

MEDICAL POLICY - 2.01.535

Temporomandibular Joint Disorder

BCBSA Ref. Policy: 2.01.21, 5.01.05

Effective Date: May 1, 2025 Last Revised: Apr. 7, 2025

Replaces: 2.01.21

RELATED MEDICAL POLICIES:

2.01.26 Prolotherapy

2.01.31 Intra-Articular Hyaluronan Injections for Osteoarthritis

5.01.512 Botulinum Toxins

7.01.588 Percutaneous Electrical Nerve Stimulation and Percutaneous

Neuromodulation Therapy

9.02.501 Orthognathic Surgery

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POLICY CRITERIA | DOCUMENTATION REQUIREMENTS | CODING RELATED INFORMATION | EVIDENCE REVIEW | REFERENCES | HISTORY

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Introduction

The temporomandibular joint (TMJ) is the joint where the jawbone connects to the skull. There is one joint on each side of the jaw. The areas of the bones forming the joint are covered with cartilage and separated by a small disk. This disk helps keep joint movement smooth.

Sometimes the disc erodes or moves out of its proper position. Arthritis may develop in the joint and damage the cartilage, or an injury can damage the joint. Regardless of the cause, TMJ disorders (TMJD) can result in pain and affect the function of the joint and the muscles that control jaw movement. TMJDs may go away without treatment, or pain relievers can be used to alleviate symptoms. This policy describes the services that the health plan covers (considers medically necessary) to diagnose and treat TMJ symptoms and disorders. On some plans, services to treat TMJ problems are limited to a specific benefit which may have a dollar limit.

Note: The Introduction section is for your general knowledge and is not to be taken as policy coverage criteria. The rest of the policy uses specific words and concepts familiar to medical professionals. It is intended for providers. A provider can be a person, such as a doctor, nurse, psychologist, or dentist. A provider also can be a place where medical care is given, like a hospital, clinic, or lab. This policy informs them about when a service may be covered.

Policy Coverage Criteria

Note: Some health plan contracts may not have benefits to cover treatment of temporomandibular joint disorder. Refer to member contract language for benefit determination.

Treatment	Medical Necessity
Diagnostic procedures	 The following diagnostic procedures may be considered medically necessary in the diagnosis of temporomandibular joint disorder (TMJD): Diagnostic x-ray, tomograms, and arthrograms Computed tomography (CT) scan or magnetic resonance imaging (MRI) (in general, CT scans and MRIs are reserved for presurgical evaluations) Cephalograms (x-rays of jaws and skull) Pantograms (x-rays of maxilla and mandible) Note: Cephalograms and pantograms should be reviewed on an individual basis.
Surgical treatments	 The following surgical treatments may be considered medically necessary in the treatment of TMJD: Arthrocentesis Manipulation for reduction of fracture or dislocation of the TMJ Arthroscopic surgery in individuals with objectively demonstrated (by physical examination or imaging) internal derangements (displaced discs) or degenerative joint disease who have failed conservative treatment Open surgical procedures (when TMJD results from congenital anomalies, trauma, or disease in individuals who have failed conservative treatment) including, but not limited to: Arthroplasties Condylectomies Meniscus or disc plication Disc removal
Nonsurgical treatments	The following nonsurgical treatments may be considered medically necessary in the treatment of TMJD:

Treatment	Medical Necessity
	 One intraoral removable prosthetic device or appliance (encompassing fabrication, insertion, adjustment) Pharmacologic treatment (e.g., anti-inflammatory, muscle relaxing, analgesic medications) Physical therapy* (e.g., CPT 97530, 97014) Trigger point injections
	*Note: These services process to the Temporomandibular Joint (TMJ) Disorder benefit when the member has this benefit.
Two-piece oral appliance	A two-piece intraoral prosthetic device or appliance is considered not medically necessary.

Treatment	Investigational
Diagnostic procedures	 The following diagnostic procedures are considered investigational in the diagnosis of TMJD: Arthroscopy of the TMJ for purely diagnostic purposes Computerized mandibular scan (measures and records muscle activity related to movement and positioning of the mandible and is intended to detect deviations in occlusion and muscle spasms related to TMJD) Electromyography (EMG), including surface EMG Joint vibration analysis Kinesiography Muscle testing Neuromuscular junction testing Range-of-motion measurements Somatosensory testing Standard dental radiographic procedures Thermography Transcranial or lateral skull x-rays; intraoral tracing or gnathic arch tracing (intended to demonstrate deviations in the positioning of the jaw that are associated with TMJD) Ultrasound imaging/sonogram
Nonsurgical treatments	The following nonsurgical treatments are considered investigational in the treatment of TMJD:

Treatment	Investigational
	Biofeedback
	Botulinum toxin A
	Dental restorations/prostheses
	Devices promoted to maintain joint range of motion and to
	develop muscles involved in jaw function
	Dextrose prolotherapy
	Electrogalvanic stimulation
	Hyaluronic acid
	 Iontophoresis
	Orthodontic services
	Percutaneous electrical nerve stimulation (PENS)
	Platelet concentrates (e.g., platelet rich plasma)
	Transcutaneous electrical nerve stimulation (TENS) (e.g., HCPCS)
	E0720)
	Ultrasound

Documentation Requirements

Submit chart notes including type of appliance and history of re-occurring TMJ disorder.

Coding

Code	Description
СРТ	
20605	Arthrocentesis, aspiration and/or injection; intermediate joint or bursa (e.g., temporomandibular, acromioclavicular, wrist, elbow or ankle, olecranon bursa)
20606	Arthrocentesis, aspiration and/or injection, intermediate joint or bursa (e.g., temporomandibular, acromioclavicular, wrist, elbow or ankle, olecranon bursa); with ultrasound guidance, with permanent recording and reporting
21010	Arthrotomy, temporomandibular joint
21050	Condylectomy, temporomandibular joint
21060	Menisectomy, partial/complete, temporomandibular joint (separate procedure)



Code	Description
21073	Manipulation of temporomandibular joint(s) (TMJ), therapeutic, requiring an
	anesthesia service (i.e., general or monitored anesthesia care)
21085	Impression and custom preparation; oral surgical splint
21089	Unlisted maxillofacial prosthetic procedure
21116	Injection procedure for temporomandibular joint arthrography
21240	Arthroplasty, temporomandibular joint, with or without autograft (includes obtaining graft)
21242	Arthroplasty, temporomandibular joint, with allograft
21243	Arthroplasty, temporomandibular joint, with prosthetic joint replacement
21480	Closed treatment of temporomandibular dislocation; initial or subsequent
21485	Closed treatment of temporomandibular dislocation; complicated (e.g., recurrent
	requiring intermaxillary fixation or splinting), initial or subsequent
21490	Open treatment of temporomandibular dislocation
29800	Arthroscopy, temporomandibular joint, diagnostic, with or without synovial biopsy
	(separate procedure)
29804	Arthroscopy, temporomandibular joint, surgical
70328	Radiologic exam, temporomandibular joint, open and closed mouth; unilateral
70330	Radiologic examination, temporomandibular joint, open and closed mouth; bilateral
70332	Temporomandibular joint arthrography, radiological supervision and interpretation
70350	Cephalogram, orthodontic
70355	Orthopantogram (e.g., panoramic x-ray)
HCPCS	
J0585	Injection, onabotulinumtoxinA, (Botox) 1 unit
J0586	Injection, abobotulinumtoxinA, (Dysport) 5 units
J0587	Injection, rimabotulinumtoxinB, (Myobloc) 100 units
J0588	Injection, incobotulinumtoxinA, (Xeomin) 1 unit
J7321	Hyaluronan or derivative, Hyalgan or Supartz, for intra-articular injection, per dose
J7323	Hyaluronan or derivative, Euflexxa, for intra-articular injection, per dose
J7324	Hyaluronan or derivative, Orthovisc, for intra-articular injection, per dose
l	



Code	Description	
J7325	Hyaluronan or derivative, Synvisc or Synvisc-One, for intra-articular injection, 1 mg	
J7326	Hyaluronan or derivative, Gel-One, for intra-articular injection, per dose	
J7327	Hyaluronan or derivative, Monovisc, for intra-articular injection, per dose	
J7328	Hyaluronan or derivative, GELSYN-3, for intra-articular injection, 0.1 mg	
J7329	Hyaluronan or derivative, Trivisc, for intra-articular injection, 1 mg	
J7331	Hyaluronan or derivative, SYNOJOYNT, for intra-articular injection, 1 mg	
J7332	Hyaluronan or derivative, Triluron, for intra-articular injection, 1 mg	
S3900	Surface electromyography (EMG)	
CDT		
D7880	Occlusal orthotic device, by report	
D7881	Occlusal orthotic device adjustment	
D7899	Unspecified TMD therapy, by report	
D7999	Unspecified oral surgery procedure, by report	

Note: CPT codes, descriptions and materials are copyrighted by the American Medical Association (AMA). HCPCS codes, descriptions and materials are copyrighted by Centers for Medicare Services (CMS). CDT codes, descriptions and materials are copyrighted by the American Dental Association (ADA).

Related Information

N/A

Evidence Review

Description

Temporomandibular joint disorder (TMJD) refers to a group of disorders characterized by pain in the temporomandibular joint and surrounding tissues. Initial conservative therapy is generally



recommended; there are also a variety of nonsurgical and surgical treatment possibilities for individuals whose symptoms persist.

Background

Diagnosis of Temporomandibular Joint Disorder

In the clinical setting, temporomandibular joint disorder (TMJD) is often a diagnosis of exclusion and involves physical examination, patient interview, and a review of dental records. Diagnostic testing and radiologic imaging are generally only recommended for individuals with severe and chronic symptoms. Diagnostic criteria for TMJD have been developed and validated for use in both clinical and research settings.^{1,2,3}

Symptoms attributed to TMJD vary and include, but are not limited to, clicking sounds in the jaw; headaches; closing or locking of the jaw due to muscle spasms (trismus) or displaced disc; pain in the ears, neck, arms, and spine; tinnitus; and bruxism (clenching or grinding of the teeth).

Treatment

For many individuals, symptoms of TMJD are short-term and self-limiting. Conservative treatments, (e.g., eating soft foods, rest, heat, ice, avoiding extreme jaw movements) and anti-inflammatory medication, are recommended before considering more invasive and/or permanent therapies (e.g., surgery).

Botulinum Toxin

Chen et al (2015) summarized the evidence assessing the efficacy of botulinum toxin A for treatment of temporomandibular joint disorders in a systematic review that included five randomized controlled trials (RCTs).⁵⁵ Sample size in all trials was 30 or less except for one study. Three of the five studies were judged to be at high-risk of bias. All studies administered a single injection of onabotulinumtoxinA or abobotulinumtoxinA and followed up on the individuals at least one month later. Four studies used a placebo (normal saline) control group and the fifth compared abobotulinumtoxinA to fascial manipulation. Data was not pooled due to heterogeneity among the trials. In a qualitative review of the studies, two of the five trials found a significant short-term (1-2 months) benefit of onabotulinumtoxinA compared with control on pain reduction.



Summary of Evidence

For individuals with suspected TMJD who receive ultrasound, surface electromyography, or joint vibration analysis, the evidence includes systematic reviews of diagnostic test studies. Relevant outcomes are test validity, and other performance measures. None of the systematic reviews found that these diagnostic techniques accurately identified individuals with TMJD and many of the studies had methodologic limitations. The evidence is insufficient to determine that the technology results in an improvement in the net health outcome.

For individuals with a confirmed diagnosis of TMJD who receive intraoral devices or appliances or pharmacologic treatment, the evidence includes randomized controlled trials (RCTs) and systematic reviews of the RCTs. Relevant outcomes are symptoms, functional outcomes, quality of life, and treatment-related morbidity. A systematic review of intraoral appliances (44 studies) and meta-analyses of subsets of these studies found a significant benefit of intraoral appliances compared with control interventions. Several studies, meta-analyses, and systematic reviews exploring the effectiveness of stabilization splints on TMJD pain revealed conflicting results. Overall, the evidence shows that stabilizing splints may improve pain and positively impact depressive and anxiety symptoms. The evidence related to pharmacologic treatment varies because studies, systematic reviews, and meta-analyses lack consistency in evaluating specific agents. Some systematic reviews have found a significant benefit of several pharmacologic treatments (e.g., analgesics, muscle relaxants, and anti-inflammatory medications [vs. placebo]), but other studies showed a lack of benefit with agents such as methylprednisolone and botulinum toxin type A. The evidence is sufficient to determine that the technology results in an improvement in the net health outcome.

For individuals with a confirmed diagnosis of TMJD who receive acupuncture, biofeedback, transcutaneous electrical nerve stimulation (TENS), orthodontic services, hyaluronic acid, platelet concentrates, or dextrose prolotherapy, the evidence includes RCTs, systematic reviews of these RCTs, and observational studies. Relevant outcomes are symptoms, functional outcomes, quality of life, and treatment-related morbidity. Systematic reviews evaluating acupuncture for TMJD have found inconsistent improvement in outcomes compared with sham or active controls. A 2023 meta-analysis of 22 RCTs failed to find improved pain or maximum mouth opening with acupuncture compared with active controls. Systematic reviews evaluating hyaluronic acid have found similar outcomes to corticosteroids or placebo. Platelet-rich plasma has been compared with hyaluronic acid in a number of systematic reviews and RCTs, but the studies are small and have methodologic limitations. The evidence is insufficient to determine that the technology results in an improvement in the net health outcome.



For individuals with a confirmed diagnosis of TMJD, who receive arthrocentesis or arthroscopy, the evidence includes RCTs, systematic reviews of RCTs, and observational studies. Relevant outcomes are symptoms, functional outcomes, quality of life, and treatment-related morbidity. One review, which included three RCTs, compared arthrocentesis or arthroscopy with nonsurgical interventions for TMJD. Pooled analyses of the RCTs found that arthrocentesis and arthroscopy resulted in superior pain reduction compared with control interventions. A network meta-analysis, which included 36 RCTs, revealed that arthroscopy and arthrocentesis improve pain control and maximum mouth opening. Two recent meta-analyses identified RCTs comparing arthrocentesis to various conservative management strategies. At six months, one analysis found improved maximum mouth opening with arthrocentesis while the other found similar outcomes between arthrocentesis and conservative treatments. Similarly, pain was improved with arthrocentesis in one analysis, but not the other. The evidence is sufficient to determine that the technology results in an improvement in the net health outcome.

For individuals who receive botulinum toxin injections for treatment of temporomandibular joint disorders, the evidence includes case series and RCTs. The relevant outcomes are symptoms, functional outcomes, medication use, and treatment-related morbidity. Generally, botulinum toxin has been evaluated in clinical settings where individuals have failed the standard of care or in whom standard of care interventions are contraindicated. However, studies using a placebo comparator that lack scientific rigor do not permit conclusions about the net health benefit of botulinum toxin. Future studies in this clinical indication should use appropriate comparators in adequately powered prospective studies using a standardized treatment dose and adequate follow-up. The evidence is insufficient to determine that the technology results in an improvement in the net health outcome.

Ongoing and Unpublished Clinical Trials

Some currently ongoing and unpublished trials that might influence this review are listed in **Table 1**.

Table 1. Summary of Key Trials

NCT No.	Trial Name	Planned Enrollment	Completion Date
Ongoing			

NCT No.	Trial Name	Planned Enrollment	Completion Date
NCT06573502	Assessment of Platelet Rich Plasma With Injectable Platelet Rich Fibrin VS Platelet Rich Plasma Vs Platelet Rich Fibrin in Management of Temporomandibular Joint Osteoarthritis	45	Jul 2025
NCT06457698	Temporomandibular Bioviscosupplementation (Platelet-Rich Plasma Combined With Hyaluronic Acid) After Double- Puncture Arthrocentesis: a Randomized Controlled Trial	50 (actual)	Dec 2024
NCT06530745	Quintuple Intra-Articular Hyaluronic Acid (HA) Improves and Platelet-Rich Plasma (PRP) Does Not Affect Mandibular Mobility in TemporomandibularJoint (TMJ) Disorders: A Controlled Trial	78	Jun 2025
NCT05989217	Conservative Therapies in the Treatment of Temporomandibular Disorders: a Randomized Controlled Clinical Trial	96	Sep 2024
NCT04884763 ^a	A Randomized, Double Blind, Placebo-Controlled Single Center Phase 2 Pilot Study to Assess the Safety and Efficacy of Off-label Subcutaneous Administration of Erenumab- aooe in Patients With Temporomandibular Disorder	30 (actual)	Jan 2024
NCT04726683	Trigger Point Dry Needling vs Injection in Patients With Temporomandibular Disorders: a Randomized Placebo- controlled Trial	64	Dec 2024
Unpublished			
NCT04936945	Comparative Study Between the Outcome of Intra-articular Injection of Platelet Rich Plasma Versus Hyaluronic Acid in Arthroscopic Management of Temporomandibular Degenerative Joint Diseases: A Randomized Clinical Trial	20	Jun 2023
NCT04298554	Comparison of Cannabinoids to Placebo in Management of Arthralgia and Myofascial Pain Disorder of the Temporomandibular Region: A Randomized Clinical Trial.	59	May 2022
NCT05027243	Outcomes of Bilateral Temporomandibular Joint Arthroscopy and the Role of a Second Intervention - Timings and Results	46	July 2021

NCT: national clinical trial. ^aDenotes industry sponsored or co-sponsored trial

Practice Guidelines and Position Statements

The purpose of the following information is to provide reference material. Inclusion does not imply endorsement or alignment with the policy conclusions.



Guidelines or position statements will be considered for inclusion if they were issued by, or jointly by, a US professional society, an international society with US representation, or National Institute for Health and Care Excellence (NICE). Priority will be given to guidelines that are informed by a systematic review, include strength of evidence ratings, and include a description of management of conflict of interest.

American Association for Dental, Oral and Craniofacial Research

In 2010 (reaffirmed in 2015), the American Association for Dental Research (now the American Association for Dental, Oral, and Craniofacial Research) policy statement recommended the following for the diagnosis and treatment of temporomandibular joint disorders (TMJDs)⁵³:

It is recommended that the differential diagnosis of TMDs [temporomandibular disorders] or related orofacial pain conditions should be based primarily on information obtained from the patient's history, clinical examination, and when indicated, TMJ [temporomandibular joint] radiology or other imaging procedures. The choice of adjunctive diagnostic procedures should be based upon published, peer-reviewed data showing diagnostic efficacy and safety. However, the consensus of recent scientific literature about currently available technological diagnostic devices for TMDs is that except for various imaging modalities, none of them shows the sensitivity and specificity required to separate normal subjects from TMD patients or to distinguish among TMD subgroups...

It is strongly recommended that, unless there are specific and justifiable indications to the contrary, treatment of TMD individuals initially should be based on the use of conservative, reversible and evidence-based therapeutic modalities. Studies of the natural history of many TMDs suggest that they tend to improve or resolve over time. While no specific therapies have been proven to be uniformly effective, many of the conservative modalities have proven to be at least as effective in providing symptomatic relief as most forms of invasive treatment...

American Society of Temporomandibular Joint Surgeons

In 2001, the American Society of Temporomandibular Joint Surgeons issued consensus clinical guidelines focused on TMJDs associated with internal derangement and osteoarthritis.⁵⁴ For diagnosis of this type of TMJD, a detailed history and, when indicated, a general physical examination was recommended. Imaging of the temporomandibular and associated structures was also recommended. Options for basic radiography to provide information on temporal bone



and condylar morphology included the use of plain films, panoramic films, and tomograms. Also recommended was imaging of the disc and associated soft tissue with magnetic resonance imaging (MRI) or arthrography. Other diagnostic procedures indicated included computed tomography, MRI, arthrography (for selected cases) and isotope bone scans.

Nonsurgical treatment was recommended as first-line therapy for all symptomatic patients with this condition. Recommended treatment options include a change in diet, nonsteroidal anti-inflammatory drugs, maxillomandibular appliances, physical therapy, injections of corticosteroids or botulinum toxin, and behavior modification. If adequate symptom relief did not occur within two to three weeks, surgical consultation was advised. The guideline stated the following surgical procedures were considered acceptable and effective for individuals with TMJDs associated with internal derangement or osteoarthritis:

- Arthrocentesis
- Arthroscopy
- Condylotomy
- Arthrotomy (prosthetic joint replacement may be indicated in selected individuals who have severe joint degeneration, destruction, or ankylosis)
- Coronoidotomy/coronoidectomy
- Styloidectomy

BMJ Rapid Recommendations

The BMJ Rapid Recommendations panel developed guidelines for the management of patients with chronic pain (≥3 months) associated with TMJD.⁵⁵ The international expert panel included representation from an academic center in the United States.

The panel favored the following therapies:

- Cognitive behavior therapy (strong recommendation)
- Therapist-assisted mobilization (strong recommendation)
- Manual trigger point therapy (strong recommendation)
- Supervised postural or jaw exercise (strong recommendation)



- Usual care including home exercises, stretching, reassurance, and education (strong recommendation)
- Manipulation (conditional recommendation)
- Supervised jaw exercise with mobilization (conditional recommendation)
- Cognitive behavior therapy with non-steroidal anti-inflammatory drugs (conditional recommendation)
- Manipulation with postural exercise (conditional recommendation)
- Acupuncture (conditional recommendation)

The panel recommended against the following therapies:

- Reversible occlusal splints (conditional recommendation)
- Arthrocentesis (conditional recommendation)
- Cartilage supplement with or without hyaluronic acid injection (conditional recommendation)
- Low level laser therapy (conditional recommendation)
- Transcutaneous electrical nerve stimulation (conditional recommendation)
- Gabapentin (conditional recommendation)
- Botulinum toxin (conditional recommendation)
- Hyaluronic acid (conditional recommendation)
- Relaxation therapy (conditional recommendation)
- Trigger point injection (conditional recommendation)
- Acetaminophen (conditional recommendation)
- Topical capsaicin (conditional recommendation)
- Biofeedback (conditional recommendation)
- Corticosteroid injection (conditional recommendation)

- Benzodiazepines (conditional recommendation)
- Beta-blockers (conditional recommendation)
- Irreversible oral splints (strong recommendation)
- Discectomy (strong recommendation)
- Non-steroidal anti-inflammatory drugs with opioids (strong recommendation)

Medicare National Coverage

There is no national coverage determination.

Regulatory Status

Since 1981, several muscle-monitoring devices have been cleared for marketing by the US Food and Drug Administration (FDA) through the 510(k) process. Some examples are:

- The BioEMG III (Bio-Research Associates)
- The GrindCare Measure (Medotech A/S)
- The K7x Evaluation System (Myotronics)
- M-Scan (Bio-Research Associates)

These devices aid clinicians in the analysis of joint sound, vibrations, and muscle contractions when diagnosing and evaluating TMJD.

FDA Product Code: KZM.

Table 2. Muscle-Monitoring Devices Cleared by the US Food and Drug Administration

Devices	Manufacturer	Date Cleared	510(k) No.	Indication
K7x Evaluation System	Myotronics, Inc	Nov 2000	K003287	Electromyography

Devices	Manufacturer	Date Cleared	510(k) No.	Indication
BioEMG IIITM	Bio-Research Associates, Inc	Feb 2009	K082927	Electromyography, Joint Vibration Recording
GrindCare Measure	Medotech A/S	Apr 2012	K113677	Electromyography, Nocturnal Bruxism
M-Scan	Bio-Research Associates	Jul 2013	K130158	Electromyography
TEETHAN 2.0	BTS S.P.A.	Dec 2016	K161716	Electromyography
GrindCare System	Sunstar Suisse S.A.	Sep 2017	K163448	Electromyography, Sleep Bruxism
Nox Sleep System	Nox Medical	Nov 2019	K192469	Electromyography, Sleep Bruxism

FDA product code: KZM

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History

Date	Comments
05/01/19	New policy, approved April 18, 2019. This policy replaces 2.01.21. Policy created with literature through December 2018. Policy statement added that two-piece intraoral prosthetic device or appliance is considered not medically necessary.
06/07/19	Interim Review, approved June 4, 2019, effective June 7, 2019. Added physical therapy and trigger point injections to medically necessary non-surgical treatments.
01/01/20	Coding update, removed CDT code D9940 as it terminated 1/1/19.
05/01/20	Annual Review, approved April 7, 2020. Policy updated with literature review through December 2019; references added. Policy statements unchanged. Added CPT code 20606, J7327, J7328, J7329, J7331 and J7332.
07/01/20	Coding update. Added code J7333.



Date	Comments
05/01/21	Annual Review, approved April 13, 2021. Policy updated with literature review through January 8, 2021; references added. Investigational policy statement modified to include platelet concentrates. Added term date to HCPC code J7333.
05/01/22	Annual Review, approved April 12, 2022. Policy updated with literature review through December 20, 2021; references added. Dextrose prolotherapy added to investigational policy statement. Removed HCPCS code J7333.
11/01/22	Coding update. Added HCPCS codes J0585, J0586, J0587 and J0588. Revised coding descriptions on CDT codes D7880 & D7999.
05/01/23	Annual Review, approved April 10, 2023. Policy updated with literature review through December 19, 2022; references added. Minor editorial refinements to policy statements; intent unchanged. Changed the wording from "patient" to "individual" throughout the policy for standardization.
05/01/24	Annual Review, approved April 8, 2024. Policy updated with literature review through December 13, 2023; references added. Policy statements unchanged.
05/01/25	Annual Review, approved April 7, 2025. Policy updated with literature review through December 10, 2024; reference added. Policy statements unchanged

Disclaimer: This medical policy is a guide in evaluating the medical necessity of a particular service or treatment. The Company adopts policies after careful review of published peer-reviewed scientific literature, national guidelines and local standards of practice. Since medical technology is constantly changing, the Company reserves the right to review and update policies as appropriate. Member contracts differ in their benefits. Always consult the member benefit booklet or contact a member service representative to determine coverage for a specific medical service or supply. CPT codes, descriptions and materials are copyrighted by the American Medical Association (AMA). ©2025 Premera All Rights Reserved.

Scope: Medical policies are systematically developed guidelines that serve as a resource for Company staff when determining coverage for specific medical procedures, drugs or devices. Coverage for medical services is subject to the limits and conditions of the member benefit plan. Members and their providers should consult the member benefit booklet or contact a customer service representative to determine whether there are any benefit limitations applicable to this service or supply. This medical policy does not apply to Medicare Advantage.

