

# **New and Emerging Technology - Coverage Status**

LOB(s):  ⊠ Commercial	State(s): ⊠ Idaho		⊠ Washington	☐ Other:
⊠ Medicare				
⊠ Medicaid	⊠ Oregon	☐ Washington		

### **Enterprise Policy**

PacificSource is committed to assessing and applying current regulatory standards, widely-used treatment guidelines, and evidenced-based clinical literature when developing clinical criteria for coverage determination. Each policy contains a list of sources (references) that serves as the summary of evidence used in the development and adoption of the criteria. The evidence was considered to ensure the criteria provide clinical benefits that promote patient safety and/or access to appropriate care. Each clinical policy is reviewed, updated as needed, and readopted, at least annually, to 84112reflect changes in regulation, new evidence, and advancements in healthcare.

Clinical Guidelines are written when necessary to provide guidance to providers and members in order to outline and clarify coverage criteria in accordance with the terms of the Member's policy. This Clinical Guideline only applies to PacificSource Health Plans, PacificSource Community Health Plans, and PacificSource Community Solutions in Idaho, Montana, Oregon, and Washington. Because of the changing nature of medicine, this list is subject to revision and update without notice. This document is designed for informational purposes only and is not an authorization or contract. Coverage determinations are Member policies differ in benefits and to the extent a conflict exists between the Clinical Guideline and the Member's policy, the Member's policy language shall control. Clinical Guidelines do not constitute medical advice nor guarantee coverage.

# **Background**

New and emerging medical and behavioral healthcare procedures, pharmaceuticals, and devices (collectively "technologies") are often prescribed by physicians and/or marketed to the public before FDA or other government agency approval, or research is available in peer-reviewed literature documenting efficacy, safety, and long-term positive outcomes.

New and emerging technologies are reviewed by the New Technologies and Operational Criteria (NTOC) Committee which is chaired by a PacificSource Medical Director. The PacificSource Behavioral Health Medical Director or behavioral healthcare professional designee is involved in the decision-making process for behavioral healthcare services. Pharmaceuticals are reviewed by the PacificSource Pharmacy and Therapeutics (P&T) committee.

The NTOC Committee bases its recommendation of coverage on review and evaluation of the following resources:

- Available peer-reviewed and evidenced-based literature
- Survey of standards of care and coverage
- Consultation with specialists and expert professionals
- PacificSource group and individual contracts
- Medicare and Medicaid requirements

This "New and Emerging Technologies – Coverage Status" policy outlines the evaluation process of new and emerging technology as well as coverage status of items considered experimental, investigational, or unproven.

#### **Procedure**

#### **Commercial**

#### **Evaluation Process**

The NTOC Committee reviews and evaluates new technology and new application of existing technology of medical and behavioral healthcare procedures and devices. NTOC Committee members represent key departments and stakeholders who have operational insight or responsibility for applying the criteria developed by the Committee. A PacificSource Medical Director chairs the NTOC Committee and ensures the Behavioral Health Medical Director or behavioral healthcare professional designee is involved in the decision-making process for behavioral healthcare services.

Agenda items for the NTOC Committee to review for coverage status are collected from multiple sources, which include, but are not limited to:

- New CPT or HCPCS codes
- New FDA approvals
- Provider inquiries
- Reports of new technology acquired by a community provider or anticipated to have widespread acceptance
- Utilization reviews and trends
- Vendor requests vendor requests for reassessment of coverage position are limited to an annual review unless there is a change in FDA status or Hayes, Inc. rating

To inform its decision-making, the NTOC Committee reviews peer-reviewed and evidence-based information, which indicates if the technology is in general use or a community standard, is under continued scientific review (testing/research), is shown to have a demonstratable benefit, or is shown to be safe and efficacious). The reviews may consist of the following:

- Technology assessment consisting of:
  - Information from appropriate government regulatory bodies such as Food and Drug Administration (FDA) and Centers for Medicare & Medicaid Services (CMS)
    - FDA approval alone, is not a basis for coverage
  - o Assessment of peer-reviewed literature and their conclusions concerning:
    - Effect of the technology on health outcomes, with emphasis on random controlled clinical trial outcomes
    - Evidence comparing new technology to established alternatives
    - Results attained outside of investigational settings, with emphasis on studies that were not underwritten by the manufacturer or other sponsor with financial interest in the service or technology

- Report on long term studies indicating improved health outcomes and clinical trials now recruiting or in process
- Information available from evidence-based resources may vary depending on treatment procedure or device. The evidenced-based resources, which relies on the judgment of experts, include the following resources, with additional resources reviewed as necessary based on the technology being reviewed:
  - Agency for Healthcare Research and Quality (AHRQ)
  - Carelon Medical Benefits Management Clinical Guidelines, formerly AIM (AIM Specialty Health) Clinical Guidelines
  - Alliance of Community Health Plans (ACHP)
  - American College of Radiology® (ACR)
  - American Hospital Formulary Service Drug Information (AHFS® DI™)
  - o Bree Collaborative Foundation for Health Care Quality
  - Centers for Disease Control and Prevention (CDC)
  - Cochrane Collaboration
  - Facts and Comparison®
  - Hayes, Inc., and Hayes Genetic Testing Evaluation Service
  - MCG Health
  - MEDLINE® (component of PubMed®)
  - Micromedex®
  - National Comprehensive Cancer Network® (NCCN)
  - National Institute for Health and Care Excellence (NICE)
  - Oregon Health Evidence Review Commission (HERC)
  - Professional Societies Recommendations and Practice Guidelines
  - UpToDate®
  - o U.S. Pharmacopeia Dispensing Information
  - Washington Health Technology Assessment (HTA)
  - Washington Health Technology Clinical Committee (HTCC)
- Survey of similar market carriers and their published coverage position and/or medical policy concerning the service of technology under review
- Utilization and authorization data, as available and applicable

The PacificSource Medical Director may seek input or consult with specialists and professionals who have expertise in the technology when additional information is needed.

The NTOC committee, which includes a Medical Director, makes a determination based on the review of the technology that results in one of two options:

Covered service, which may include developing specific clinical guidelines criteria

- Deemed experimental, investigational, or unproven (E/I/U) and added to the "New and Emerging Technologies – Coverage Status" policy or as an E/I/U item to an existing policy related to the technology reviewed
  - A technology may be indicated for archival (i.e., remains unproven but only requires a review upon request) when the following conditions are met:
    - The scientific evidence does not support clinical efficacy of the technology demonstrated by ONE of the following:
      - There is no general use within the medical community, i.e., the technology does not meet community standards
      - There have not been clinical studies for five or more years, indicating the technology is no longer under scientific review
      - The technology has not been proven to have a demonstratable benefit or been shown to be safe and efficacious
    - Claims utilization indicates PacificSource has not received a request for a period of at least two years
  - The unproven technology which has been determined to be archivable, will be moved to an excel sheet (NTOC Agenda Review Schedule) on the <u>Special Function SharePoint site</u> and noted as archived in the modification history along with associated codes

### **Quality Oversight**

The "New and Emerging Technologies – Coverage Status" policy will be reviewed at least annually.

In addition, an annual report summarizing the NTOC review activity is presented to our Clinical Quality and Utilization Management (CQUM) committee, which consists of external providers and PacificSource Medical Directors, for review.

#### Medicaid

PacificSource Community Solutions (PCS) follows Guideline Notes 172 and 173 of the OHP Prioritized List of Health Services for guidance on New and Emerging Technology. In the absence of OHP guidance, PCS will follow this policy.

#### **Medicare**

PacificSource Medicare follows CMS guidelines and criteria. In the absence of CMS guidelines and criteria, PacificSource Medicare will follow internal policy for determination of coverage and medical necessity.

# **Experimental/Investigational/Unproven Determinations for Coverage Status**

The new and emerging medical technologies, in the following list have been determined by the NTOC Committee to be experimental, investigational, or unproven (E/I/U) and therefore are not covered. The impact of these technologies on health outcomes has not been established as the current scientific evidence is either not yet sufficient or does not support clinical efficacy.

The NTOC Committee has the authority to add new technologies or revise the determinations listed below based on additional review of current scientific evidence, advice, and recommendations by the NTOC Committee.

## **Experimental/Investigational/Unproven Determinations for Coverage Status**

The following list of codes are for informational purposes only and may not be all-inclusive. Deleted codes and codes which are not effective at the time the service is rendered may not be eligible for reimbursement.

PROCEDURE	CPT HCPCS	COVERAGE STATUS
AcuDetox (Auricle Acupuncture) treatment for prevention, treatment, and harm reduction of substance use.	No Specific Code 97810 97811	Not Covered (Experimental/Investigational/Unproven)
		(Refer to Contract for coverage as well)
Allogeneic haemopoietic stem cell transplantation (HSCT) for Crohn's Disease	No Specific Code 36240	Not Covered (Experimental/Investigational/Unproven)
Artificial Iris (Custom Flex) for the Treatment of Aniridia (e.g. Custom Flex Artificial Iris)	C1839	Not Covered (Experimental/Investigational/Unproven)
Artificial Retina prosthesis device (Retinal Prosthesis) (e.g., the Angus I & II)	C1842 C1841 L8608	Not Covered (Experimental/Investigational/Unproven)
Avise CTD, Avise SLE, Avise MCV, Avise MTX, SLE-key-Rule Out Test - Prognostic tests for Systemic Lupus Erythematosus (SLE) and Connective Tissue diseases	0312U, 0062U	Not Covered (Experimental/Investigational/Unproven)
Cala Trio (Cala Health, Inc.) for Treatment of Essential Tremor – (also referred to as transcutaneous afferent patterned stimulation (TAPS)) (e.g.; Cala One and Cala Trio)	E0734; A4542	Not Covered (Experimental/Investigational/Unproven)
Coil embolization of hemorrhoids (Emborrhoid technique) embolization of the hemorrhoidal arteries.	No Specific Code 37241, 37244	Not Covered (Experimental/Investigational/Unproven)
Continuous Passive Motion (CPM) for Knees and all other indications	E0935, E0936	Not Covered (Experimental/Investigational/Unproven) (Legacy Employee Health Plan covers for knees)

PROCEDURE	СРТ	COVERAGE STATUS
	HCPCS	
Cryoablation for chronic rhinitis	No Specific Code	Not Covered
(allergic or nonallergic) (e.g., ClariFix device)	31243	(Experimental/Investigational/Unproven)
Cranial Electrotherapy Stimulation	E0732, A4596	Not Covered
(CES), also known as cranial		(Experimental/Investigational/Unproven)
electrical stimulation, transcranial		
electrical stimulation (e.g., Alpha		
Stim Device, CES Ultra ) for all indications		
	No specific Code	National
Deep Brain Stimulation (e.g. Reclaim DBS) for Obsessive-	CPT Codes:	Not Covered
compulsive disorder and comorbid	61863 61864	(Experimental/Investigational/Unproven)
psychiatric disorders.	61867 61868	
	61880 61850	
	61860	
	HCPC Codes: L8680 L8685	
	L8686 L8687	
	L8688	
DISC Nucleoplasty	S2348	Not Covered
(Radiofrequency Coblation) a		(Experimental/Investigational/Unproven)
percutaneous disc decompression		
(PDD) or radiofrequency coblation		
-used to treat herniated discs (e.g., ArthroCare System, Per-D Spine		
Wand)		
EarliPoint™ Evaluation -	No specific Code	Not Covered
diagnosing and assessing autism	97110 97112	(Experimental/Investigational/Unproven)
in children 16 to 30 months old		(Experimental/investigational/emprevent)
Esophageal cooling device (e.g.,	No Specific Code	Not Covered
ensoETM) for all indications	43499 C1889	(Experimental/Investigational/Unproven)
External Trigeminal Nerve	E0733 A4541	Not Covered
Stimulation (eTNS) System (e.g.		(Experimental/Investigational/Unproven)
Monarch; NeuroSigma) indication		
ADHD in children aged 7 to 12		
years	20000	
Extracorporeal shock wave	28890	Not Covered
therapy (ESWT)		(Experimental/Investigational/Unproven)
Gastric Emptying Breath Test -	0106U	Not Covered
(Cairn 13C-Spirulina Gastric		(Experimental/Investigational/Unproven)
Emptying Breath Test (GEBT)) -		
measurement of the rate of gastric emptying in adults		
emptymy m addits		

PROCEDURE	CPT HCPCS	COVERAGE STATUS
Gait Analysis (computerized)	96000, 96001	Not Covered
(Motion Analysis) for all indications.		(Experimental/Investigational/Unproven)
Ganglion Impar block for	No Specific Code	Not Covered
Coccydynia, pelvic pain, and all other indications	64999	(Experimental/Investigational/Unproven)
Geniculate artery embolization	No Specific Code	Not Covered
(GAE) (embolization of the knee) – treatment of osteoarthritis related	37242	(Experimental/Investigational/Unproven)
knee pain		Approved for coverage of knee hemarthrosis
Genova's GI Effects	No Specific Code	Not Covered
Comprehensive Profile analyzes for all indications	81599, 89240, 81479	(Experimental/investigational/Unproven)
Glycine Receptor Alpha1 IgG, Cell	O431U	Not Covered
Binding Assay (GLYCC) test (e.g. Stiff Person Syndrome)		(Experimental/Investigational/Unproven)
Implantable shock absorber (e.g.,	No Specific Code	Not Covered
MISHA knee system) for all indications	L8699 27599	(Experimental/investigational/Unproven)
Intense Pulse Light Therapy for	No Specific Code	Not Covered
the treatment of dry eyes from meibomian gland dysfunction	17999	(Experimental/investigational/Unproven)
Interactive Metronome Training	No Specific Code	Not Covered
(e.g., attention deficit hyperactivity disorder or any other indication)	97110, 97112	(Experimental/Investigational/Unproven)
Interferential Current Stimulation	S8130, S8131	Not Covered
Therapy (IFS/ICT) (electrical stimulation) for all indications.		(Experimental/Investigational/Unproven)
Intra-arterial Infusion	No Specific Code	Not Covered
(Embolization) of imipenem/cilastatin sodium (IPM/CS) refractory interphalangeal (DIP) joint-Osteoarthritis (OA) and all other indications.	37242 20999	(Experimental/Investigational/Unproven)
	C1761	Not Covered
Intravascular Lithotripsy (IVL) – (Shockwave intravascular lithotripsy system (IVL)) Coronary		Not Covered (Experimental/Investigational/Unproven)
iovera (Pacira Biosciences Inc.)	No Specific Code	Not Covered
System (cold injection) for Knee Osteoarthritis	64640,	(Experimental/Investigational/Unproven)

PROCEDURE	CPT HCPCS	COVERAGE STATUS
IsoPSA blood-based test to assess	0359U	Not Covered
prostate cancer risk		(Experimental/Investigational/Unproven)
Jaw Motion Rehab System (e.g.,	E1700, E1701, E1702	Not Covered
TheraBite Jaw Motion Rehabilitation System, OraStretch press, and Dynasplint Trismus System) – all indications	E1702	(Experimental/Investigational/Unproven)
Latera Nasal Implant (Absorbable nasal implants) for all indications	No Specific Code 30999 30468	Not Covered (Experimental/Investigational/Unproven)
LipiView II and LipiScan Ocular	No Specific Code	Not Covered
Surface Interferometer images	92285	(Experimental/Investigational/Unproven)
with Dynamic Meibomian Imaging (DMI)		Refer to Category III policy for "T" codes
Lymphaticovenous Anastomosis	No Specific Code	Not Covered
(LVA) (Lymphovenous bypass or shunt) for lymphedema	38999, 38308	(Experimental/Investigational/Unproven)
Magnetic Resonance Neurography	No Specific Code	Not Covered
(MRN), (also known as Magnetic	76498	(Experimental/Investigation/Unproven)
Resonance Neurogram or MR Imaging of the Peripheral Nerves (PNI)) for all indications		(Carelon does not review)
MicroGen DX qPCR & NGS Test	0112U	Not Covered
(MicroGenDX) (also known as Next Generation testing) - test for bacterial/fungi Infection (outpatient only)		(Experimental/Investigational/Unproven)
MI-HEART Ceramides Risk Score	0119U	Not Covered
(measures the risk of adverse cardiovascular events) – for all indications		(Experimental/Investigational/Unproven)
MiraDry System aka Microwave or	No Specific Code	Not Covered
electromagnetic energy (microwave thermolysis) for all indications of hyperhidrosis	17999	(Experimental/Investigational/Unproven)
Nd: YAG laser vitreolysis (YAG	No Specific Code 67031, 67299	Not Covered
Reflex Laser Systems, Ellex) - treatment of vitreous floaters or any other indication		(Experimental/Investigational/Unproven)
Navigated transcranial magnetic	No Specific Code	Not Covered
stimulation (nTMS) - presurgical planning	64999	(Experimental/Investigational/Unproven)

PROCEDURE	CPT HCPCS	COVERAGE STATUS
Neurofilament light chain (NfL) testing – for all indications	0361U	Not Covered (Experimental/Investigational/Unproven)
Nociceptive Trigeminal Inhibition Tension Suppression System (NTI- TSS or NTI-tension suppression system) for Headaches and Migraines	No Specific Code 21110, D7880, D8210	Not Covered (Experimental/Investigational/Unproven)
Nodify CDT, Nodify XL2 (BDX- XL2); EarlyCDT Test for the risk of malignancy of a lung nodule and all other indications	0360U; 0080U	Not Covered (Experimental/Investigational/Unproven)
Non-Pneumatic Active Dynamic Compression (Dayspring (Koya Medical Inc.)) for Treatment of Lymphedema	No Specific Code E0678, E0679, E0680, E0681, E0682	Not Covered (Experimental/Investigational/Unproven)
Occipital Nerve Stimulation (ONS) - intended to prevent migraines headaches and all other headaches	No Specific Code 64553; 61885; 61886; 64999	Not Covered (Experimental/Investigational/Unproven)
Occipital Nerve Decompression Surgery (ONS) (Also known as Peripheral Occipital nerve decompression surgery or migraine surgery) for Migraine Headaches	No Specific Code 64716, 64722, 64999, 64640	Not Covered (Experimental/Investigational/Unproven)
OSSIOfiber Trimmable Nail System/Compression Screws - Orthopedic Surgeries fixation devices for all indications.	No Specific Code L8699	Not Covered (Experimental/Investigation/Unproven) (May be included in surgical procedure)
Ovarian or Internal Iliac Vein Embolization for Treatment of Pelvic Congestion Syndrome	No Specific Code 37241	Not Covered (Experimental/Investigational/Unproven)
Patency Capsule Testing (e.g., PillCam Patency System, Agile Patency System/Capsule) for all indications	No specific code, 91299	Not Covered (Experimental/Investigational/Unproven)
Percutaneous ultrasonic ablation (e.g., Tenex percutaneous ultrasonic ablation procedures) - for all indications	No Specific Code 20999, 27599, 17999, 25999	Not Covered (Experimental/Investigational/Unproven) (May be coded depending on the area of treatment)

PROCEDURE	СРТ	COVERAGE STATUS
Percutaneous electrical nerve stimulation (PENS)/percutaneous neuromodulation therapy (PNT) all indications (e.g., chronic musculoskeletal or neuropathic	No specific code E1399, 64999	Not Covered (Experimental/Investigational/Unproven)
pain) Percutaneous transluminal	92972	Not Covered
coronary lithotripsy (List separately in addition to code for primary procedure)		(Experimental/Investigational/Unproven)
PortableConnect/ ROMTech (ROMTech Portable Connect Rehab Adaptive Device) (e.g., postsurgical rehabilitative exercises) for all indications	No Specific Code E1399	Not Covered (Experimental/Investigational/Unproven)
Prescription Digital Therapeutics (PDT) (prescription-only software to manage medical disorders or diseases)	No Specific Code 99199, E1399, T1505, A9291, A9999, A9292	Not Covered (Experimental/Investigational/Unproven)
Pudendal Nerve Blocks (Injections) ( e.g., Chronic Pelvic Pain, Pudendal neuralgia (PN))	64430	Not Covered (Experimental/Investigation/Unproven)  Note: Does not apply to the use of pudendal nerve blocks in obstetrics and other operative pelvic procedures;
Pulsed electromagnetic stimulation (PEMF)/SoftPulse Targeted Electromagnetic field therapy (tPEMF) (e.g., OthroCor Active Knee System, SoftPulse Device) for all indications	E0761	Not Covered (Experimental/Investigation/Unproven)
Quest AD-Detect Test - Alzheimer's Testing	0346U	Not Covered (Experimental/Investigational/Unproven)
Radiofrequency Intradiscal Biacuplasty (IBD) (e.g., Bialys TransDiscal System or Biacuplasty) (referred to as (TIPs) for all indications	No Specific Code 22899	Not Covered (Experimental/Investigational/Unproven)
Radiofrequency Nasal Valve w/VivAer Device – Treatment of nasal airway obstruction	30469	Not Covered (Experimental/Investigational/Unproven)

PROCEDURE	CPT HCPCS	COVERAGE STATUS
Reverse Axillary Lymphatic Mapping - all indications	No Specific Code 38999, C9756, 92240	Not Covered (Experimental/Investigational/Unproven)
RhinAer Procedure (Aerin Medical) (posterior nasal nerve ablation using radiofrequency) for Treatment of Chronic Rhinitis	No Specific Code 31242 30999	Not Covered (Experimental/Investigational/Unproven)
Robotic Assisted In-Home Therapy Device/ Systems (e.g., Motus Hand/Foot Robotic System, IpsilHand) for all indications	E0739; E0738	Not Covered (Experimental/Investigational/Unproven)
Selective Myectomy for all indications	No Specific Code 21499	Not Covered (Experimental/Investigational/Unproven)
ShuntCheck® to detect fluid flow in CSF shunt	62252	Not Covered (Experimental/Investigational/Unproven)
Sphenopalatine Ganglion Nerve Block for any indication including, but may not be limited to, headaches or trigeminal neuralgia.	No Specific Code 64505	Not Covered (Experimental/Investigational/Unproven)
Subacromial Balloon Spacer Implantation (InSpace) Rotator cuff repair or any other indication	No Specific Code 29999, C9781	Not Covered (Experimental/Investigational/Unproven)
Supraorbital Nerve Block Injection w/Ultrasound (SON) - for all indications (e.g., Migraine Headaches suboccipital neuralgia)	No Specific Code 64400 64999	Not Covered (Experimental/Investigational/Unproven)
Thermal Capsular Shrinkage Therapy (e.g.; Thermal capsulorrhaphy or Electrothermal Shrinkage) for all orthopedic indications	No Specific Code 29999, S2300, 25320	Not Covered (Experimental/Investigational/Unproven)
TissueCypher Barrett's Esophagus Assay (e.g., diagnosis and management of Barrett's esophagus and other esophageal disorders)	0108U	Not Covered (Experimental/Investigational/Unproven)
Trabecular Bone Score (TBS) Bone Mineral Density Measurement) to Predict Fracture Risk in Postmenopausal Patients	77089, 77090, 77091, 77092	Not Covered (Experimental/Investigational/Unproven)

PROCEDURE	CPT HCPCS	COVERAGE STATUS
Transcutaneous Vagal Nerve Stimulator (t-VNS) and Vagus Nerve Stimulation (VNS non- implantable) (e.g., depression/Bipolar/psychological disorders and all other indications)	E0735	Not Covered (Experimental/Investigational/Unproven)
Vagus Nerve Stimulation (Non- implantable, noninvasive tVNS, gammaCore Sapphire Device) for cluster and migraine headaches	E0735	Not Covered (Experimental/Investigation/Unproven)
Vascular Lymph Node Transfer (VLNT) also called lymph node transfer (LNT); Lymphatic By-pass Procedure) for all indications	No Specific Code 38999, 38308	Not Covered (Experimental/Investigational/Unproven)
Vertebral axial decompression (e.g., Lordex; VAX-D; DRX, and DRS System), (mechanized spinal distraction therapy or non-surgical traction device) for back pain	S9090	Not Covered (Experimental/Investigational/Unproven)
Virtual Reality Cognitive Behavioral Health therapy device	E1905	Not Covered (Experimental/Investigational/Unproven) (All "T" code related to Virtual Reality denied per Category III Policy)
Whole Body Vibration Platform (Galileo Plate) for all indications	No Specific Code E1399	Not Covered (Experimental/Investigational/Unproven)
Wireless GI Motility Monitoring Capsule Testing (e.g., SmartPill Mobility Testing System) – for all indications	91112	Not Covered (Experimental/Investigational/Unproven)

<sup>\*</sup> HCPCS codes, descriptions and materials are copyrighted by Centers for Medicare and Medicaid Services (CMS).

## **Related Policies**

Category III Current Procedural Terminology (CPT) Codes

Medical Necessity Reviews

## **Appendix**

**Policy Number:** 

**Effective:** 1/1/2021 **Next review:** 2/1/2025

<sup>\*</sup> CPT® codes, descriptions and materials are copyrighted by the American Medical Association (AMA).

Policy type: Enterprise

Author(s):

Depts.: Health Services

Applicable regulation(s): NC Social Security Act Section §1862 (a), 42 CFR 411.15(o), NCQA UM 10(A)(B) — Evaluation of New Technology; Guideline Notes 172 and 173 of the OHP Prioritized List of Health Services for guidance on New and Emerging Technology; WAC 284-44-043, WAC 284-46-507, IDS 41-3930, 41-5903, MCA 33-32-103, ARM 37.82.102(c)QA UM 10(A)(B) — Evaluation of New Technology; Guideline Notes 172 and 173 of the OHP Prioritized List of Health Services for guidance on New and Emerging Technology; WAC 284-44-043, WAC 284-46-507, IDS 41-3930, 41-5903, MCA 33-32-1003, ARM 37.82.102(c)

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