of treatment, the organization issues the determination early enough to allow the patient to request a review and receive a decision before the reduction or termination occurs; and (4)

(b) For requests to extend a current course of treatment, the organization issues the determination within: (No Weight)

(i) 24 hours of the request for a utilization management determination, if it is a case involving urgent care and the request for extension was received at least 24 hours before the expiration of the currently certified period or treatments; or (4)

(ii) 72 hours of the request for a utilization management determination, if it is a case involving urgent care and the request for extension was received less than 24 hours before the expiration of the currently certified period or treatments. (4)

**Notice of Certification Decisions**

**HUM - 20 - Certification Decision Notice and Tracking**

For certifications, the organization: (No Weight)

(a) Has a process for notification of the attending physician or other ordering provider, facility rendering service, and patient: (4)

(b) Includes tracking information (such as reference number) in the notice of certification; and (3)
Upon request from the attending physician or other ordering provider, facility rendering service, or patient, provides written notification of any certification. (4)

**HUM - 21 - Continued Certification Decision Requirements**

Confirmation of certification for continued hospitalization or services includes the number of extended days or units of service, the next anticipated review point, the new total number of days or services approved, and the date of admission or onset of services. (3)

**Notice of Non-Certification Decisions**

**HUM - 22 - Written Notice of Non-Certification Decisions and Rationale**

For non-certifications, the organization issues written notification of the non-certification decision to the patient and attending physician or other ordering provider or facility rendering service that includes: (No Weight)

(a) The principle reasons for the determination not to certify; (1)

(b) A statement that the clinical rationale used in making the non-certification decision will be provided, in writing, upon request; and (4)

(c) Instructions for: (No Weight)

   (i) Initiating an appeal of the non-certification; and (Mandatory)

   (ii) Requesting a clinical rationale for the non-certification. (Mandatory)

**HUM - 23 - Clinical Rationale for Non-Certification Requirements**

Upon request from the patient, attending physician, or other ordering provider or facility rendering service, the organization provides specific clinical rationale upon which the non-certification was based. (4)
UM Procedures

HUM - 24 - Reversal of Certification Determinations

The organization does not reverse a certification determination unless it receives new information that is relevant to the certification and that was not available at the time of the original certification. (4)

HUM - 25 - Frequency of Continued Reviews

The organization ensures that the frequency of reviews for the extension of initial determinations is based on the severity or complexity of the patient’s condition or on necessary treatment and discharge planning activity (i.e., not routinely conducted on a daily basis). (4)

Information Upon Which UM is Conducted

HUM - 26 - Scope of Review Information

The organization, when conducting routine prospective review, concurrent review, or retrospective review: (No Weight)

(a) Accepts information from any reasonably reliable source that will assist in the certification process; (2)

(b) Collects only the information necessary to certify the admission, procedure or treatment, length of stay, or frequency or duration of services. (2)

(c) Does not routinely require hospitals, physicians, and other providers to numerically code diagnoses or procedures to be considered for certification, but may request such codes, if available; (4)

(d) Does not routinely request copies of all medical records on all patients reviewed; (4)

(e) Requires only the section(s) of the medical record necessary in that specific case to certify medical necessity or appropriateness of the admission or extension of stay, frequency or duration of service, or length of anticipated inability to return to work; and (4)
(f) Administers a process to share all clinical and demographic information on individual patients among its various clinical and administrative departments that have a need to know, to avoid duplicate requests for information from enrollees or providers. (3)

**HUM - 27 - Prospective and Concurrent Review Determinations**

For prospective review and concurrent review, the organization bases review determinations solely on the medical information obtained by the organization at the time of the review determination. (4)

**HUM - 28 - Retrospective Review Determinations**

For retrospective review, the organization bases review determinations solely on the medical information available to the attending physician or ordering provider at the time the medical care was provided. (4)

**HUM - 29 - Lack of Information Policy and Procedure**

The organization implements policies and procedures to address situations in which it has insufficient information to conduct a review. Such policies and procedures provide for: (4)

(a) Procedural time frames that are appropriate to the clinical circumstances of the review (i.e., prospective, concurrent, retrospective reviews); (4)

(b) Resolution of cases in which the necessary information is not provided to the organization within specified time frames; and (4)

(c) Processes by which the organization issues an administrative non-certification due to lack of information. (4)

**Appeals Considerations**

**HUM - 30 - Non-Certification Appeals Process**

The organization maintains a formal process to consider appeals of non-certifications that includes: (No Weight)
(a) The availability of *standard appeal* for non-urgent cases and *expedited appeal* for cases involving urgent care; and (Mandatory)

(b) Written policies and procedures that: (No Weight)

   (i) Clearly describe the *appeal* process, including the right to appeal of the patient, provider, or facility rendering service; (Mandatory)

   (ii) Provide for explicit time frames for each stage of the *appeal* resolution process; and (Mandatory)

   (iii) Are available, upon request, to any patient, provider, or facility rendering service. (Mandatory)

**HUM - 31 - Appeals Process**

As part of the *appeals* process: (No Weight)

(a) The organization provides the patient, provider, or facility rendering service the opportunity to submit written comments, documents, records, and other information relating to the case; (Mandatory)

(b) Takes all such information into account during the *appeals* process without regard to whether such information was submitted or considered in the initial consideration of the case; and (Mandatory)

(c) In instance of a first level *appeal*, the organization implements the decision of the first level clinical *appeal* if it overturns the initial denial. (Mandatory)

**HUM - 32 - Appeal Peer Reviewer Qualifications**

*Appeals considerations* are conducted by *health professionals* who: (No Weight)

(a) Are clinical peers; (Mandatory)

(b) Hold an active, unrestricted *license* to practice medicine or a health profession; (Mandatory)
(c) Are board-certified (if applicable) by: (No Weight)

(i) A specialty board approved by the American Board of Medical Specialties (doctors of medicine); or (3)

(ii) The Advisory Board of Osteopathic Specialists from the major areas of clinical services (doctors of osteopathic medicine); (No Weight)

(d) Are in the same profession and in a similar specialty as typically manages the medical condition, procedure, or treatment as mutually deemed appropriate; and (Mandatory)

(e) Are neither the individual who made the original non-certification, nor the subordinate of such an individual. (Mandatory)

**HUM - 33 - Expedited Appeals Process Timeframe**

Expedited appeals are completed with verbal notification of determination to the requesting party within 72 hours of the request followed by a written confirmation of the notification within 3 calendar days to the patient and attending physician or other ordering provider or facility rendering service. (Mandatory)

**HUM - 34 - Standard Appeals Process Timeframe**

Standard appeals are completed, and written notification of the appeal decision issued, within 30 calendar days of the receipt of the request for appeal to the patient and attending physician or other ordering provider or facility rendering service. (Mandatory)

**HUM - 35 - Written Notice of Upheld Non-Certifications**

For appeals determinations, the organization issues written notification of the adverse appeal decision to the patient and attending physician or other ordering provider or facility rendering service that includes: (No Weight)

(a) The principal reasons for the determination to uphold the non-certification; (4)

(b) A statement that the clinical rationale used in making the appeal decision will be provided, in writing, upon request; and (4)
(c) Information about additional appeal mechanisms available, if any. (4)

**HUM - 36 - Appeal Record Documentation**

The organization maintains records for each appeal that includes: (No Weight)

(a) The name of the patient, provider, and/or facility rendering service; (3)

(b) Copies of all correspondence from the patient, provider, or facility rendering service and the organization regarding the appeal; (3)

(c) Dates of appeal reviews, documentation of actions taken, and final resolution; (3)

(d) Minutes or transcripts of appeal proceedings (if any); and (3)

(e) Name and credentials of the clinical peer that meets the qualifications in Standard UM 32. (3)