

Health Evidence Review Commission

May 19, 2022 1:30 PM - 4:30 PM

Online Meeting

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Section 1.0 Call to Order

AGENDA HEALTH EVIDENCE REVIEW COMMISSION

Online meeting

May 19, 2022 1:30-4:30 pm

(All agenda items are subject to change and times listed are approximate)

#	Time	ltem	Presenter	Action Item
1	1:30 PM	Call to order	Kevin Olson	
2	1:35 PM	Approval of minutes (3/10/2022)	Kevin Olson	Х
3	1:40 PM	Director's report	Jason Gingerich	
4	1:45 PM	Value-based Benefits Subcommittee report See <u>VbBS agenda</u> for specific topics	Ariel Smits	Х
6	2:45 PM	Coverage Guidance Topic: Bariatric Procedures (Review the research plan for an upcoming report on weight loss procedures) • Scope Statement	Ariel Smits	х
5	3:00 PM	Coverage Guidance Topic: PANDAS/PANS (OCD and other symptoms developed after a case of strep in children) • Coverage guidance • Prioritized List changes	Val King Ariel Smits	Х
8	4:25 PM	Next steps • Schedule next meeting – 8/11/2022 Virtual	Kevin Olson	
9	4:30 PM	Adjournment	Kevin Olson	

Note: Public comment will be taken on each topic per HERC policy at the time at which that topic is discussed.

MINUTES

HEALTH EVIDENCE REVIEW COMMISSION Online Meeting March 10, 2022

Members Present: Kevin Olson, MD, Chair; Holly Jo Hodges, MD, MBA, Vice-Chair; Devan Kansagara, MD; Adriane Irwin, PharmD, Kathryn Schabel, MD; Max Kaiser, DO; Deborah Espesete, LAc, MAcOM, MPH; Cris Pinzon, MPH, BSN, BS, RN; Stacy Geisler, DDS, PhD; Ben Hoffman, MD.

Members Absent: Leslie Sutton; Mike Collins; Lynnea Lindsey, PhD.

Staff present: Ariel Smits, MD, MPH; Jason Gingerich; Daphne Peck.

Also Attending: Valerie King MD MPH & Shauna Durbin, (OHSU Center for Evidence-based Policy); Alison Little, MD MPH; Amanda Trujillo; Catherine Sweeney; Dana Pursley-Haner (Sherman County); Gary Hansen (RespirTech); Kristty Zamora-Polanco (Oregon Health Authority); I walker; Lisa Kouzes, DC; Maria Gonzalez-Cress; Melanie Ewald; Miriam McDonell, MD; Obinna Oleribe; Siobhan Hess; Stephanie.

Call to Order

Kevin Olson, Chair of the Health Evidence Review Commission (HERC), called the meeting to order; roll was called. A quorum of members was present at the meeting. Each Commissioner gave a brief introduction.

Minutes Approval

MOTION: To approve the minutes of the 11/18/2021 meeting as presented. CARRIES 11-0.

Director's Report

Legislative session

Gingerich reported there were no changes that affected the Commission this short session.

Plain language summary

He pointed out a new section to the meeting materials and asked for feedback. Gingerich said during the Value-based Benefits Subcommittee (VbBS) report we will pause to reflect on those new statements as they come up.

Required trainings

Gingerich said there are required trainings for Commissioners to take between March 15 and December 31, 2022. An email will be delivered soon with details.

Early and Periodic Screening, Diagnostic and Treatment (EPSDT) waiver

He said that EPSDT waiver that has been in effect since the 1990s has been dropped from Oregon's 1115 waiver application based on public feedback. Other OHA staff are preparing for the change by looking at operational changes and HERC staff are reviewing the unfunded region to identify things that

should be considered for reprioritization. The implementation for this change is January 2024, though some changes have already been made to the List and other changes will be recommended in coming months.

Membership

- Regina Dehen resigned from VbBS. There is an opening for a naturopath. Recruitment will begin soon.
- Committee appointments
 - Evidence-based Guidelines Subcommittee (EbGS)
 - Appoint Dr. Ben Hoffman
 - Commissioner and a pediatrician
 - Appoint Dr. Miriam (Mimi) McDonnell
 - An obstetrician/gynecologist and a public health officer
 - Dr. Alison Little as Vice Chair, to be nominated and appointed at EbGS
 - Long serving member of the Commission; she will retire from the Commission at the year's end.
 - o Dr. Stacy Geisler will serve on the Oral Health Advisory Panel; no vote is required

MOTION: To Appoint Dr. Hoffman to EbGS. CARRIES: 11-0.

MOTION: To Appoint Dr. McDonnell to EbGS. CARRIES: 11-0.

Value-based Benefits Subcommittee (VbBS) Report on Prioritized List Changes Meeting materials pages 60-137

Ariel Smits reported the VbBS met earlier in the day, 3/10/2022. She summarized the subcommittee's recommendations.

RECOMMENDED CODE MOVEMENT (changes to the 10/1/2021 Prioritized List unless otherwise noted)

- Move the diagnosis code for inflammatory joint diseases associated with autoimmune gut disease from an unfunded to a funded line
- Add the procedure code for platelet rich plasma injections to an unfunded line
- Add a procedure code to allow minimally invasive ablation of small renal tumors to the funded renal cancer line
- Add the CPT codes for gait analysis and surface electromyography to an unfunded line
- Delete the diagnosis code for extra toes from an unfunded line and left only on a funded line
- Add the procedure code for dorsal rhizotomy to a funded line to pair with spastic cerebral palsy
- Make a variety of straightforward coding changes

ITEMS CONSIDERED BUT NO RECOMMENDATIONS FOR CHANGES MADE

- No change was made in the non-coverage of mid-foot fusion for foot arthritis
- No change was made in the non-coverage of treatment of actinic keratoses
- No change was made to the non-coverage of sensory integration therapy

RECOMMENDED GUIDELINE CHANGES (changes to the 10/1/2021 Prioritized List unless otherwise noted)

• Edit the chemodenervation guideline to include two additional lines with chemodenervation codes

- Add a new guideline indicating that pelvic congestion syndrome is a non-funded syndrome and does not pair with various vein procedures
- Edit the breast reconstruction after breast cancer surgery guideline to clarify that reconstruction is also covered after lumpectomy.
- Delete two guidelines regarding breast screening and extensively edit one guideline to indicate when breast MRI is a covered service
- Add a new guideline outlining when ablation of renal tumors is covered
- Edit the lower urinary tract symptoms guideline to clarify when procedures are covered
- Add a new guideline regarding dorsal rhizotomy
- Make several straightforward guideline note changes

2024 Biennial Review

- Delete the agenesis of lung line effective 1/1/2024
- Delete the spastic diplegia line effective 1/1/2024

MOTION: To accept the VbBS recommendations on *Prioritized List changes* as stated. See the VbBS minutes of 3/10/2022 for a full description. Carries: 11-0.

Coverage Guidance Topic: High-Frequency Chest Wall Oscillation Devices Meeting materials handout, pages 2-43

Gingerich said a team member discovered a significant error in the coverage guidance during preparations for today's meeting. The error resulted in a key piece of evidence having its confidence level downgraded.

Valerie King, MD MPH said the evidence in question was a confusing Cochrane review where most of the evidence for cystic fibrosis was found. In that review, there was one randomized control trial (RCT) where the Cochrane Review authors had obtained some unpublished data from the authors of one included RCT. A non-eligible comparator was mistakenly used. It was on this issue of hospitalizations that took the confidence of evidence from being low to very low. This could possibly change the conclusions about cystic fibrosis and may influence extrapolations of that evidence to bronchiectasis.

Hoffman said chest physical therapy (PT) is hard to do and is time consuming.

Gingerich said the Commission could choose to send the edited coverage guidance out to public comment or return it to the subcommittee for further review and study.

MOTION: To return the coverage guidance to EbGS for further review. Carries 11-0.

Public Comment

There was no public comment.

Adjournment

Meeting adjourned at 3:30 pm. Next meeting will be from 1:30-4:30 pm on Thursday, May 19, 2022, by Zoom, online.



Value-based Benefits Subcommittee Recommendations Summary For Presentation to:

Health Evidence Review Commission on March 10, 2022

For specific coding recommendations and guideline wording, please see the text of the 3/10/2022 VbBS minutes.

RECOMMENDED CODE MOVEMENT (changes to the 10/1/2021 Prioritized List unless otherwise noted)

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- Add the procedure code for platelet rich plasma injections to an unfunded line
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- Make a variety of straightforward coding changes

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- Edit the breast reconstruction after breast cancer surgery guideline to clarify that reconstruction is also covered after lumpectomy.
- Delete two guidelines regarding breast screening and extensively edit one guideline to indicate when breast MRI is a covered service
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- Make several straightforward guideline note changes

2024 Biennial Review

- Delete the agenesis of lung line effective 1/1/2024
- Delete the spastic diplegia line effective 1/1/2024

VALUE-BASED BENEFITS SUBCOMMITTEE

Virtual Meeting March 10, 2022 8:00 AM – 1:00 PM

Members Present: Kevin Olson, MD, Chair; Holly Jo Hodges, MD, MBA, Vice-chair; Cris Pinzon, MPH, BSN, BS, RN; Brian Duty, MD; Adriane Irwin, PharmD; David Saenger, MD.

Members Absent: Kathryn Schabel, MD; Mike Collins.

Staff Present: Ariel Smits, MD, MPH; Jason Gingerich; Daphne Peck.

Also Attending: Dawn Mautner, MD; Kristty Zamora-Polanco and Senna Towner (Oregon Health Authority); Jenna Oh; I walker; Lisa Kouzes; Maria Gonzalez-Cress; Obinna Oleribe; Shauna Durbin and Val King MD MPH (Center for Evidence Based Health Policy); siobhan hess

> Roll Call/Minutes Approval/Staff Report

The meeting was called to order at 8:00 am and roll was called. A quorum of members was present at the meeting. Minutes from the November 18, 2021 VbBS meeting were reviewed and approved.

Gingerich gave an update on Early and Periodic Screening, Diagnostic and Treatment (EPSDT) changes that are anticipated to be put into place on 1/1/2024, as well as the impact of recent changes to Statement of Intent 4 around allowing otherwise nonfunded services to be covered if they would benefit a child in terms of growth, development or ability to participate in school. He mentioned a recent CMS letter with requirements that treatments related to "long COVID" should be covered when medically necessary even if they wouldn't otherwise be covered, which is in some ways similar to the EPSDT changes coming in 2024.

Gingerich made announcements of membership changes. He also introduced HERC staff trial of plain language summaries to certain issues summaries in today's meeting materials and asked for member and public feedback.

Smits reviewed the errata document, as well as the January 1, 2022 placement of newly ACIP-approved pneumococcal vaccine CPT codes on a funded line per expressed HERC intent.

> Topic: Straightforward/Consent Agenda

Discussion: There was discussion on the following items:

- 1) CPT 87913 (COVID genotyping). Olson asked whether there was a pressing reason to add this code to the Diagnostic Procedure File as it is not currently required for clinical care and is subject to misuse. Smits noted that the code could be added to the COVID line to only pair with COVID infection. Gingerich noted that there were federal rules regarding COVID testing that would need to be consulted if this test was not covered. The group agreed to the staff recommended placement on the Diagnostic Procedure File, but requested that staff periodically audit use and bring this information to the HERC for possible action if overused.
- 2) Newborn home visits: Gingerich noted that these services are a carve-out and do not have cost to the CCOs.

Recommended Actions:

- 1) Add M62.81 (Muscle weakness (generalized)) to the dysfunction lines 71,292,345 and 377
- 2) Remove N96 (Recurrent pregnancy loss) from line 658 GENITOURINARY CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY
 - a. Advise HSD to add N96 to the Diagnostic Workup File
- 3) Remove H02.73 family (Vitiligo of eyelid and periocular area) from line 654 SENSORY ORGAN CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY
 - a. Add H02.73 family to lines 426 SEVERE INFLAMMATORY SKIN DISEASE and 656 DERMATOLOGICAL CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY
- 4) Remove K22.10 (Ulcer of esophagus without bleeding) from line 513 ESOPHAGITIS AND GERD; ESOPHAGEAL SPASM; ASYMPTOMATIC DIAPHRAGMATIC HERNIA
 - a. Add K22.10 to line 56 ULCERS, GASTRITIS, DUODENITIS, AND GI HEMORRHAGE
- 5) Remove M35.00 (Sjogren syndrome, unspecified) from line 510 DYSFUNCTION OF NASOLACRIMAL SYSTEM IN ADULTS: LACRIMAL SYSTEM LACERATION
 - a. Add M35.00 to line 330 SYSTEMIC SCLEROSIS; SJOGREN'S SYNDROME
- 6) Remove L49.7 (Exfoliation due to erythematous condition involving 70-79 percent of body surface) from lines 57 SEVERE BURNS and 127 MODERATE BURNS
 - a. Add L49.7 to line 504 ERYTHEMATOUS CONDITIONS
- 7) Remove H70.1 (Chronic mastoiditis) and H70.9 families (Unspecified mastoiditis) from line 476 CHRONIC OTITIS MEDIA; OPEN WOUND OF EAR DRUM
 - a. Add H70.1 and H70.9 families to line 170 ACUTE MASTOIDITIS
- 8) Change the title of line 482 to MILD/MODERATE LICHEN PLANUS
- Remove D78.02 (Intraoperative hemorrhage and hematoma of the spleen complicating other procedure) from line 529 DISORDERS OF FUNCTION OF STOMACH AND OTHER FUNCTIONAL DIGESTIVE DISORDERS
 - a. Add D78.02 to line 285 COMPLICATIONS OF A PROCEDURE ALWAYS REQUIRING TREATMENT
- 10) Remove B33.2 family (Viral endocarditis, myocarditis, pericarditis, cardiomyopathy) from line 615 OTHER VIRAL INFECTIONS
 - a. Add B33.2 family to line 81 MYOCARDITIS, PERICARDITIS, AND ENDOCARDITIS
- 11) Remove H16.31 (Corneal abscess) family from line 473 KERATOCONJUNCTIVITIS
 - a. Add H16.31 family to line 244 CORNEAL ULCER; SUPERFICIAL INJURY OF EYE AND ADNEXA

- 12) Add HCPCS C9761 (Cystourethroscopy, with ureteroscopy and/or pyeloscopy, with lithotripsy, and ureteral catheterization for steerable vacuum aspiration of the kidney, collecting system, ureter, bladder, and urethra if applicable) to lines 49 CONGENITAL HYDRONEPHROSIS, 180 URETERAL STRICTURE OR OBSTRUCTION; HYDRONEPHROSIS; HYDROURETER, and 352 URINARY SYSTEM CALCULUS
- 13) Add 67515 (Injection of medication or other substance into Tenon's capsule) to lines 370 AMBLYOPIA and 393 STRABISMUS WITHOUT AMBLYOPIA AND OTHER DISORDERS OF BINOCULAR EYE MOVEMENTS; CONGENITAL ANOMALIES OF EYE; LACRIMAL DUCT OBSTRUCTION IN CHILDREN
- 14) Remove 17000 (Destruction (eg, laser surgery, electrosurgery, cryosurgery, chemosurgery, surgical curettement), premalignant lesions (eg, actinic keratoses); first lesion) from lines 373 ACNE CONGLOBATA AND ACNE FULMINANS, 453 SEVERE CYSTIC ACNE, 522 ROSACEA; MILD/MODERATE ACNE
- 15) Add N48.82 (Acquired torsion of penis) to line 424 COMPLICATIONS OF A PROCEDURE USUALLY REQUIRING TREATMENT
- 16) Modify GN73 as shown in Appendix A
- 17) Add the following CPT codes to line 3 PREVENTION SERVICES WITH EVIDENCE OF EFFECTIVENESS
 - a. 91308 Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]) vaccine, mRNA-LNP, spike protein, preservative free, 3 mcg/0.2 mL dosage, diluent reconstituted, tris-sucrose formulation, for intramuscular use
 - b. 0081A Immunization administration by intramuscular injection of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]) vaccine, mRNA-LNP, spike protein, preservative free, 3 mcg/0.2 mL dosage, diluent reconstituted, tris-sucrose formulation; first dose
 - c. 0081B Second dose
 - d. 91309 Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]) vaccine, mRNA-LNP, spike protein, preservative free, 50 mcg/0.5 mL dosage, for intramuscular use
 - e. 0094A Immunization administration by intramuscular injection of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]) vaccine, mRNA-LNP, spike protein, preservative free, 50 mcg/0.5 mL dosage, booster dose
- 18) Add CPT 87913 (Infectious agent genotype analysis by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]), mutation identification in targeted region(s)) to the Diagnostic Procedure File
- 19) Modify Diagnostic Guideline D27 as shown in Appendix A
- 20) Add CPT 99502 (Home visit for newborn care and assessment) to line 3 PREVENTION SERVICES WITH EVIDENCE OF EFFECTIVENESS
- 21) Add CPT 99501 (Home visit for postnatal assessment and follow-up care) to line 1 PREGNANCY
- 22) Add the S86.11 family (Strain of other muscle(s) and tendon(s) of posterior muscle group at lower leg level) to line 376 DISRUPTIONS OF THE LIGAMENTS AND TENDONS OF THE ARMS AND LEGS, EXCLUDING THE KNEE, RESULTING IN SIGNIFICANT INJURY/IMPAIRMENT
- 23) Add the S46.00 family (Unspecified injury of muscle(s) and tendon(s) of the rotator cuff of shoulder) to line 417 DISORDERS OF SHOULDER, INCLUDING SPRAINS/STRAINS GRADE 4 THROUGH 6 and 608 SPRAINS AND STRAINS OF ADJACENT MUSCLES AND JOINTS, MINOR
 - a. Remove the S46.00 family from line 634 SUPERFICIAL WOUNDS WITHOUT INFECTION AND CONTUSIONS

24) Add the S46.09 family (Other injury of muscle(s) and tendon(s) of the rotator cuff of shoulder), S46.19 family (Other injury of muscle, fascia and tendon of long head of biceps), S46.29 family (Other injury of muscle, fascia and tendon of other parts of biceps), S46.39 family (Other injury of muscle, fascia and tendon of triceps), S46.89 family (Other injury of other muscles, fascia and tendons at shoulder and upper arm level), and S46.99 family (Other injury of unspecified muscle, fascia and tendon at shoulder and upper arm level) to lines 376, 417, and 608 and remove from line 634

25) Table:

Code	Add to line	Delete from line
S56.00 family (Unspecified injury of flexor	376 DISRUPTIONS OF THE	634
muscle, fascia and tendon of right thumb at	LIGAMENTS AND TENDONS OF	SUPERFICIAL
forearm level)	THE ARMS AND LEGS,	WOUNDS
	EXCLUDING THE KNEE,	WITHOUT
	RESULTING IN SIGNIFICANT	INFECTION AND
	INJURY/IMPAIRMENT	CONTUSIONS
	608 SPRAINS AND STRAINS OF	
	ADJACENT MUSCLES AND JOINTS,	
	MINOR	
S56.09 family (Other injury of flexor muscle,	376	634
fascia and tendon of right thumb at forearm	608	
level)		
S56.19 family (Other injury of flexor muscle,	376	634
fascia and tendon of index finger at forearm	608	
level)		
S56.20 family (Unspecified injury of other	376	634
flexor muscle, fascia and tendon at forearm	608	
level)		
S56.29 family (Other injury of other flexor	376	634
muscle, fascia and tendon at forearm level)	608	
S56.39 family (Other injury of extensor or	376	634
abductor muscles, fascia and tendons of	608	
thumb at forearm level)		
S56.49 family (Other injury of extensor	376	634
muscle, fascia and tendon of middle finger at	608	
forearm level)		
S56.59 family (Other injury of other extensor	376	634
muscle, fascia and tendon at forearm level)	608	
S56.89 family (Other injury of other muscles,	376	634
fascia and tendons at forearm level)	608	
S66.00 family (Unspecified injury of long flexor	376	634
muscle, fascia and tendon of thumb at wrist	608	
and hand level)		
S66.09 family (Other specified injury of long	376	634
flexor muscle, fascia and tendon of thumb at	608	
wrist and hand level)		

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S66.10 family (Unspecified injury of flexor	376	634
muscle, fascia and tendon of index finger at	608	
wrist and hand level)		
S66.19 family (Other injury of flexor muscle,	376	634
fascia and tendon of index finger at wrist and	608	
hand level)		
S66.20 family (Unspecified injury of extensor	376	634
muscle, fascia and tendon of thumb at wrist	608	
and hand level)		
S66.29 family (Other specified injury of	376	634
extensor muscle, fascia and tendon of thumb	608	
at wrist and hand level)		
S66.30 family (Unspecified injury of extensor	376	634
muscle, fascia and tendon of other finger at	608	
wrist and hand level)		
S66.39 family (Other injury of extensor	376	634
muscle, fascia and tendon of index finger at	608	
wrist and hand level)		
S66.40 family (Unspecified injury of intrinsic	376	634
muscle, fascia and tendon of thumb at wrist	608	
and hand level)		
S66.49 family (Other specified injury of	376	634
intrinsic muscle, fascia and tendon of thumb	608	
at wrist and hand level)		
S66.50 family (Unspecified injury of intrinsic	376	634
muscle, fascia and tendon of index finger at	608	
wrist and hand level)		
S66.59 family (Other injury of intrinsic muscle,	376	634
fascia and tendon of index finger at wrist and	608	054
hand level)	000	
S76.09 family (Other specified injury of	376	634
muscle, fascia and tendon of hip)	608	054
S76.10 family (Unspecified injury of	376	634
quadriceps muscle, fascia and tendon)	608	034
S76.20 family (Unspecified injury of adductor	376	634
muscle, fascia and tendon of thigh)	608	034
		624
S86.00 family (Unspecified injury of right Achilles tendon)	376 608	634
·		624
S86.09 (Other specified injury of Achilles	376	634
tendon), \$96.00 family (Unspecified injury of	608	
muscle and tendon of long flexor muscle of		
toe at ankle and foot level)	276	624
S96.09 family (Other injury of muscle and	376	634
tendon of long flexor muscle of toe at ankle	608	
and foot level)		

S96.10 family (Unspecified injury of muscle	376	634
and tendon of long extensor muscle of toe at	608	
ankle and foot level)		
S96.19 family (Other specified injury of muscle	376	634
and tendon of long extensor muscle of toe at	608	
ankle and foot level)		
S96.20 family (Unspecified injury of intrinsic	376	634
muscle and tendon at ankle and foot level)	608	
S96.29 family (Other specified injury of	376	634
intrinsic muscle and tendon at ankle and foot	608	
level)		

- 26) Add the S76.29 family (Other injury of adductor muscle, fascia and tendon of right thigh), S76.39 family (Other specified injury of muscle, fascia and tendon of the posterior muscle group at thigh level), S86.19 (Other injury of other muscle(s) and tendon(s) of posterior muscle group at lower leg level), S86.29 (Other injury of muscle(s) and tendon(s) of anterior muscle group at lower leg level) and S86.39 (Other injury of muscle(s) and tendon(s) of peroneal muscle group at lower leg level) to lines 376, 432 INTERNAL DERANGEMENT OF KNEE AND LIGAMENTOUS DISRUPTIONS OF THE KNEE, RESULTING IN SIGNIFICANT INJURY/IMPAIRMENT, and 608 and remove from line 634
- 27) Add HCPCS C97640-C9767 (Revascularization, endovascular, open or percutaneous, lower extremity artery(ies), except tibial/peroneal; with intravascular lithotripsy and transluminal stent placement) to line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS
 - a. Modify GN173 as shown in Appendix A
- 28) Remove ICD-10-CM F98.3 (Pica of infancy and childhood) from line 631 PICA
- 29) Rename line 631 PICA IN ADULTS
- 30) Make no change in the non-pairing of mid-foot arthrosis with foot arthritis

MOTION: To approve the recommendations stated in the consent agenda. CARRIES 6-0.

> Topic: Chemodenervation (botulinum toxin) guideline update

Discussion: There was no discussion about this topic.

Recommended Actions:

1) Modify GN219 as shown in Appendix A

MOTION: To recommend the guideline note changes as presented. CARRIES 6-0.

> Topic: Enteropathic arthropathies

Discussion: There was no discussion about this topic.

Recommended Actions:

- 1) Remove ICD-10-CM M07.6 code family (enteropathic arthropathy) from line 659 MUSCULOSKELETAL CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY.
- 2) Add ICD-10-CM M07.6 family (enteropathic arthropathy) to line 46 RHEUMATOID ARTHRITIS AND OTHER INFLAMMATORY POLYARTHROPATHIES

MOTION: To recommend the code changes as presented. CARRIES 6-0.

> Topic: Erythropoietin in chronic kidney disease

Discussion: There was concern about the proposed addition of coverage of ICD-10-CM D63.1 (Anemia in chronic kidney disease). This code is listed in coding guidelines as "epo resistant anemia." It also does not specify what level of renal dysfunction is required for treatment. Staff were instructed to clarify this topic and bring back to a future meeting.

> Topic: Pelvic congestion syndrome

Discussion: There was no discussion about this topic.

Recommended Actions:

 Add a new guideline note to line 532 CHRONIC PELVIC INFLAMMATORY DISEASE, PELVIC PAIN SYNDROME, DYSPAREUNIA as shown in Appendix B

MOTION: To recommend the guideline note changes as presented. CARRIES 6-0.

> Topic: Platelet rich plasma

Discussion: Smits introduced the summary document. Olson requested that when prior coverage guidances are referenced in a review, that a link to or a copy of that coverage guidance be provided.

Recommended Actions:

- 1) Add CPT 0232T (Injection(s), platelet rich plasma, any site, including image guidance, harvesting and preparation when performed) to line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS
 - a. Modify GN173 as shown in Appendix A

MOTION: To recommend the code and guideline note changes as presented. CARRIES 6-0.

> Topic: Breast reconstruction after lumpectomy

Discussion: There was no discussion about this topic.

Recommended Actions:

1) Modify Guideline Note 79 as shown in Appendix A

MOTION: To recommend the guideline note changes as presented. CARRIES 6-0.

> Topic: Breast MRI guidelines

Discussion: Smits introduced the summary document. Olson expressed concerns that situations recommended by NCCN and American Society of Breast Surgeons, such as multifocal disease, lobular breast cancer, extremely dense breasts or discrepancies in tumor size between imaging studies. In these cases, MRI can help determine whether a patient is a candidate for a lumpectomy rather than a mastectomy, or whether a patient requires a bilateral mastectomy. The new breast MRI guideline was modified to include such coverage.

Gingerich suggested deleting the reference to the breast MRI coverage guidance from the new guideline as the coverage guidance has been retired. This was accepted without discussion.

Recommended Actions:

- 1) Retire the following Coverage Guidances
 - a. Breast Cancer Screening in Women at Above Average Risk
 - b. PET For Breast Cancer (recently revised PET coverage criteria)
 - c. MRI for Breast Cancer Diagnosis (last affirmed 2016)
 - d. MRI for Breast Cancer Screening (outdated)
- 2) Delete Diagnostic Guideline D9 and Guideline Note 26
- 3) Revise Diagnostic Guideline D6 as shown in Appendix A

MOTION: To recommend the guideline note changes as amended. CARRIES 6-0.

> Topic: Actinic keratoses

Discussion: There was no discussion about this topic.

Recommended Actions:

 Make no change in the placement of ICD-10 L57.0 (Actinic keratoses) on line 627 BENIGN NEOPLASMS OF SKIN AND OTHER SOFT TISSUES.

> Topic: Radiofrequency ablation and cryotherapy for select renal cell cancers

Discussion: Smits introduced the summary document. There was some discussion regarding whether to cover these procedures for renal cell cancers up to 4 cm. The group decided that the major guidelines recommended under 3cm and that size was kept in the proposed new guideline.

Recommended Actions:

- 1) Add CPT 50592 (Ablation, 1 or more renal tumor(s), percutaneous, unilateral, radiofrequency) and 50593 (Ablation, renal tumor(s), unilateral, percutaneous, cryotherapy) to line 214 CANCER OF KIDNEY AND OTHER URINARY ORGANS
 - a. Advise HSD to remove CPT 50593 from the Ancillary Procedures File
 - b. Delete CPT 50592 from line 662/GN173 as shown in Appendix A
- 2) Add a new guideline to line 214 as shown in Appendix B

MOTION: To recommend the code and guideline note changes as presented. CARRIES 6-0.

> Topic: Clarification of the lower urinary tract symptoms (LUTS) guideline

Discussion: There was no discussion about this topic.

Recommended Actions:

1) Modify GN145 as shown in Appendix A

MOTION: To recommend the guideline note changes as presented. CARRIES 6-0.

> Topic: Sensory integration therapy

Discussion: There was no discussion about this topic.

Recommended Actions:

1) Update the GN173 entry as shown in Appendix A

MOTION: To recommend the guideline note changes as presented. CARRIES 6-0.

> Topic: Congenital foot deformity code review

Discussion: Smits introduced the summary document. Hodges requested that staff consult with orthopedics or other subject matter experts to ensure that the coding changes proposed are appropriate. Staff will consult experts and bring this topic back to a future meeting for further discussion.

> Topic: Gait analysis and surface electromyography

Discussion: There was no discussion about this topic.

Recommended Actions:

- Add CPT 96000-96004 (Comprehensive computer-based motion analysis by video-taping and 3D kinematics; Dynamic surface electromyography) to line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS
 - a. Advise HSD to remove these codes from the Ancillary and Diagnostic Procedures files
- 2) Modify GN173 as shown in Appendix A

MOTION: To recommend the code and guideline note changes as presented. CARRIES 6-0.

> Topic: Polydactyly clarification

Discussion: There was no discussion about this topic.

Recommended Actions:

- Remove ICD-10-CM Q69.9 (Polydactyly, unspecified) from line 579 CAVUS DEFORMITY OF FOOT;
 FLAT FOOT; POLYDACTYLY AND SYNDACTYLY OF TOES
- 2) Rename line 579 CAVUS DEFORMITY OF FOOT; FLAT FOOT; POLYDACTYLY AND SYNDACTYLY OF TOES

MOTION: To recommend the code changes as presented. CARRIES 6-0.

> Topic: 2024 Biennial Review: agenesis of lung

Discussion: There was no discussion about this topic.

Recommended Actions:

Effective 1/1/2024:

1) Delete Line 647 AGENESIS OF LUNG

MOTION: To recommend the biennial review change as presented. CARRIES 6-0.

> Topic: 2024 Biennial Review: Dorsal rhizotomy for spastic diplegic cerebral palsy

Discussion: There was no discussion about this topic.

Recommended Actions:

Effective October 1 2022:

- 1) Add CPT 63185 and 63190 (laminectomy with rhizotomy) to line 292 NEUROLOGICAL DYSFUNCTION IN POSTURE AND MOVEMENT CAUSED BY CHRONIC CONDITIONS
- 2) Adopt the new guideline shown in Appendix B for line 292
- Strike through line 491 SPASTIC DIPLEGIA Treatment: RHIZOTOMY for the 10/1/22 Prioritized List

Effective 1/1/2024:

1) Delete Line 491 SPASTIC DIPLEGIA

MOTION: To recommend the code and guideline note changes and Biennial Review changes as presented. CARRIES 6-0.

Public Comment:

No additional public comment was received.

> Issues for next meeting:

- -Coding for erythropoietin in chronic kidney disease
- -Congenital foot deformity review

> Next meeting:

May 19, 2022; Virtual meeting

> Adjournment:

The meeting adjourned at 11:15 AM.

Revised Guideline Notes

DIAGNOSTIC GUIDELINE D6, BREAST CANCER SCREENING IN ABOVE AVERAGE RISK WOMEN BREAST MRI

Annual screening mammography and annual screening MRI are covered only for women at above-average risk of breast cancer. This coverage, beginning at 30 years of age, includes women who have one or more of the following:

- Greater than 20% lifetime risk of breast cancer
- BRCA1 or BRCA2 gene mutation, or who have not been tested for BRCA but have a first-degree relative who is a BRCA carrier
- •A personal history or a first-degree relative diagnosed with Bannayan-Riley-Ruvalcaba syndrome, Cowden syndrome, or Li-Fraumeni syndrome
- Other germline gene mutations known to confer a greater than 20% lifetime risk of breast cancer

For women with a history of high dose chest radiation (≥ 20 Gray) before the age of 30, annual screening MRI and annual screening mammography are covered beginning 8 years after radiation exposure or at age 25, whichever is later.

For women with both a personal history and a family history of breast cancer which give a greater than 20% lifetime risk of breast cancer, annual mammography, annual breast MRI and annual breast ultrasound are covered.

- A) Annual breast MRI screening for high-risk patients
 - 1) For individuals with a genetic mutation known to confer a greater than 20% lifetime risk of breast cancer (e.g. BRCA1, BRCA2, Bannayan-Riley-Ruvalcaba syndrome, Cowden syndrome, or Li-Fraumeni syndrome), beginning 10 years prior to when the youngest family member was diagnosed with breast cancer (but not prior to age 25 years) or age 40 years, whichever comes first
 - 2) For individuals who received high dose chest radiation (≥ 20 Gray) between the ages of 10 and 30 years beginning 8 years after radiation exposure or at age 25, whichever is later
 - 3) For individuals with a lifetime risk of ≥ 20% as defined by models that are largely dependent on family history, beginning 10 years prior to when the youngest family member was diagnosed with breast cancer (but not prior to age 25 years) or age 40 years, whichever comes first
- B) Evaluation of possible breast cancer
 - To search for occult breast cancer in patients with Paget's disease of the nipple or in patients with axillary node metastasis when clinical examination and conventional breast imaging fail to detect a primary breast cancer
 - 2) For the further evaluation of suspicious clinical or imaging findings that remain indeterminate after complete mammographic and sonographic evaluations in lesions that do not meet criteria for breast biopsy
- C) Preoperative breast MRI
 - 1) <u>for patients with recently diagnosed breast cancer who qualify for MRI screening based</u> <u>on the high-risk criteria in section A above</u>

- 2) For determining the extent of cancer or presence of multi-focal or multi-centric tumor or the presence of contralateral cancer, in patients with a proven breast cancer and associated clinical or conventional indeterminate imaging findings suspicious for malignancy. This may include patients with invasive lobular carcinoma or extremely dense breast tissue (limiting mammographic sensitivity), or when there are significant discrepancies in the estimated tumor size as measured on clinical exam, mammogram, and ultrasound
- D) Evaluation of suspected breast implant rupture
 - 3) Breast MRI is covered for evaluation of suspected breast implant rupture, if the MRI findings will aid the decision-making for implant removal or aid the diagnostic evaluation of indeterminate clinical or conventional imaging findings in patients with implants

For women with increased breast density, supplemental screening with breast ultrasound, MRI, or digital breast tomosynthesis is not covered.

Breast PET-CT scanning and breast-specific gamma imaging are not covered for breast cancer screening.

For surveillance for a treated breast cancer, see Guideline Note 26 BREAST CANCER SURVEILLANCE.

The development of this guideline note was informed by a HERC <u>coverage guidance</u>. See <u>https://www.oregon.gov/oha/HPA/DSI-HERC/Pages/Evidence-based Reports.aspx</u>

DIAGNOSTIC GUIDELINE D27, SARS-COV-2 (COVID-19) TESTING

Testing for SARS-CoV-2 (COVID-19) virus RNA or viral antigen is a covered diagnostic service. <u>Testing for viral variants/mutations</u> (CPT 87913) is only covered when required to guide patient treatment.

Antibody testing for SARS-CoV-2 (COVID-19; CPT 86413, 86328 or 86769) is covered as diagnostic only when such testing meets the following criteria:

- A) Testing is done using tests that have FDA Emergency Use Authorization (EUA) or FDA approval; AND
- B) Testing is used as part of the diagnostic work up in hospitalized patients of
 - 1) Acute COVID-19 infection in a patient with a previous negative COVID-19 antibody test and a negative COVID-19 RNA or viral antigen test; OR
 - 2) Complications of COVID-19 infection, such as myocarditis, coagulopathy, or multisystem inflammatory syndrome in children (MIS-C) or multisystem inflammatory syndrome in adults (MIS-A).

GUIDELINE NOTE 26, BREAST CANCER SURVEILLANCE

Line 191

History and physical exam is indicated every 3 to 6 months for the first three years after primary therapy, then every 6-12 months for the next 2 years, then annually thereafter.

Mammography is indicated annually, and patients treated with breast-conserving therapy, initial mammogram of the affected breast should be 6 months after completion of radiotherapy.

No other surveillance testing is indicated.

For ongoing screening for a new breast cancer, see Guideline Note 2006 BREAST CANCER SCREENING IN ABOVE AVERAGE RISK WOMEN.

GUIDELINE NOTE 73, PENILE ANOMALIES

Lines 424,433,571,658

Congenital anomalies of the penis (ICD-10-CM Q54.4, Q55.5 and Q55.6) are included on Line 434 only when they

- A. Are associated with hypospadias, OR
- B. Result in documented urinary retention, OR
- C. Result in repeated urinary tract infections, OR
- D. Result in recurrent infections such as meatitis or balanitis, OR
- E. Involve 35 degrees of curvature or greater for conditions resulting in lateral or ventral curvature, OR
- F. Involve 60 degrees of rotation or greater for conditions resulting in penile torsion, OR
- G. Involve aplasia/congenital absence of the penis.

Otherwise, these diagnoses are included on Line 658.

Acquired anomalies of the penis (ICD-10-CM N48.82, N48.83, N48.89 or T81.9XXA) are included on Line 424 only when they are the result of a prior penile procedure AND either

- A. Result in a skin bridge, OR
- B. Result in a buried penis, OR
- C. Are associated with hypospadias, OR
- D. Result in documented urinary retention, OR
- E. Result in repeated urinary tract infections, OR
- F. Result in recurrent infections such as meatitis or balanitis, OR
- G. Involve 35 degrees of curvature or greater for conditions resulting in lateral or ventral curvature, OR
- H. Involve 60 degrees of rotation or greater for conditions resulting in penile torsion.

Otherwise, these diagnoses are included on Line 571 or Line 658.

GUIDELINE NOTE 79, BREAST RECONSTRUCTION

Line 191

Breast reconstruction is only covered after mastectomy, <u>or lumpectomy that results in a significant deformity or asymmetry</u>, as a treatment for breast cancer or as prophylactic treatment for the prevention of breast cancer in a woman who qualifies under Guideline Note 3, and must be completed within 5 years of initial mastectomy <u>or lumpectomy</u>.

Breast reconstruction may include contralateral reduction mammoplasty (CPT 19318) or contralateral mastopexy (CPT 19316). Mastopexy is only to be covered when contralateral reduction mammaplasty is inappropriate for breast reconstruction and mastopexy will accomplish the desired reconstruction result.

GUIDELINE NOTE 145, TREATMENTS FOR BENIGN PROSTATE ENLARGEMENT WITH LOWER URINARY TRACT SYMPTOMS

Line 327

For men with lower urinary tract symptoms (LUTS) due to benign prostate enlargement, surgical procedures are included on these lines only if symptoms are severe, and if drug treatment and conservative management options have been unsuccessful or are not appropriate. hyperplasia (BPH), surgical procedures are included on this line for patients with one of the following:

- A) Refractory urinary retention; OR
- B) Recurrent urinary tract infections due to BPH; OR
- C) Recurrent bladder stones or gross hematuria due to BPH; OR
- D) Severe symptoms (International Prostate Symptom Score (IPSS) of 20-35) in patients who are not candidates for drug treatment due to intolerable side effects or have failed combination therapy with an alpha-blocker and 5-alpha reductase inhibitor for at least 3 months.

Prostatic urethral lift procedures (CPT 52441, 52442, HCPCS C9739, C9740) are included on Line 327 when the following criteria are met:

- Age 50 or older
- Estimated prostate volume < 80 cc
- IPSS ≥ 13
- No obstructive median lobe of the prostate identified on cystoscopy at the time of the procedure

The following interventions for benign prostate enlargement are not included on Line 327 due to lack of evidence of effectiveness:

- Botulinum toxin
- HIFU (High Intensity Focused Ultrasound)
- TEAP (Transurethral Ethanol Ablation of the Prostate)
- Laser coagulation (for example, VLAP/ILC)
- Prostatic artery embolization

The development of this guideline note was informed by a HERC <u>coverage guidance</u>. See <u>https://www.oregon.gov/oha/HPA/DSI-HERC/Pages/Evidence-based-Reports.aspx</u>

GUIDELINE NOTE 219, CHEMODENERVATION

Lines 292,327,351,362,378,393,410,<u>500,</u>517,<u>526</u>

Inclusion of chemodenervation on the Prioritized List has the following limitations for the lines specified below:

Line 292 NEUROLOGICAL DYSFUNCTION IN POSTURE AND MOVEMENT CAUSED BY CHRONIC CONDITIONS

Chemodenervation with botulinum toxin injection (CPT 64642-64647) is included on this line for treatment of upper and lower limb spasticity (ICD-10-CM codes G24.02, G24.1, G35, G36.0, I69.03- I69.06 and categories G71, and G80-G83)

Line 327 FUNCTIONAL AND MECHANICAL DISORDERS OF THE GENITOURINARY SYSTEM INCLUDING BLADDER OUTLET OBSTRUCTION

Chemodenervation of the bladder (CPT 52287) is included on this line only for treatment of idiopathic detrusor over-activity or neurogenic detrusor over-activity (ICD-10-CM N32.81) in

patients who have not responded to or been unable to tolerate at least two urinary incontinence antimuscarinic therapies (e.g. fesoterodine, oxybutynin, solifenacin, darifenacin, tolterodine, trospium). Treatment is limited to 90 days, with additional treatment only if the patient shows documented positive response. Positive response to therapy is defined as a reduction of urinary frequency of 8 episodes per day or urinary incontinence of 2 episodes per day compared to baseline frequency.

Line 351 STRABISMUS DUE TO NEUROLOGIC DISORDER

Chemodenervation with botulinum toxin injection (CPT 67345) is included on this line for the treatment of strabismus due to other neurological disorders (ICD-10-CM H50.89).

Line 362 DYSTONIA (UNCONTROLLABLE); LARYNGEAL SPASM

Chemodenervation with botulinum toxin injection (CPT 64612, 64616) is included on this line only for treatment of blepharospasm (ICD-10-CM G24.5), spasmodic torticollis (ICD-10-CM G24.3), and other fragments of torsion dystonia (ICD-10-CM G24.9).

Line 378 ESOPHAGEAL STRICTURE; ACHALASIA

Chemodenervation with botulinum toxin injection (CPT 43201) is included on this line for treatment of achalasia (ICD-10 K22.0).

Line 393 STRABISMUS WITHOUT AMBLYOPIA AND OTHER DISORDERS OF BINOCULAR EYE MOVEMENTS; CONGENITAL ANOMALIES OF EYE; LACRIMAL DUCT OBSTRUCTION IN CHILDREN

Chemodenervation with botulinum toxin injection (CPT 67345) is included on this line for the treatment of strabismus due to other neurological disorders (ICD-10-CM H50.89).

Line 410 MIGRAINE HEADACHES

Chemodenervation for treatment of chronic migraine (CPT 64615) is included on this line for prophylactic treatment of adults who meet all of the following criteria:

- A) have chronic migraine defined as headaches on at least 15 days per month of which at least 8 days are with migraine
 - B) has not responded to or have contraindications to at least three prior pharmacological prophylaxis therapies (e.g. beta-blocker, anticonvulsant or tricyclic antidepressant)
 - C) their condition has been appropriately managed for medication overuse
 - D) treatment is administered in consultation with a neurologist or headache specialist.

Treatment is limited to two injections given 3 months apart. Additional treatment requires documented positive response to therapy. Positive response to therapy is defined as a reduction of at least 7 headache days per month compared to baseline headache frequency.

<u>Line 500 SIALOLITHIASIS, MUCOCELE, DISTURBANCE OF SALIVARY SECRETION, OTHER AND UNSPECIFIED DISEASES OF SALIVARY GLANDS</u>

<u>Chemodenervation with botulinum toxin injection (CPT 64611) is included on this line for the treatment of excessive salivation.</u>

Line 517 DISORDERS OF SWEAT GLANDS

Chemodenervation with botulinum toxin injection (CPT 64650, 64653) is included on this line for the treatment of axillary hyperhidrosis and palmar hyperhidrosis (ICD-10-CM L74.52, R61).

Line 526 CHRONIC ANAL FISSURE

<u>Chemodenervation with botulinum toxin injection (CPT 46505) is included on this line for the treatment of anal fissures.</u>

GUIDELINE NOTE 173, INTERVENTIONS THAT ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS FOR CERTAIN CONDITIONS

Line 662

The following Interventions are prioritized on Line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS:

Procedure	Intervention Description	Rationale	Last Review
Code			
<u>0232T</u>	Injection(s), platelet rich plasma,	Insufficient evidence of	March 2022
	any site, including image guidance,	effectiveness	
	harvesting and preparation when		
	performed		
C9764-C9767	Revascularization, endovascular,	Insufficient evidence of	March 2022
C9772-C9775	open or percutaneous, <u>lower</u>	effectiveness	
	extremity artery(ies)		
	tibial/peroneal artery(ies), with		
	intravascular lithotripsy		
50592	Radiofrequency ablation, 1 or	Insufficient evidence of	December
	more renal tumor(s)	effectiveness	2005
97533	Sensory integrative techniques to	Insufficient evidence of	August 2010
	enhance sensory processing and	effectiveness	
	promote adaptive responses to		March 2022
	environmental demands		
96000-96004	Comprehensive computer-based	Insufficient evidence of	March 2022
	motion analysis by video-taping	<u>effectiveness</u>	
	and 3D kinematics		
	<u>Dynamic surface</u>		
	electromyography		

Appendix B

New Guideline Notes

GUIDELINE NOTE XXX PELVIC CONGESTION SYNDROME

Line 532

Pelvic congestion syndrome is included on this line using ICD-10-CM N94.89. This condition does not pair with any vein embolization procedures due to lack of evidence of effectiveness.

GUIDELINE NOTE XXX THERMAL ABLATION OF RENAL CELL CARCINOMA

Line 214

Thermal ablation (e.g., cryosurgery, radiofrequency ablation; CPT 50592, 50593) is included on this line only when:

- 1) The patient has biopsy-confirmed stage T1 renal cell cancer of <3 cm size; AND
- 2) The patient either has a surgically inoperable tumor(s) or is a poor candidate for standard treatments (i.e., nephrectomy).

GUIDELINE NOTE XXX DORSAL RHIZOTOMY FOR SPASTIC CEREBRAL PALSY

Line 292

Dorsal rhizotomy (CPT 63185 and 63190) is only included on this line for patients who meet ALL of the following criteria:

- A) Has spastic diplegic cerebral palsy (ICD-10-CM G80.1); AND
- B) Is a child aged 2 to 10 years; AND
- C) Has good intrinsic lower extremity motor power, but is limited in ambulation by spasticity; AND
- D) Has the functional capacity and motivation to participate in post-operative rehabilitation; AND
- E) Has failed or been unable to tolerate other conservative treatment (e.g., pharmacotherapy, orthopedic management, physical therapy); AND
- F) Has no contraindications to the procedure (e.g., significant scoliosis, progressive neurological disorders, severe fixed joint deformities)

MINUTES

Evidence-based Guidelines Subcommittee

Online meeting April 7, 2022 2:00-5:10pm

Members Present: Devan Kansagara, MD, Chair; Alison Little, MD, MPH; Lynnea Lindsey, PhD; Leslie Sutton (departed at 3:15); Max Kaiser, DO; Leda Garside, RN, MBA; Lisa Kouzes, DC; Abigail Khan, MD; Ben Hoffman, MD; Mimi McDonnell, MD.

Staff Present: Ariel Smits, MD, MPH; Jason Gingerich; Daphne Peck.

Also Attending: Val King MD, MPH, Shauna Durbin & Bethany Godlewski (OHSU Center for Evidence-based Policy); Aaron Trimble, MD; Ben Botkin (Lund Report); Bobbie Clark; Carrie Woodman; Cathy Daraee; Christina Cronin-Vejar; Christine Fallabel; Derek Rogalsky, MD; Erin Scow; Gary Hansen; Greg Showell; Jaymey Sweeney; Jennifer Gore; Joey Razzano; Kimberly Goddard (Representative Prusak Office); Kym McCornack; Melanie Ewald; Meryam; Paria Zarrinnegar; Paul Ryan; Dr. Alison Christy (Providence Pediatric Neurology); Sarah Lemley; Tim Kelly; Val Halpin; Wendy Nawara; Yarisel.

1. Call to Order

Devan Kansagara called the meeting of the Evidence-based Guidelines Subcommittee (EbGS) to order at 2:00 pm. A quorum of members was present at the meeting.

2. Minutes Review

Minutes from the 12/2/2021 meeting were reviewed and approved 10-0.

3. Staff Report

Gingerich read aloud the orientation statement.

Kansagara welcomed two new members to the subcommittee (McDonnell and Hoffman). The rest of the subcommittee members gave brief introductions.

Kansagara nominated Alison Little to be Vice-Chair of the subcommittee. Little gave a brief introduction, including that she is anticipating to retire at the end of 2022. A motion was made to elect Alison Little as Vice-Chair of the subcommittee. **Motion was approved 10-0**. Gingerich noted that given Little's upcoming retirement, he asked members to indicate their interest for the position in 2023.

4. Review Public Comment: PANDAS/PANS

Kansagara asked the staff to walk through the public comment disposition. Smits summarized the key feedback received from the public during the comment period; similar concerns were grouped together and addressed at the beginning of the public disposition document. There were many comments submitted and Smits reviewed proposed responses to each one. Changes proposed included requiring one pediatric subspecialist for referral for IVIG instead of requiring two subspecialists; expanding access to specialists to include adult providers for adolescents; expanding access to providers to include more provider types; clarifying the length of an appropriate trial of lesser-invasive therapy; allowing use of telemedicine and teleconsultations.

Kansagara outlined the major difference between this proposed recommendation and the recommendation from the last meeting is lowering the requirement from two pediatric subspecialists down to requiring one subspecialist.

Gingerich introduced the three appointed experts for this topic by reading their biographical statements:

Dr. Alison Christy is the Clinical Director for Providence Pediatric Neurology at Providence St. Vincent Medical Center. She is a pediatric neuroimmunologist and her areas of expertise include neuroimmunological disorders, pediatric neurology and movement disorders. She has given multiple talks on the topic of PANDAS/PANS at professional conferences in Portland. She is director of the Doernbecher Immune Brain Disorders Clinic. Christy has no relevant conflicts of interest to disclose.

Dr. Michael Daines is Associate Professor and Division Chief of Pediatric Allergy, Immunology and Rheumatology at University of Arizona. He is also the Co-Director for the Children's Post Infectious Autoimmune Encephalitis Center of Excellence in Tucson, AZ. His specialties are pediatric allergy and immunology. Daines is the lead investigator of an FDA-approved phase 3 clinical trial for IVIG in the treatment of PANS. His division also oversees several active research projects related to PANDAS/PANS and has a registry for patients and family members. Dr. Daines has received industry funding from Octapharma for the design of the Phase 3 IVIG trial (paid to the university). He has also received travel reimbursements from the PACE Foundation, a PANDAS/PANS advocacy organization.

Dr. Paria Zarrinnegar is Assistant Professor of Psychiatry at OHSU, joining in 2018. She is a board-certified psychiatrist who specializes in biopsychosocial assessment among children and adolescents. Zarrinnegar has no relevant conflicts of interest to disclose.

Kansagara invited public testimony, and limited the testimony to three minutes per person.

Public testimony

<u>Sarah Lemley, Director of the Northwest PANDAS PANS Network</u>: Ms. Lemley said she has no conflicts. She thanked the subcommittee for their deliberations on this topic. She urged the committee to adopt Option 2 and said that Option 1 imposes cumbersome barriers for families who are already stretched thin. Requiring one pediatric subspecialist would align Oregon with most other states in allowing a treatment pathway for IVIG.

<u>Paul Ryan, PACE Foundation (PANDAS/PANS advocacy group)</u>: Mr. Ryan said he has no conflicts. He urged the subcommittee to vote for Option 2 and to expand the provider type eligible to treat this population to include rheumatologists and infectious disease specialists. He thanked the committee for their time.

<u>Christina Cronin-Vejar, parent</u>: Ms. Cronin-Vejar said she has no conflicts. She described her child's disease and treatment history, including tonsillectomy. She said her family considered IVIG but could not afford the treatment. She said there is a significant lack of providers who know about PANDAS/PANS and even fewer who are comfortable treating this population. She urged the subcommittee to vote for Option 2 and said these children deserve appropriate medical treatment.

<u>Kym McCornack, parent</u>: Ms. McCornack thanked the subcommittee. She described her daughter's disease and treatment history, including accessing IVIG and subsequent symptom remission. She urged the subcommittee to listen to the family experiences and vote for Option 2.

Sutton asked what other states' coverage of PANDAS/PANS treatments look like and the use of telehealth and/or teleconsultations/provider-to-provider consultations. Mr. Ryan said there is a lot of variety across the eight states but that telehealth has grown in its uptake since the COVID public health emergency. Ryan stated that depends on the subspecialist and whether they decide to see the child inperson or consult with the referring provider. Hoffman said that a physical exam can be a very important component of evaluation, especially for PANDAS/PANS and how complicated these diseases are, and being sure that use of plasmapheresis is only ever prescribed appropriately. Kansagara mentioned the testimony of another testifier from the December meeting and how one child was ultimately diagnosed with Wilson's disease. Kansagara said that e-consults typically consist of a brief review of chart notes and labs and do not allow for the type of in-depth evaluation that a video visit would afford. Lindsey agreed with Kansagara saying that an e-consult does not include an interaction with the child and that the intent of the proposed recommendation is that a subspecialist evaluate the child experiencing PANDAS/PANS.

Gingerich presented the two options that staff had prepared. Kansagara highlighted the differences between the two options. Little asked the appointed experts what their feedback was. Christy described a recent patient story and her consultations with three other pediatric subspecialists within a short period of time, emphasizing that once access to one pediatric subspecialist is achieved, subsequent interactions with subspecialists can be accomplished within a short time frame. Christy noted Daines' testimony from the last meeting was that he only treats about 5% of his patients with IVIG, and that her concern with Option 2 is that it would lead to overtreatment and misdiagnoses. Smits asked if modifying the recommendation to requiring one pediatric subspecialist as well as one e-consult with a second subspecialist would alleviate her concern and Christy responded that that modification seemed reasonable. Kansagara said that is what Option 1(b) currently accomplishes and that the concern from the members of the public, as well as the dearth of available providers, from the past few meetings indicates that requiring multiple subspecialists is too onerous to accomplish.

Khan stated that her concern with allowing use of e-consults at all makes her uncomfortable, stating that e-consultations are typically done with conditions that can be diagnosed in a relatively straightforward manner with a high degree of confidence using objective data, and that is not the case with these conditions. She stated that if the issue is limited resources and a limited provider pool, then e-consults could pose a higher risk to patients and families. Kansagara clarified that Option 1 (b) would require one subspecialist visit with the child and that the second subspecialist encounter could be done

through an e-consult. Khan said that she would expect an e-consult to maintain the same standards of practice, regardless of if it is the first opinion or the second. Kansagara agreed and questioned if the second e-consult becomes superfluous because it wouldn't offer a detailed examination to provide any added benefit.

Kouzes stated a reasonable family physician would seek out opinions from other specialists and that enforcing that process is not necessary. She stated that if the assumption is that the providers are reasonable, then having one primary care physician and the single subspecialist consult is adequate. Hoffman said that part of the risk is, from his own experience in Portland, is that there is potential of abuse and as long as there is oversight, competent primary care physicians and subspecialists should be enough. Kouzes responded that she understands the risk of abuse but that the policy should assume that licensing boards will be the appropriate venue to address abuse. Hoffman agreed. Kansagara said the requirement for one subspecialist to weigh has less to do with the concerns about malpractice than a nuaced decision that requires expertise and there is a real medical value in having at least one subspecialist perform a detailed examination given that some of these treatments have potential harms.

Kaiser said he prefers Option 1 because the diagnostic criteria are evolving for this subset of diseases, the understanding of the disease is evolving, and the understanding of the efficacy of treatments is evolving. These diseases differ than most well-defined conditions that the subcommittee discusses, and given the vulnerability of this population, Option 1 is a good balance. Hoffman said if he is concerned about a child having PANDAS/PANS, he will utilize treatment options within his scope, which does not include plasmapheresis or IVIG. He said if he calls Dr. Christy and she agrees with IVIG, he would feel comfortable proceeding with that treatment plan. Given that, he prefers Option 2. Godlewski said the medical polcieis for Aetna, Moda and Regence Blue Cross Blue Shield all say that IVIG is considered investigational for PANDAS/PANS. McDonnell said she agrees with Khan that adding a provider-toprovider or e-consult would not be a useful requirement. Kansagara agreed and said it would be difficult to meaningfully use e-consults for these conditions. Kouzes and Kaiser discussed whether patients would find subspecialists first or if their primary care provider would be willing to reach out to experts. Hoffman said that the second consultation is a barrier. Lindsey said she does not feel the group does not have a good grasp of how e-consults are defined, and she is worried that the coverage guidance might be unclear as a result. Kansagara said the use of e-consults will differ based on the practice setting, and informally polled the subcommittee members where they stand between the two options. Most members preferred advancing Option 2 to the Commission for consideration. Kouzes made recommendations to adjust the wording of Option 2(b) to include allergists and infectious disease specialists as well as some other edits.

A motion was made to refer the draft coverage guidance with Option 2 as modified to the Health Evidence Review Commission for review. **Motion approved 7-1 (Nay: Kaiser, Abstained: Little, Absent: Sutton).**

DRAFT HERC Coverage Guidance

Tonsillectomy, adenoidectomy, adenotonsillectomy, and prophylactic antibiotic therapy are not recommended for coverage to treat pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) and pediatric acute-onset neuropsychiatric syndrome (PANS) (weak recommendation).

Up to 3 monthly immunomodulatory courses of intravenous immunoglobulin (IVIG) therapy or therapeutic plasma exchange are recommended for coverage to treat PANDAS and PANS (*weak recommendation*) when both of the following are met:

- A) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- B) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of IVIG or plasma exchange. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.

Note: Other treatments (corticosteroids, SSRIs, NSAIDs, short-course antibiotics, and behavioral therapies) were included in an initial version of this report. These therapies were determined to be beyond the scope of a HERC coverage guidance, as these therapies are commonly used for many indications and are not typically subject to utilization control. Only treatments subject to coverage criteria were retained for the final version of this report.

5. Review Coverage Guidance: High-Frequency Chest Wall Oscillation Devices

Gingerich introduced Dr. Aaron Trimble, the appointed expert for this topic, by reading aloud his biographical statement:

Dr. Aaron Trimble is Assistant Professor in Pulmonary and Critical Care Medicine at Oregon Health and Science University. He has expertise in pulmonology and conducts research in cystic fibrosis and mucociliary clearance. He prescribes high-frequency chest wall oscillation devices for patients with cystic fibrosis and bronchiectasis and is also part of the adult CF clinic at OHSU. He has received grant funding from the Cystic Fibrosis Foundation to study high-frequency chest wall oscillation devices. He has also received research funding and food/travel/beverages for his work on CF medications.

Kansagara outlined the process for reviewing the draft coverage guidance for the new members. Gingerich presented the GRADE table as King summarized the evidence overview for the draft coverage guidance, including revisions made to the hospitalizations and pulmonary exacerbations outcomes for the cystic fibrosis GRADE table. This resulted in rating the hospitalization outcome from having a 'low' confidence of evidence to a 'very low' confidence of evidence. Smits summarized the changes made to the rationale and balance of benefits and harms tables as a result of the downgraded confidence of evidence.

Smits summarized the non-cystic fibrosis bronchiectasis GRADE table, stating that staff relied on expert testimony as well as extrapolated evidence for cystic fibrosis to make a recommendation for coverage for this population. Smits presented the option of recommending against coverage for non-cystic fibrosis bronchiectasis. There were no changes in recommendations made for the other two conditions included in the draft coverage guidance.

Kansagara summarized the reasons that the subcommittee initially voted to recommend coverage for cystic fibrosis, including that children need treatment options when manual chest physiotherapy is not available. McDonnell asked why chest wall oscillation devices would be offered to patients with cystic fibrosis if chest physiotherapy failed (criterion A). Trimble responded by saying that the heterogeneity of patients with cystic fibrosis undergoing different treatments will result in differential outcomes for individual patients. Therefore, failing chest physiotherapy does not mean the vest option will fail, as some patients respond to the vest better than others. Hoffman agreed and added that even if a caregiver is capable of performing chest physiotherapy, it is onerous to children and their families. Trimble responded that there are more adults than children with cystic fibrosis now that life expectancy has increased and that adults with cystic fibrosis do not have parents who are capable of performing manual therapy and might other treatment options.

Smits reviewed the recommendations for each of the four conditions. Smits asked if the group wanted to change the recommendation for the cystic fibrosis condition. Lindsey said the testimony heard in prior meetings was compelling enough that the weaker evidence does not change her decision to vote to recommend coverage. Little stated that she previously voted not to recommend coverage and will continue to do so. Smits then surveyed the group if they wanted to change their recommendation for non-cystic fibrosis bronchiectasis. McDonnell noted the paucity of evidence for this condition. Smits reviewed the recommendations for chronic obstructive pulmonary disease and neuromuscular diseases. Kansagara stated that the neuromuscular diseases group contains a heterogenous array of individuals with various disorders and given the small size of each subpopulation, it is very hard to study and highly unlikely that future evidence would emerge, necessitating the reliance on expert testimony. Trimble said that chest wall oscillation is designed to clear the airway and that people with neuromuscular diseases often have issues with aspiration.

Gingerich presented the HERC Guidance Development Framework and Kansagara discussed the flowchart to help organize the various factors the group needs to consider when making coverage decisions. The group discussed costs associated with vest purchases and claims data that can provide context for the decision. Kansagara asked the group what they felt about keeping the COPD recommendation unchanged; there was no discussion. He then surveyed the group about the neuromuscular diseases condition; there was no discussion. Kansagara asked the group about the cystic fibrosis recommendation; there was no discussion. Kansagara asked the group about the non-cystic fibrosis bronchiectasis recommendation; Lindsey stated she would vote for non-coverage for this

coniditon. Hoffman agreed with Lindsey. Kouzes said that posting the non-coverage recommendation for comment might generate more public comment and provide more insight for the next meeting.

Smits revised the rationale table for non-cystic fibrosis bronchiectasis, stating that the larger population makes it more feasible to expect that a trial could be conducted to generate future research.

Trimble stated that he has concerns of equity if the subcommittee relies on a no-vote to generate public comment to refine the decision-making for the next meeting. He does not have any confidence that the patients he sees will or are able to engage in this process, citing language and socioeconomic barriers. Garside said she agrees with Trimble's health equity concerns. Gingerich asked if Trimble could submit formal comment during the public comment period that summarizes his sentiment to ensure that important factor is considered at the next meeting. Smits asked Trimble to encourage his colleagues to also engage in the coverage guidance process. There was no public comment.

A motion was made to refer the draft coverage guidance as modified for a formal 21-day public comment period. **Motion approved 8-1 (Nay: Little, Absent: Sutton).**

DRAFT HERC Coverage Guidance

High-frequency chest wall oscillation devices are recommended for coverage for patients with cystic fibrosis (*weak recommendation*) when there is documentation of frequent exacerbations requiring antibiotics, frequent hospitalization, or rapidly declining lung function measured by spirometry, despite either:

- A) having received chest physiotherapy and positive expiratory pressure therapy, OR
- B) documentation that chest physiotherapy and positive expiratory pressure devices are not tolerated or not available (e.g., inability of a caregiver to perform chest physiotherapy).

High-frequency chest wall oscillation devices are not recommended for coverage for patients with non-cystic fibrosis bronchiectasis (*weak recommendation*).

High-frequency chest wall oscillation devices are recommended for coverage for patients with neuromuscular disease resulting in chronic lung disease (*weak recommendation*) when there is evidence of chronic lung infection, despite either:

- A) having received chest physiotherapy and positive expiratory pressure therapy, OR
- B) documentation that chest physiotherapy and positive expiratory pressure devices are not tolerated, contraindicated, or not available (e.g., inability of a caregiver to perform chest physiotherapy).

High-frequency chest wall oscillation devices are not recommended for coverage for patients with chronic obstructive pulmonary disease (weak recommendation).

6. Review Scope Statement: Bariatric Procedures

Gingerich presented the scope statement. Smits summarized the document. Kansagara invited public testimony.

Public Testimony

<u>Greg Showell</u>: Mr. Showell said he is a registered nurse and a bariatric program coordinator in Corvallis. He said these bariatric interventions are needed and lowering the BMI to 30 expands access to care. He said the bariatric surgery candidates undergo a rigorous screening process to ensure that patients are good candidates for these procedures.

<u>Derek Rogalsky</u>: Dr. Rolgalsky is a bariatric surgery in Coos Bay and states he has no conflicts. He said he performs his procedures at an MBSAQIP-accredited center. He agrees that bariatric surgery should be expanded to adolescents. He had minor comments regarding the scope statement. He encouraged the study designs to be expanded beyond RCTs to include prospective cohort studies. He briefly summarized the evidence profiles of various bariatric procedures.

McDonnell asked about including a question regarding revisional bariatric procedures; King said that could be added as a subgroup for Key Question 3. The group discussed eligible study designs that could be included.

A motion was made to refer the scope statement as modified to the Health Evidence Review Commission for review. **Motion approved 9-0 (Nay: 0, Absent: Sutton).**

7. Review scope statement: Continuous glucose monitors

Continuous glucose monitors scope statement was tabled until the next meeting due to time constraints.

8. Adjournment

The meeting was adjourned at 5:10 pm. The next meeting is scheduled for June 2 from 2:00-5:00 pm online.

Section 2.0 VBBS Report

Color Key

Topics under development
Upcoming discussion topics
Reviewed but no changes planned
Already approved changes

			Planned	Summary of change (or recommended change, decision not	Larger
Request source	Topic Description	Meeting Date	Imp. Date	to change)	cost
Staff review	Deformities of upper body and all limbs	tbd		Review with orthopedics expert	
	Congenital anomalies of knee (Knee				
Staff review	problems since birth)	tbd		Review with orthopedics expert	
	Genitourinary with minimal or no			Co	
	treatment required (genital and urinary				
Staff review	organs)	tbd		Review with urology expert	
				Consider adding insomnia above the funding line for	
	Sleep disorders other than sleep apnea			cognitive behavioral therapy for insomnia (CBTi). Consider	
Staff review	(including insomnia)	tbd		role of medication.	
Staff (Val King)	Temporomandibular Joint Syndrome (TMJ) (Pain and dysfunction in the jaw joint and muscles controlling jaw movement)	tbd		Needs evidence review for medical and surgical treatments	
HSD nurse	movementy	tou		Proposal to add to covered nerve lesion line with ulnar nerve	
reviewer	Median and radial nerve lesions	8/11/2022		lesions	
	Benign neoplasm of the digestive system (Surgery for an abnormal growth found in the stomach or	3			
Staff review	intestines)	5/19/2022		Proposal to add benign carcinoid tumors to funded region	
	Bilateral bone anchored hearing aids (BAHA) (A specific type of hearing aid				
HSD	for children)	5/19/2022		Proposal to expand coverage from unilateral to bilateral	
	Scrotal varices (An enlargement of the veins within the skin that holds the			Already on line 327 as well as line 548 with no guideline.	
Staff review	testicles (scrotum))	5/19/2022		Propose to remove from line 548 and change name of line	
Staff review	Other complications of a procedure	5/19/2022		Propose to rename line "Minor" as diagnoses are minor	

			Planned	Summary of change (or recommended change, decision not	Larger
Request source	Topic Description	Meeting Date	Imp. Date	to change)	cost
	Anemias due to kidney diseases			0	
	(erythropoietin) (A drug to treat low			Recommend clarifying coverage of erythropoietin for non-	
Staff review	blood count caused by kidney disease)	5/19/2022		end stage kidney disease	
	Conduct disorder/impulse disorders (A			Working with partners, will review with Behavioral Health	
Dr. Hoffman	, , ,	tbd		Advisory panel (BHAP).	
	Somatic symptoms line (Extreme				
	feelings and anxiety about physical			Review with BHAP for any need for reprioritization of one or	
Staff review	symptoms)	tbd		more diagnoses or of entire line	
o. 66	- 6 66			Under review with ortho and podiatry, likely August 2022	
Staff review	Deformities of foot	tbd		review	
	Physical therapy for minor				
	musculoskeletal conditions (Injuries and				
	disorders that affect the human body's				
	movement or muscles, tendons,				
Staff review	ligaments, nerves, discs, blood vessels, etc.)			Limited benefit; would be very difficult to implement	
Stall Teview	etc.)			limited benefit, would be very difficult to implement	
	Allergic rhinitis (Nasal allergies/Hay		,	No change; little impact on health except when comorbidity	
Dr. Hoffman	fever)			or growth/development/school exceptions apply	
Dr. Homman	Angiodema (Swelling (edema) of the	-60		or growthy developmenty school exceptions apply	
	lower layer of skin and tissue just under			Removed unfunded duplicate line (no substantive change,	
Dr. Hoffman		11/18/2021	1/1/2022	was already covered)	
		,,	_, _,	No change made; serious benign neoplasms are on line 401;	
Dr. Hoffman	Benign bone neoplasm			Guideline 137 clarifies which are covered.	
	Congenital anomalies of female genital			No change: Diagnoses on this line have no treatment. Other	
Dr. Hoffman	tract excluding vagina			anomalies that require repair are on funded line(s)	
				No change; primary care and preferred medications should	
Dr. Hoffman	Dermatophytoses (ringworm, etc.)			be sufficient for these conditions	
				No change: Primary care and preferred medications	
Dr. Hoffman	Diaper rash			(nystatin) should be sufficient	

			Planned	Summary of change (or recommended change, decision not	Larger
Request source	Topic Description	Meeting Date	Imp. Date	to change)	cost
				No change; primary care and preferred medications (NSAIDS,	
Dr. Hoffman	Dysmenorrhea			birth control) should be sufficient for these conditions	
				No change; primary care and preferred meds should be	
				sufficient for these conditions. Rare exceptions can be	
Dr. Hoffman	Hodeolum/chalazeon			considered through existing processes	
				No change; primary care and preferred medications should	
Dr. Hoffman	Mild eczema			be sufficient for these conditions	
				No change; primary care and preferred medications should	
Dr. Hoffman	Mild psoriasis			be sufficient for these conditions	
				No change: Primary care and preferred medications should	
Dr. Hoffman	Minor burns			be sufficient	
	Pica (Persistent eating of non-food			No change: Removed ambiguity of coverage for pica in	
	items (for example clay, wool, lead,			children (should have already been in funded region),	
	wood) at an age when it is considered			renamed line to clarify that the unfunded line is "Pica in	
Advocates	to be developmentally inappropriate)	3/10/2022	10/1/2022	adults"	
				No change; primary care and preferred medications should	
Dr. Hoffman	Symptomatic urticaria			be sufficient for these conditions	
				Liver angiosarcoma has a very poor prognosis with any	
	Angiosarcoma of liver; intrahepatic bile			treatment (6 months even with surgery). Per NIH, the only	
Staff review	duct carcinoma			treatment of bile duct carcinoma is palliative care	
Staff review	Central retinal artery occlusion	Z)		Reviewed; no effective treatment is available	
				Only microotia (ICD10 17.2) might be considered to move to	
		1		funded line and most treatment recommendations are only	
	25			to repair for costmetic reasons. Severe microotia (grade 3	
	Congenital ear anomalies without			and 4) would have hearing impairment and the hearing	
Dr. Hoffman	hearing impairment			issues are addressed on line 311	
	Co				
				Cognitive behavioral therapy would be available with	
	Conversion disorders F44.4-7, include			another underlying disorder such as depression. No other	
Dr. Hoffman	non-epilectic seizures			treatment for actual disorder indicated	

_			Planned	Summary of change (or recommended change, decision not	_
Request source	Topic Description	Meeting Date	Imp. Date	to change)	cost
Staff review	Cysts of Bartholin's gland and vulva			N75.1 (Abscess of Bartholin's gland) is included on line 205. Cysts typically have no symptoms and do not need treatment	
Stall Teview	Cysts of Bartholli s gland and vulva			Treatment is directed at underlying diseases, which appear	
Staff review	Enophthalmos			in funded region	
Dr. Haffman	Infectious mononucleosis			Primary care should be sufficient; there is no treatment for	
Dr. Hoffman	Miscellaneous rare congenital			this condition	
Staff review	anomalies			Individual consideration will be required	
Staff review	Nasal polyps			and saline. Surgery indicated if causing chronic sinusitis due to blockage of sinus ostia (would be covered on chronic sinusitis line)	
Staff review	Personality disorders			No effective treatment	
Staff review	Secondary and ill-defined neoplasms			Treatment should be targeted to primary cancer, which would be covered.	
	Thrombosed and complicated		47	Generally treated with fiber and observation. Could be	
Staff review	hemorrhoids			addressed based on individual review	
Staff review	Tension headaches			Primary care and NSAIDs are effective treatments.	
Staff review	Esophageal ulcer	3/10/2022	10/1/2022	Added to funded region	
Dr. Hoffman	Foreign body in digestive tract	3/10/2022	1/1/2022	Had already been addressed prior to the concern raised, but implementation was pending	
Staff review	Generalized muscle weakness	3/10/2022	10/1/2022	Added to funded region	
				Working on implementation issues; addition to funded	
HSD Staff	Handicapping malocclusion	11/18/2021	1/1/2023	region planned for 1/1/2023	х
CCO	Dorsal rhizotomy	3/10/2022	10/1/2022	Added to funded region	Х
Staff review	Corneal abcess	3/10/2022	10/1/2022	Added to funded region	
	155			Change name of line to reflect mild/moderate; severe forms	
Staff review	Lichen planus	3/12/2020	10/1/2022	on funded line as defined by Guideline Note 21	
Staff review	Mastoiditis	3/12/2020	10/1/2022	Added to funded region	
Dr. Hoffman	Nightmare disorder	11/18/2021	1/1/2022	Added to funded region	
Dr. Hoffman	Oral candidiasis (thrush)	8/12/2021	10/1/2021	Added to funded region for feeding problems in newborns line	

Below the Line Review Summary

			Planned	Summary of change (or recommended change, decision not	Larger
Request source	Topic Description	Meeting Date	Imp. Date	to change)	cost
				Clarified coverage criteria for acquired vs congenital	
	Phimosis (acquired penile			anomalies of the penis. Added to funded region for acquired	
Dr. Hoffman	complications, circumcision etc)	10/7/2021	1/1/2022	anomalies.	
Staff review	Polydactyly	3/12/2020	10/1/2022	Clarified earlier decision to confirm in funded region	
				Created new criteria for septoplasty, clarified conditions for	
	Rhinoplasty/septoplasty/ deviated			coverage. Some new coverage and new limitations for	
Public	septum	8/12/2021	10/1/2022	services that would be cosmetic.	
Advocates	Selective mutism	11/18/2021	1/1/2022	Moved to funded anxiety line	
Staff review	Sjogren syndrome	3/10/2022	10/1/2022	Added to funded region	
Staff review	Tendon and ligament injuries	3/10/2022	10/1/2022	Added to funded region for full tears	
	Viral endocarditis, myocarditis,				
Staff review	pericarditis, cardiomyopathy	3/10/2022	10/1/2022	Added to funded region	
				Added vitiligo as a funded condition. Affects children's social	
Staff review	Vitiligo	10/7/2021	1/1/2022	function	х
Staff review	Acquired torsion of penis	3/10/2022	10/1/2022	Added to funded region	
Staff review	Agenesis of lung	3/10/2022	10/1/2022	Added to funded region for supportive care	
				Added path to coverage for treatments supporting growth,	
EPSDT	Child growth and development	11/18/2021	1/1/2022	development and participation in school for children	х
Staff review	Chronic pancreatitis		1/1/2022	Already merged for 2022 before this review	
Staff review	Vitiligo of eyelid	3/10/2022	10/1/2022		

Visual Field Testing

<u>Question</u>: Should various diagnoses be paired with visual field testing or should visual field testing be made diagnostic?

Question source: Julie Falardeau, MD, OHSU Ophthalmology

<u>Issue</u>: Dr. Falardeau requested consideration of coverage/pairing of multiple diagnoses of visual field testing with a variety of ophthalmologic diseases. Visual field testing is used to determine if a patient has blind spots or visual limitations from eye or central nervous system disease. Visual field testing is coded with CPT 99201-99285, 92002-92014, 92081-92083, and 92133. Visual field testing is on 60+ ophthalmology lines. Evaluation and management and emergency room services (9920—99285) are already on the diagnostic procedures file, so should be covered regardless of the presenting diagnosis.

Dr. Falardeau's specific requests for consideration are listed below:

- 1) Optic neuritis: (ICD-10-CM H46.XX). These diagnoses are currently on line 650 INTRACRANIAL CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY. These conditions result in loss of vision temporarily or permanently based on a variety of causes. Optic neuritis is a common presentation of multiple sclerosis.
 - a. Evidence
 - i. De Lott 2022, review of optic neuritis
 - High-dose corticosteroids, both oral and intravenous (IV), are the most commonly used treatment for acute ON. A meta-analysis of three randomized controlled trials found no benefit in visual acuity recovery at 1 month, 6 months, and 1 year based on the dose or duration of oral treatment. A meta-analysis of two trials comparing placebo to IV corticosteroids of over 3000 mg total also found no significant improvement in visual acuity, contrast sensitivity, or visual field at 6 months
 - the only benefit of corticosteroids was hastened visual recovery within the first 2 weeks, which is the primary indication for treatment.
 Secondary analyses of the trial data suggest that this early benefit is only about 1–2 lines of Snellen acuity
- 2) Pairing of visual field testing (CPT 92081-92083, 92133) with
 - a. Papilledema (ICD-10-CM H47.1X) which is on line 650 INTRACRANIAL CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY [Note ICD-10-CM H47.10 (Unspecified papilledema) is on the Diagnostic Workup File]. Dr. Falardeau writes that in about 10% of papilledema patients, surgery is required to prevent further vision loss. Such surgery would be for the underlying condition causing the papilledema (which would be covered). The most reliable way to monitor optic nerve function is with visual field testing.
 - b. Optic disc atrophy (ICD-10-CM H47.2X) which is on line 654 SENSORY ORGAN CONDITIONS WITH NO OR MINIMALLY EFFECTIVE TREATMENTS OR NO TREATMENT NECESSARY. Dr. Falardeau also notes that optic atrophy may require visual field testing particularly for determination of whether vision is sufficient for driving. Optic atrophy has no treatment; treatment is aimed at the underlying condition (glaucoma, MS, etc.).
 - c. Visual field defect (ICD-10-CM H53.4X) which is on the DIAGNOSTIC WORKUP FILE.

Visual Field Testing

3) Pairing of visual field testing with optic nerve/chiasm tumors (ICD-10 C72.3X). Ophthalmology office visit CPT codes are already on line 294 CANCER OF BRAIN AND NERVOUS SYSTEM

HERC staff recommendation:

- 1) Remove visual field testing (CPT 92081-92083, 92133) from all current lines
 - Advise HSD to add CPT 92081-92083, 92133) to the Diagnostic Procedure File (Codes 99203-99285 for evaluation and management and emergency visits are already on this file).
 - b. There is a possibility for overutilization; however, these procedures are already found on 60+ ophthalmology lines

Plain Language Summary:

<u>Background:</u> An adenoidectomy is an operation done to remove your adenoids, which are a part of the immune system and are located in the back of the nose. The current guideline allows adenoid removal when a second set of ear tubes are placed in children 4 years and older. The proposed update allows for adenoid removal with the first set of ear tubes in some cases.

<u>Should OHP cover this treatment?</u> Yes, staff recommends adenoid removal should be allowed under certain conditions during initial ear tube placement because the benefits outweigh the risk of harms and concern for repeated surgery with anesthesia.

<u>Question</u>: Should the current guideline regarding coverage of adenoidectomy be updated to agree with the current American Academy of Otolaryngology-Head and Neck Surgery (AAO-HNS) guideline?

Question source: Peggy Kelley, MD, Director of Pediatric ENT at Providence Health Plans

<u>Issue</u>: The current guideline regarding adenoidectomy restricts this procedure to "Adenoidectomy is not indicated at the time of first pressure equalization tube insertion. It may be indicated in children aged 4 and older who are having their second set of tubes." This wording was based on the 2016 AAO guideline. AAO has updated their guideline for 2022 and includes indications for adenoidectomy with the first set of tympanostomy tubes in certain circumstances. Dr. Kelley is requesting that HERC reconsider our guideline wording.

From Dr. Kelley

I would like to submit updated guidelines for the indication for adenoidectomy for children getting PE Tubes. The issue I would like to address is Guideline note 5. Guideline Note 51 states, "Adenoidectomy is not indicated at the time of first pressure equalization tube insertion. It may be indicated in children aged 4 and older who are having their second set of tubes." This information was the guideline more than 5 years ago.

The 2016 guideline from the Academy of Otolaryngology Head & Neck surgery was: A new recommendation against adenoidectomy for a primary indication of OME in children <4 years old, including those with prior tympanostomy tubes, unless a distinct indication exists (nasal obstruction, chronic adenoiditis) This leaves in place the recommendation for adenoidectomy for children >age 4 years with nasal symptoms AND/Causing the otitis media with effusion.

The most recent guideline link is <u>Clinical Practice Guideline</u>: <u>Tympanostomy Tubes in Children-American Academy of Otolaryngology-Head and Neck Surgery (AAO-HNS) (entnet.org)</u>. This is the 2022 guideline. It confirms the indication for adenoidectomy with Tubes if there are nasal symptoms and clarifies that the purpose is to try to minimize the need for repeated surgery. From the executive Summary: A new option for the clinician to perform adenoidectomy as an adjunct to tympanostomy tube insertion for children with symptoms directly related to the adenoid (adenoid infection or nasal obstruction) or in children aged 4 years or older to reduce future incidence of recurrent otitis media or the need for repeat tube insertion.

Current Prioritized List status

СРТ	Code description	Current Placement
42820	Tonsillectomy and	42 CLEFT PALATE WITH AIRWAY OBSTRUCTION
	adenoidectomy; younger than	47 DEEP ABSCESSES, INCLUDING APPENDICITIS AND
	age 12	PERIORBITAL ABSCESS
		64 CONGENITAL ANOMALIES OF UPPER ALIMENTARY
		TRACT, EXCLUDING TONGUE
		202 SLEEP APNEA, NARCOLEPSY AND REM BEHAVIORAL DISORDER
		368 STREPTOCOCCAL SORE THROAT AND SCARLET FEVER;
		VINCENT'S DISEASE; ULCER OF TONSIL; UNILATERAL
		HYPERTROPHY OF TONSIL
		551 CHRONIC DISEASE OF TONSILS AND ADENOIDS
42821	Tonsillectomy and	42,47,64,202,368,551
	adenoidectomy; age 12 or over	
42830	Adenoidectomy, primary;	42, 47, 202,
	younger than age 12	311 HEARING LOSS - AGE 5 OR UNDER
		446 HEARING LOSS - OVER AGE OF FIVE
		466 CHRONIC SINUSITIS
		476 CHRONIC OTITIS MEDIA; OPEN WOUND OF EAR
		DRUM
		551
42831	Adenoidectomy, primary; age 12	42,47,202,476,551
	or over	
42835	Adenoidectomy, secondary;	42,47,202,311,446,466,476,551
	younger than age 12	
42836	Adenoidectomy, secondary; age	42,47,202,476,551
	12 or over	

GUIDELINE NOTE 51, CHRONIC OTITIS MEDIA WITH EFFUSION

Lines 311,446,476

Antibiotic and other medication therapy (including antihistamines, decongestants, and nasal steroids) are not indicated for children with chronic otitis media with effusion (OME) (without another appropriate diagnosis).

Patients with specific higher risk conditions (including craniofacial anomalies, Down's syndrome, and cleft palate, or documented speech and language delay) along with hearing loss and chronic otitis media with effusion are intended to be included on Line 311 or Line 446 for children up to and including age 7. Otherwise hearing loss associated with chronic otitis media with effusion (without those specific higher risk conditions) is only included on Line 476.

For coverage to be considered on Line 311, Line 446 or Line 476, there should be a 3 to 6 month watchful waiting period after diagnosis of otitis media with effusion, and if documented hearing loss is greater than or equal to 25dB in the better hearing ear, tympanostomy surgery may be indicated, given short- but not long- term improvement in hearing. Formal audiometry is indicated for children with chronic OME present for 3 months or longer. Children with language delay, learning problems, or

significant hearing loss should have hearing testing upon diagnosis. Children with chronic OME who are not at risk for language delay (such as those with hearing loss <25dB in the better hearing ear) or developmental delay should be reexamined at 3- to 6-month intervals until the effusion is no longer present, significant hearing loss is identified, or structural abnormalities of the eardrum or middle ear are suspected.

Adenoidectomy is not indicated at the time of first pressure equalization tube insertion. It may be indicated in children aged 4 and older who are having their second set of tubes.

Removal of retained tympanostomy tubes requiring anesthesia (CPT code 69424) or as an office visit, is included on Line 424 as a complication, pairing with ICD-10-CM H74.8.

The development of this guideline note was informed by a HERC <u>coverage guidance</u>. See https://www.oregon.gov/oha/HPA/DSI-HERC/Pages/Evidence-based-Reports.aspx

Practice guideline:

- 1) Rosenfeld 2022: Updated AAO clinical practice guideline for tympanostomy tubes in children
 - a. Adjuvant adenoidectomy:
 - i. Clinicians may perform adenoidectomy as an adjunct to tympanostomy tube insertion for children with symptoms directly related to the adenoids (adenoid infection or nasal obstruction) OR in children aged 4 years or older to potentially reduce future incidence of recurrent otitis media or the need for repeat tube insertion
 - Strength of recommendation: <u>option</u> based on randomized controlled trials, meta-analyses, and population-level studies, with a balance of benefits and harms.
 - iii. Aggregate evidence quality: Grade B, based on RCTs for persistence of OME post-surgically, rate of repeat tube insertion, and hearing outcomes; observational studies regarding the rate of tube reinsertion and hearing outcomes; and meta-analyses on the benefit of adenoidectomy in patients greater than 4 years of age as compared with those younger than 4 years of age
 - iv. Level of confidence in evidence: High for symptoms related to adenoids and children over the age of 4 years; medium for role as primary treatment in select populations and role in second tube insertion procedures in patients younger than 4 years
 - Benefits: Optimize management of adenoid-related disease (nasal obstruction, bacterial infection, chronic rhinitis); reduce need for further surgery and anesthesia; optimize hearing outcomes; decreased persistence of MEE after surgery.
 - vi. Risks, harms, costs: Surgical risks of adenoidectomy, additional anesthetic risk related to need for intubation during procedure, bleeding, hypernasality, velopharyngeal insufficiency, nasopharyngeal scarring/stenosis, Grisel's syndrome, longer recovery
 - vii. Benefit-harm assessment: Equilibrium (balance) of benefits vs harms

HERC staff recommendation:

- 1) Modify GN51 as shown below
 - a. Adds line 424 to the guideline as this line is mentioned in the guideline wording
 - b. Modifies wording regarding adenoidectomy based on 2022 AAO guideline

GUIDELINE NOTE 51, CHRONIC OTITIS MEDIA WITH EFFUSION

Lines 311,424,446,476

Antibiotic and other medication therapy (including antihistamines, decongestants, and nasal steroids) are not indicated for children with chronic otitis media with effusion (OME) (without another appropriate diagnosis).

Patients with specific higher risk conditions (including craniofacial anomalies, Down's syndrome, and cleft palate, or documented speech and language delay) along with hearing loss and chronic otitis media with effusion are intended to be included on Line 311 or Line 446 for children up to and including age 7. Otherwise hearing loss associated with chronic otitis media with effusion (without those specific higher risk conditions) is only included on Line 476.

For coverage to be considered on Line 311, Line 446 or Line 476, there should be a 3 to 6 month watchful waiting period after diagnosis of otitis media with effusion, and if documented hearing loss is greater than or equal to 25dB in the better hearing ear, tympanostomy surgery may be indicated, given short- but not long- term improvement in hearing. Formal audiometry is indicated for children with chronic OME present for 3 months or longer. Children with language delay, learning problems, or significant hearing loss should have hearing testing upon diagnosis. Children with chronic OME who are not at risk for language delay (such as those with hearing loss <25dB in the better hearing ear) or developmental delay should be reexamined at 3- to 6-month intervals until the effusion is no longer present, significant hearing loss is identified, or structural abnormalities of the eardrum or middle ear are suspected.

Adenoidectomy is not indicated at the time of first pressure equalization tube insertion. It may be indicated in children aged 4 and older who are having their second set of tubes. included on these lines at the time of tympanostomy tube insertion for children under age 4 with symptoms directly related to the adenoids (adenoid infection or nasal obstruction) OR in children aged 4 years or older.

Removal of retained tympanostomy tubes requiring anesthesia (CPT code 69424) or as an office visit, is included on Line 424 as a complication, pairing with ICD-10-CM H74.8.

The development of this guideline note was informed by a HERC <u>coverage guidance</u>. See <u>https://www.oregon.gov/oha/HPA/DSI-HERC/Pages/Evidence-based-Reports.aspx</u>

Plain Language Summary:

<u>Background:</u> Bone anchored hearing aids (BAHA) is a specific type of hearing aid who have "normal" hearing in one ear. This topic looks at covering this hearing aid for children who have hearing loss is both ears.

<u>Should OHP cover BAHA for children who have hearing loss in both ears?</u> Staff recommends OHP cover this treatment based on expert opinion.

Question: Should bone anchored hearing aids (BAHA) be covered for bilateral use in children?

Question source: Medical management committee as HSD

<u>Issue</u>: Currently, GN 103 limits coverage of bone-anchored hearing aid system (BAHA) to patients with normal hearing in one ear (with or without a hearing aid). Recently, MMC received a request for coverage for bilateral BAHA in a child with bilateral stenosis of the ear canal. Because GN 103 limited use to one ear and required the other ear to have normal hearing, the child was denied BAHA implantation. When coverage of BAHA was discussed in 2014 and 2015, only unilateral use was discussed based on the literature reviewed and on expert pediatric ENT opinion.

The BAHA is a hearing aid which uses the principle of bone conduction. In normal hearing, sound may be transmitted to the inner ear both by air (through the external ear canal) or through the bones of the skull. In individuals who are unable to hear using air conduction, either due to a congenital malformation of the ear canal or due to chronic ear infection, a hearing aid which utilizes bone conduction is the most appropriate.

Current Prioritized List status

The following codes are on lines 311 HEARING LOSS - AGE 5 OR UNDER and 446 HEARING LOSS - OVER AGE OF FIVE:

CPT **69714** (Implantation, osseointegrated implant, skull; with percutaneous attachment to external speech processor)

CPT **69716** (Implantation, osseointegrated implant, skull; with magnetic transcutaneous attachment to external speech processor)

CPT **69717** (Revision or replacement (including removal of existing device), osseointegrated implant, skull; with percutaneous attachment to external speech processor)

CPT **69719** (Revision or replacement (including removal of existing device), osseointegrated implant, skull; with magnetic transcutaneous attachment to external speech processor)

GUIDELINE NOTE 103, BONE ANCHORED HEARING AIDS

Lines 311,446

Bone anchored hearing aids (BAHA, CPT 69714, 69715; HCPCS L8690-L8694) are included on these lines when the following criteria are met:

- A) The patient is aged 5-20 years for implanted bone anchored hearing aids; headband mounted BAHA devices may be used for children under age 5
- B) Treatment is for unilateral severe to profound hearing loss when the contralateral ear has normal hearing with or without a hearing aid
- c) Traditional air amplification hearing aids and contralateral routing of signal (CROS) hearing aid systems are not indicated or have been tried and are found to be not effective

D) Implantation is unilateral.

Use of BAHA for treatment of tinnitus is not covered.

GUIDELINE NOTE 143, TREATMENT OF UNILATERAL HEARING LOSS

Lines 311,446

Unilateral hearing loss treatment is Included on these lines only for children aged 20 and younger with the following conditions:

- 1. For mild to moderate sensorineural unilateral hearing loss (defined as 26-70 dB hearing loss at 500, 1000 and 2000 Hz), first line intervention should be a conventional hearing aid, with second line therapy being contralateral routing of signal (CROS) system
- 2. For severe to profound unilateral sensorineural hearing loss (defined as 71 dB hearing loss or greater at 500, 1000 and 2000 Hz), first line therapy should be a contralateral routing of signal (CROS) system with second line therapy being a bone anchored hearing aid (BAHA). BAHA SoftBand therapy may be first line therapy for children under age 5 or patients with severe ear deformities (e.g. microstia, severe canal atresia).

Cochlear implants are not included on these lines for unilateral hearing loss per Guideline Note 31 COCHLEAR IMPLANTATION.

Evidence

- 1) Mandavia 2017, systematic review on BAHA
 - a) N=39 studies
 - Eighteen articles were retrospective case series; three were case reports; eight were prospective case series; four were systematic reviews; four were narrative reviews; one was a Delphi study, one was a consensus document.
 - b) Using the GRADE criteria, the quality of evidence was classified as of 'very low quality'
- 2) Janssen 2012, a systematic review of BAHA for bilateral permanent conductive hearing loss
 - a) N=11 studies
 - i) All observational
 - b) In most studies, comparisons between unilateral and bilateral BAHA were intra-subject.
 - c) Bilateral BAHA provided audiologic benefit compared to unilateral BAHA (improved thresholds for tones [2 studies], speech in quiet [5 studies] and in noise [3 studies], and improved localization/lateralization [3 studies]) and patients' perceived subjective benefit from bilateral BAHA (3 studies).
 - d) Disadvantages of bilateral BAHAs included listening in noise in some conditions (3 studies), presumed additional cost, and presumed increase in adverse event risk.
 - e) **Conclusion:** Bilateral BAHA provided additional objective and subjective benefit compared to unilateral BAHA; however, there was a limited number of studies available with good quality evidence.

Other payer policies

1) NHS 2013

- a) Criteria for unilateral implantation BAHA will be funded when assessment by a multidisciplinary team leads to a clear recommendation of a BAHA AND confirms all of the criteria below.
 - i) The patient has one of the following:
 - (a) Permanent bilateral conductive or mixed hearing loss.
 - (ь) Bilateral conductive or mixed hearing loss where one ear works better than the other, but clinicians would have considered two air conducting hearing aids (ACHAs) if the type of hearing loss had not precluded their use.
 - (c) Unilateral conductive hearing loss with ear canal stenosis that is unlikely to benefit from meatoplasty; or who have had revision surgery and failed to tolerate ACHA.
 - (a) Profound unilateral sensorineural hearing loss
 - ii) AND The patient is clinically unsuitable for other medical or surgical treatments.
 - iii) Otological indications supporting the use of BAHA include:
 - (a) Congenital malformation of the middle/external or microtia.
 - (ь) Chronically draining ear that does not allow the use of an air conducting hearing aid.
 - (c) Patients with bilateral conductive hearing loss due to ossicular disease (and not appropriate for surgical correction) or unable to be aided by conventional air conducting devices.
 - iv) AND The following audiological criteria should be met: Conduction or mixed hearing loss with a bone conduction pure tone average (0.5, 1, 2, 3 kHz) threshold up to 45 dBHL for the Devino or BP 100,55dB for the Intenso and 70 dB for Cordelle II (Body

- Processor). In the advent of new processors being released manufacturers audiological recommendations should be followed.
- v) Air conduction pure tone average not better than 40 dB (for Adults).
- vi) A maximum speech discrimination score better than 60% when using a phonetically balanced word list.

b) AND

- i) The patient has had preoperative counselling, and has realistic expectations about the benefits and limitations of BAHA. They must be prepared to maintain their device in the long term.
- ii) The patient will be able to keep the area around the fixture clean, either on their own or with help from other people.
- iii) There are no contraindications for BAHA.
- c) In children with binaural congenital hearing loss, intervention should take place as early in life as possible; BAHA may be provided on a headband until the child is old enough for surgery. The minimum age for first surgery, as identified by the equipment manufacturer, is three years. It is recommended that implant surgery be performed in two stages in children of up to 10 years of age. In children with bilateral conductive hearing loss; clinicians may consider bilateral BAHA if a decision is made that this would provide children with the best hearing environment in the classroom situation, following multidisciplinary clinical assessment by the BAHA team.

2) Aetna 2022:

- a) Aetna considers fully or partially implantable bone-anchored hearing aids (BAHAs) or temporal bone stimulators medically necessary prosthetics for persons aged 5 years and older with a unilateral or bilateral conductive or mixed conductive and sensorineural hearing loss who have any of the following conditions, where the condition prevents restoration of hearing using a conventional air-conductive hearing aid and who meet the audiologic criteria below:
 - i) Congenital or surgically induced malformations of the external ear canal or middle ear (such as aural atresia); or
 - ii) Dermatitis of the external ear, including hypersensitivity reactions to ear moulds used in air conduction hearing aids; *or*
 - iii) Hearing loss secondary to otosclerosis in persons who can not undergo stapedectomy; *or*
 - iv) Severe chronic external otitis or otitis media; or
 - v) Tumors of the external ear canal and/or tympanic cavity; or
 - vi) Other conditions in which an air-conduction hearing aid is contraindicated.

b) Audiologic criteria:

- i) Unilateral implant: Conductive or mixed (conductive and sensorineural) hearing loss with pure tone average bone conduction threshold values measured at 0.5, 1, 2, and 3 kHz less than or equal to 45 dB HL (BAHA Attract, BAHA Divino, BAHA BP100, Baha 4, Bonebridge Bone Conduction Implant, Cochlear Osia, Cochlear Osia 2, Cochlear Osia B1300 system, and Sophono Alpha System), 55 dB HL (BAHA 5 Power, Baha 5 Super Power Sound Processor, BAHA Intenso, Ponto Plus Power) or 65 dB HL (BAHA Cordelle II).
- ii) Bilateral implant: Moderate-to-severe bilateral symmetric conductive or mixed (conductive and sensorineural) hearing loss, meeting above-listed bone conduction thresholds in both ears. Symmetric bone conduction threshold is defined as less than:

- (a) 10 dB average difference between ears (measured at 0.5, 1, 2 and 4 kHz) or less than 15 dB difference at individual frequencies (BAHA Divino, Ponto Plus, Ponto Plus Power, Ponto Pro, Sophono Alpha System); *or*
- (ъ) 10 dB average difference between ears (measured at 0.5, 1, 2, and 3 kHz), or less than a 15 dB difference at individual frequencies (BAHA Attract, BAHA BP100, BAHA 4, BAHA 5 Power, Baha 5 Super Power Sound Processor, BAHA Cordelle II, BAHA Intenso, Bonebridge Bone Conduction Implant, Cochlear Osia, Cochlear Osia 2, and Cochlear Osia B1300 system).
- iii) Aetna considers an implantable BAHA for conductive or mixed hearing loss experimental and investigational when criteria are not met because of insufficient evidence in the peer-reviewed published medical literature.
- iv) Aetna considers the use of an implantable BAHA medically necessary in persons with unilateral sensorineural hearing loss (single-sided deafness, i.e., deafness in one ear while the other ear has normal hearing).

3) Idaho Medicaid 2022

- a) Bone-Anchored Hearing Aid Bone-Anchored Hearing Aid (BAHA) is covered for participants under 21 with a prior authorization when medically necessary. It is recommended that participants over the age of five, trial a soft band BAHA before surgery is scheduled. The participant must meet one of the following criteria:
 - The participant is diagnosed with ear canal atresia, no ear canals, and unable to wear an ear mold;
 - ii) The participant is diagnosed with microtia, very small ear canal, and unable to wear an ear mold;
 - iii) The participant has persistently discharging ears and is unable to use air conduction aid:
 - iv) The participant has an ear condition made worse with ear molds; or
 - v) Audiology test results indicate a pure tone average bone conduction threshold of up to 65 dB. Purchase of an auditory non-osseo integrated sound processor includes the headband in its reimbursement

Expert input:

Dr. Peggy Kelley, Providence pediatric ENT clinical director

I would request adding as in indication: Temporary bilateral conductive hearing loss in patients with cleft palate and middle ear effusions until their palate is repaired and PE Tubes can be placed. This would be BAHA on headband. (We no longer place PE tubes until the palate is closed as the ears just constantly drain until the palate is closed.) The BAHA is preferred over a traditional hearing aid in this type of patient. The BAHA corrects the conductive component only which means that there is no overcorrection or hearing damage that would result to the hearing nerve. It is possible for the native hearing to fluctuate with occasional clearing the middle ear and changes in the viscosity of the fluid in the middle ear until the palate is repaired. A traditional hearing aid is set for a specific hearing loss which may be too little or two much. The BAHA working across the bone is not influenced by the middle ear fluctuations.

HERC staff summary

The evidence for bilateral BAHA is very limited, with some benefits seen when compared to unilateral BAHA but increased cost and risk. Private payers and evidence based payers, such as NHS, are covering bilateral BAHA for children with bilateral conductive or mixed conductive/sensorineural hearing loss. The current requirement to have normal hearing in one ear for unilateral implantation is not reflected in any other payer policy reviewed, and per our expert is out of date. Experts recommend coverage of bilateral BAHA in certain situations.

HERC staff recommendation:

1) Modify GN103 as shown below

GUIDELINE NOTE 103, BONE ANCHORED HEARING AIDS

Lines 311,446

Bone anchored hearing aids (BAHA, CPT 69714, 69715; HCPCS L8690-L8694) are included on these lines when the following criteria are met:

- A) The patient is aged 5-20 years for implanted bone anchored hearing aids; headband mounted BAHA devices may be used for children under age 5; AND
- B) Treatment is for unilateral severe to profound hearing loss when the contralateral ear has normal hearing with or without a hearing aid
- c) Traditional air amplification hearing aids and contralateral routing of signal (CROS) hearing aid systems are not indicated or have been tried and are found to be not effective
- D) Implantation is unilateral.
- E) The patient has one of the following:
 - Permanent bilateral conductive or mixed hearing loss (for example, congenital malformation of the middle/external ear, microtia, or ossicular disease) unable to be aided by conventional air conducting devices, OR
 - 2) <u>Unilateral conductive hearing loss with ear canal stenosis or ear canal atresia that is unlikely to benefit from surgery; OR</u>
 - 3) <u>Profound unilateral sensorineural hearing loss when the contralateral ear has normal</u> hearing with or without a hearing aid; OR
 - 4) Temporary bilateral conductive hearing loss in patients with cleft palate and middle ear effusions until their palate is repaired and tympanostomy tubes can be placed (for BAHA headband only); AND
- F) The patient is clinically unsuitable for other medical or surgical treatments.

Use of BAHA for treatment of tinnitus is not covered.

Consent Agenda Issues—May 2022

Code	Code Description	Line(s) Involved	Issue	Recommendation(s)
11960	Insertion of tissue expander(s) for other than breast, including subsequent expansion	Burn lines	OHSU plastic surgery requested that tissue expanders be paired with a variety of disfiguring diagnoses that might require extra	Remove 11960 and 11971 from all current Prioritized List lines.
11971	Removal of tissue expander without insertion of implant		tissue to close the wound from the repair. For example, patients with large congenital nevi frequently have tissue expanders placed to create extra normal tissue for use in closing the incision when the congenital nevi is excised. HERC staff recommend placing these codes on the Ancillary file to be used with various covered procedures as needed.	Advise HSD to add 11960 and 11971 to the Ancillary Procedures File
B4100	Food thickener, administered orally, per ounce	CALL	B4100 is not on any list. This is a standard treatment for dysphagia.	Advise HSD to add B4100 to the Ancillary Procedure File
58559- 58563	Hysteroscopy with various surgical procedures	1 PREGNANCY	A series of CPT codes were added to line 1 in 2017 which were all thought to represent hysterectomy procedures. Five of these codes actually represent hysteroscopy with various procedures such as removal of fibroid or lysis of adhesions which are not appropriate during pregnancy. All of these codes are currently on other, appropriate lines.	Remove 58559-58563 from line 1

Consent Agenda Issues—May 2022

Code	Code Description	Line(s) Involved	Issue	Recommendation(s)
61538	Craniotomy with elevation of bone flap; for lobectomy, temporal lobe, with electrocorticography during surgery	174 GENERALIZED CONVULSIVE OR PARTIAL EPILEPSY WITHOUT MENTION OF IMPAIRMENT OF CONSCIOUSNESS	CPT 61537 (the same come without electrocorticography) appears on line 174. A CCO requested that 61538 be added to the epilepsy surgery line	Add 61538, 61539, and 61781 to line 174
61539	Craniotomy with elevation of bone flap; for lobectomy, other than temporal lobe, partial or total, with electrocorticography during surgery		In addition, CPT 61539 is missing from line 174 when all similar codes (61540-61543) appear on that line	
61781	Stereotactic computer-assisted (navigational) procedure; cranial, intradural		Lastly, CPT 61781 appears on 6 other lines with intracranial surgery but not on line 174. The same CCO requested consideration of addition to line 174	
		572 OTHER COMPLICATIONS OF A PROCEDURE	This line contains only diagnoses for minor complications. HERC staff recommends changing the line title to reflect this.	Change the name of line 572 to OTHER MINOR COMPLICATIONS OF A PROCEDURE
186.1	Scrotal varices	327 FUNCTIONAL AND MECHANICAL DISORDERS OF THE GENITOURINARY SYSTEM INCLUDING BLADDER OUTLET OBSTRUCTION 548 SUBLINGUAL, SCROTAL, AND PELVIC VARICES	ICD-10 I86.1 is on both a covered and an uncovered line with no guideline. Staff recommends removing from lower line to reduce confusion.	Delete I86.1 from line 548 Rename line 548 SUBLINGUAL , SCROTAL, AND PELVIC VARICES

Consent Agenda Issues—May 2022

Code	Code Description	Line(s) Involved	Issue	Recommendation(s)
90759	Hepatitis B vaccine (HepB), 3- antigen (S, Pre-S1, Pre-S2), 10	3 PREVENTION SERVICES WITH EVIDENCE OF EFFECTIVENESS	CPT 90759 was placed on the Excluded File when reviewed as a	Add 90759 to line 3
	mcg dosage, 3 dose schedule		new code in November, 2021 as it was not in the ACIP/CDC vaccine	Advise HSD to remove 90759 from the
			schedule. The OHA immunization sector staff have notified HERC staff that this vaccine	Excluded File
			(PreHevbrio) will be on the 2023 ACIP/CDC vaccine schedule as one	
			option for hepatitis B vaccination in adults. This code will need to	
			be opened by June 2022 as CDC plans to allow it for use at that	
47562	Laparoscopy, surgical;	641 GALLSTONES WITHOUT	Two cholecystectomy codes are	Add 47562 and 47563
7/302	cholecystectomy	CHOLECYSTITIS	missing from line 641.	to line 641
47563	Laparoscopy, surgical; cholecystectomy with cholangiography			

COVID-19 Related Codes May 2022

Issues:

- 1) CDC and CMS released 3 new ICD-10 codes for COVID vaccination status effective April 1, 2022
- 2) New HCPCS codes were released for COVID vaccinations done by dentists. There are also 7 HCPCS codes that are already in effect for dentists to use. Staff recommends all of these be placed on line 3. See: https://www.ada.org/-/media/project/ada-organization/ada/ada-org/files/publications/cdt/covid-19 vaccinationprocedurecodeguidance v2 2022mar.pdf?rev=b7ee2f99437246a8b6188d1f6e35 789f&hash=2A13A6E543730634A59B6EA1B6A456DC
- 3) New CPT codes were released in early May for the Sanofi-GSK vaccine, which will be effective with FDA EUA/approval. Additionally, a new CPT code was released for the booster dose of the new Pfizer tri-sucrose formulation vaccine
- 4) It has come to HERC staff attention that multiple HCPCS codes have been released for the administration of Evusheld (tixagevimab/cilgavimab), a monoclonal antibody injection for patients who are unable to receive a COVID-19 vaccine for medical reasons or whose immune systems may not respond robustly to COVID-19 vaccination. This therapy should be on line 3 as a preventive measure.
- 5) There are also HCPCS codes for bebtelovimab and bamlinivimab injections, which are monoclonal antibody products for treatment of COVID. Similar therapies are on line 399 INFLUENZA, COVID-19 AND OTHER NOVEL RESPIRATORY VIRAL ILLNESS

HERC staff recommendations:

1) Advise HSD to place the new COVID-related ICD-10 codes as shown below

ICD-10	Code Description	Recommended
Code		Placement
Z28.310	Unvaccinated for COVID-19	Informational File
Z28.311	Partially vaccinated for COVID-19	Informational File
Z28.39	Other under-immunization status [non-COVID vaccines]	Informational File

2) Add the following HCPCS codes to line 3 PREVENTION SERVICES WITH EVIDENCE OF EFFECTIVENESS or line 399 INFLUENZA, COVID-19 AND OTHER NOVEL RESPIRATORY VIRAL ILLNESS

HCPCS	Code Description	Recommended
Code		Placement
D1708	Pfizer-BioNTech Covid-19 vaccine administration – third dose	3
D1709	Pfizer-BioNTech Covid-19 vaccine administration – booster dose	3
D1710	Moderna Covid-19 vaccine administration – third dose	3
D1711	Moderna Covid-19 vaccine administration – booster dose	3
D1712	Janssen Covid-19 vaccine administration - booster dose	3

COVID-19 Related Codes May 2022

HCPCS	Code Description	Recommended
Code		Placement
D1713	Pfizer-BioNTech Covid-19 vaccine administration tris-sucrose pediatric – first dose	3
D1714	Pfizer-BioNTech Covid-19 vaccine administration tris-sucrose pediatric – second dose	3
M0220	Injection, tixagevimab and cilgavimab, for the pre-exposure prophylaxis only, for certain adults and pediatric individuals (12 years of age and older weighing at least 40kg) with no known sarscov-2 exposure, who either have moderate to severely compromised immune systems or for whom vaccination with any available covid-19 vaccine is not recommended due to a history of severe adverse reaction to a covid-19 vaccine(s) and/or covid-19 vaccine component(s), includes injection and post administration monitoring	3
M0221	Injection, tixagevimab and cilgavimab, for the pre-exposure prophylaxis only, for certain adults and pediatric individuals (12 years of age and older weighing at least 40kg) with no known sarscov-2 exposure, who either have moderate to severely compromised immune systems or for whom vaccination with any available covid-19 vaccine is not recommended due to a history of severe adverse reaction to a covid-19 vaccine(s) and/or covid-19 vaccine component(s), includes injection and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	3
Q0220	Injection, tixagevimab and cilgavimab, for the pre-exposure prophylaxis only, for certain adults and pediatric individuals (12 years of age and older weighing at least 40kg) with no known sarscov-2 exposure, who either have moderate to severely compromised immune systems or for whom vaccination with any available covid-19 vaccine is not recommended due to a history of severe adverse reaction to a covid-19 vaccine(s) and/or covid-19 vaccine component(s), 600 mg.	ANCILLARY PROCEDURES FILE
Q0221	Injection, tixagevimab and cilgavimab, for the pre-exposure prophylaxis only, for certain adults and pediatric individuals (12 years of age and older weighing at least 40kg) with no known sarscov-2 exposure, who either have moderate to severely compromised immune systems or for whom vaccination with any available covid-19 vaccine is not recommended due to a history of severe adverse reaction to a covid-19 vaccine(s) and/or covid-19 vaccine component(s), 300 mg.	ANCILLARY PROCEDURES FILE
Q0222	Injection, bebtelovimab, 175 mg	ANCILLARY PROCEDURES FILE
M0222	Intravenous injection, bebtelovimab, includes injection and post administration monitoring	399

COVID-19 Related Codes May 2022

HCPCS Code	Code Description	Recommended Placement
M0223	Intravenous injection, bebtelovimab, includes injection and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	399
Q0239	Injection, bamlanivimab, 700 mg	ANCILLARY PROCEDURES FILE
M0239	Intravenous infusion, bamlanivimab-xxxx, includes infusion and post administration monitoring	399

3) Add the following CPT codes to line 3 PREVENTION SERVICES WITH EVIDENCE OF EFFECTIVENESS

CPT Code	Code Description	Recommended Placement
91310	Severe acute respiratory syndrome coronavirus 2	3 PREVENTION SERVICES WITH
	(SARS-CoV-2) (coronavirus disease [COVID-19])	EVIDENCE OF EFFECTIVENESS
	vaccine, monovalent, preservative free, 5 mcg/0.5 mL	9
	dosage, adjuvant AS03 emulsion, for intramuscular	Pending FDA approval/EUA
	use	
0104A	Immunization administration by intramuscular	3 Pending FDA approval/EUA
	injection of severe acute respiratory syndrome	
	coronavirus 2 (SARS-CoV-2) (coronavirus disease	
	[COVID-19]) vaccine, monovalent, preservative free, 5	
	mcg/0.5 mL dosage, adjuvant AS03 emulsion, booster	
	dose	
0074A	Immunization administration by intramuscular	3 PREVENTION SERVICES WITH
	injection of severe acute respiratory syndrome	EVIDENCE OF EFFECTIVENESS
	coronavirus 2 (SARS-CoV- 2) (coronavirus disease	
	[COVID-19]) vaccine, mRNA-LNP, spike protein,	
	preservative free, 10 mcg/0.2 mL dosage, diluent	
	reconstituted, tris-sucrose formulation; booster dose	

May 2022

Items Discussed with Leadership with No Changes Recommended

- 1) Yttrium-90 treatment for colorectal cancer metastatic to the liver
 - a. In November 2018, this indication was reviewed and not found to have supporting evidence. An AHRQ review from 2012 that found it ineffective for this indication. Sirtex requested a re-review based on two new registry studies. HERC staff literature review found a recently-published RCT (Malcahy 2021) showing no survival benefit with Y-90 compared to conventional chemotherapy for this indication.
- 2) Breast tomosynthesis (3D Mammography)
 - a. Dr. Nathalie Johnson requested a re-review of breast tomosynthesis, which is on line 662/GN173. USPSTF is currently doing an in-depth review of screening modalities for breast cancer, including breast tomosynthesis. HERC staff advised waiting until the USPSTF review is completed before addressing this topic.
- 3) Equine psychotherapy
 - a. Psychotherapy is already covered and uses the same codes whether or not horses are used in the therapeutic setting. Staff review indicates insufficient evidence to support adding hippotherapy for any condition or modality.
- 4) Prescription therapeutic smart phone apps
 - a. This is a new area of medicine with a rapidly developing research base. The MED project is going to undertake a systematic review of this technology in the next year. Freespira requested a review of their app for opioid use treatment. HERC staff advised waiting until the MED review is completed before addressing this topic.
- 5) Smoking cessation and elective surgery
 - a. Dr. Amy Henninger requested that Guideline A4 be deleted because it
 disproportionately affects people of color and other groups with high smoking rates.
 This guideline was last reviewed in August 2021, and HERC considered deleting the
 guideline because of equity concerns and decided against deletion.

Temporary Urethral Stents May 2022

Plain Language Summary:

<u>Background:</u> Urethral stents are small tubes inserted into the urine duct (ureter) to treat or prevent a blockage that prevents the flow of urine from the kidney to the bladder. Last reviewed in 2015, temporary stents were not covered because there was not enough research to show they worked.

Should OHP cover this treatment? No, there is still not enough data to show benefit for temporary stent

Question: Should temporary urethral stents continue to be on line 662/GN173?

Question source: HERC staff

<u>Issue:</u> Temporary urethral stents were last reviewed in 2015 and found to be investigational. The CPT code for placement of such stents (CPT 53855) was placed on line 662/GN173. HERC staff recently identified a HCPCS code for this procedure that was overlooked in the 2015 review and continues to be Ancillary: HCPCS C9769 (Cystourethroscopy, with insertion of temporary prostatic implant/stent with fixation/anchor and incisional struts). Prior to adding this HCPCS code to the GN173 entry, staff rereviewed the evidence on this topic as it has been more than 5 years since last review.

Temporary stents are designed primarily for short-term use in the treatment of symptomatic benign prostatic hyperplasia (BPH), for a duration of 6 months to 3 years. Temporary stents are made of non-absorbable material, which prevents epithelial ingrowth and therefore allows easy removal. However, this may lead to unintended migration. Some temporary stents are biodegradable, so that they break down into small fragments, which are excreted through the urethra over time. Temporary stents are also very commonly used after urologic procedures, such as stone removal. Alternatives to temporary urethral stents include permanent stents and long-term Foley catherization. Both of these alternatives are covered on the BPH line or as an Ancillary therapy.

CPT 52282 (Cystourethroscopy, with insertion of permanent urethral stent) is on lines 214 CANCER OF KIDNEY AND OTHER URINARY ORGANS, 271 CANCER OF BLADDER AND URETER, 327 FUNCTIONAL AND MECHANICAL DISORDERS OF THE GENITOURINARY SYSTEM INCLUDING BLADDER OUTLET OBSTRUCTION, and 511 BENIGN NEOPLASM OF KIDNEY AND OTHER URINARY ORGANS.

Evidence

- 1) Sountoulides 2019, review of treatment for BPH
 - a. Overall, there is a diversity of stents design in terms of length, lumen diameter and material. In addition, the available studies are very small with short follow up, significant attrition rate and different definitions of efficacy. Therefore, there is a lack of robust data.
 - b. The only temporary stent reviewed with the Spanner stent, which has two studies (N=30, 43). In the study of 43 men, 63% of the patients had an unsatisfactory outcome due to immediate or delayed retention or elective stent removal because of severe symptoms.

Temporary Urethral Stents May 2022

Expert guidelines

- 1) American Urological Association 2021, Management of lower urinary tract symptoms attributed to benign prostatic hyperplasia
 - a. Does not mention temporary urethral stents in the treatment recommendations

Other payer policies

1) Aetna and Wellmark BCBS consider temporary urethral stents (CPT 53855 and HCPCS C9769) to be investigational

HERC staff recommendations:

- 1) Add HCPCS C9769 (Cystourethroscopy, with insertion of temporary prostatic implant/stent with fixation/anchor and incisional struts) to line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS
- 2) Modify GN173 as shown below

GUIDELINE NOTE 173, INTERVENTIONS THAT ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS FOR CERTAIN CONDITIONS

Line 662

The following Interventions are prioritized on Line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS:

Procedure	Intervention Description	Rationale	Last Review
Code			
53855	Temporary prostatic stents	Insufficient evidence of	October, 2015
<u>C9769</u>		effectiveness	
	5		May 2022

Plain Language Summary:

<u>Background</u>: Gastric neurostimulators are an implanted device used for diabetic stomach problems known as gastroparesis. Gastroparesis is a condition where the normal movement of food from the stomach to the small intestine is drastically slowed or has stopped. The devices have a limited form of FDA approval as a Humanitarian Use Device.

<u>Should OHP cover gastric neurostimulators?</u> Staff recommends OHP covering them. Though there are limited studies, evidence was found that for patients who were not able to take medication. Covering this would standardize the coordinated care organization review process for exceptions.

Question: Where should gastric neurostimulators be placed on the Prioritized List?

Question source: Holly Jo Hodges, CCO medical director

<u>Issue</u>: Gastric neurostimulators were removed from line 662/GN173 in October 2020 and placed in the Excluded File due to the fact that they did not have full FDA approval (humanitarian use approval only). Recently, a provider requested authorization to use a gastric neurostimulator and the relevant CCO requested HERC review of this technology.

Gastroparesis is a condition in which the normal movement of food from the stomach to the small intestine is drastically slowed or has stopped. This can lead to nausea and vomiting. Gastroparesis is frequently associated with diabetes. Gastric electrical stimulation (GES) is a treatment that sends weak electrical signals to the nerves and smooth muscles in the lower stomach. This treatment helps decrease nausea and vomiting caused by gastroparesis. A small battery-powered device is surgically placed in the skin in the lower belly area. Wires are then placed in the area to be stimulated.

Alternative treatments for gastroparesis include eating small meals, avoiding fizzy or high fiber foods, medications that stimulate stomach activity such as metoclopramide or domperidone, gastric or jejunostomy tube feeding, or parenteral nutrition.

Current FDA approval status: Humanitarian Device Exemption (HDE). This means that the FDA has found that the device "will not expose patients to an unreasonable or significant risk of illness or injury, and the probable benefit to health from using the device outweighs the risk of illness or injury." HUD devices are exempt from FDA effectiveness requirements and in many cases cannot be sold for profit.

Current Prioritized List status

The following codes are on the Excluded File:

CPT 43647 Laparoscopy, surgical; implantation or replacement of gastric neurostimulator electrodes, antrum

CPT 43648 Laparoscopy, surgical; revision or removal of gastric neurostimulator electrodes, antrum

CPT 43881 Implantation or replacement of gastric neurostimulator electrodes, antrum, open

CPT 43882 Revision or removal of gastric neurostimulator electrodes, antrum, open

HCPCS E0765 FDA approved nerve stimulator, with replaceable batteries, for treatment of nausea

ICD-10-CM K31.84 (Gastroparesis) is on line 529 DISORDERS OF FUNCTION OF STOMACH AND OTHER FUNCTIONAL DIGESTIVE DISORDERS

ICD-10-CM E11.43 (Type 2 diabetes mellitus with diabetic autonomic (poly)neuropathy) has the subdiagnosis of "Type 2 diabetes mellitus with diabetic gastroparesis" and is on line 27 TYPE 2 DIABETES MELLITUS

ICD-10-CM E10.43 (Type 1 diabetes mellitus with diabetic autonomic (poly)neuropathy) has the subdiagnosis of "Type 1 diabetes mellitus with diabetic gastroparesis" and is on line 8 TYPE 1 DIABETES MELLITUS

Evidence

- 1) NICE 2014 Gastroelectrical stimulation for gastroparesis
 - a. Current evidence on the efficacy and safety of gastric electrical stimulation for gastroparesis is adequate to support the use of this procedure
 - b. Efficacy
 - i. A meta-analysis of 4 studies including 169 patients with diabetic gastroparesis treated by gastroelectrical stimulation (part of a systematic review of 601 patients) reported improvement in total symptom severity score (weighted mean difference 8.96 [95% confidence interval {CI} 6.1 to 11.8]; p<0.00001) and a statistically significant improvement in gastric emptying at 4 hours (assessed using standardised radionucleotide scans of a solid meal: weighted mean difference 13.0 [95% CI 7.4 to 18.6]; p<0.00001). Subgroup analysis showed that the improvement was statistically significant in patients with diabetic or idiopathic gastroparesis but not in patients with post-surgical gastroparesis
 - ii. A meta-analysis of 3 studies including 58 patients with idiopathic gastroparesis treated by gastroelectrical stimulation reported improvement in total symptom severity score (weighted mean difference 7.5 [95% CI 5.4 to 9.7]; p<0.00001).
 - iii. In a systematic review of 364 patients, a meta-analysis of 4 studies including 75 patients with gastroparesis treated by gastroelectrical stimulation reported no statistically significant change in weight (weighted mean difference 3.7 [95% CI –0.2 to 7.6])
 - iv. In the systematic review of 364 patients, a meta-analysis of 8 studies including 184 patients with gastroparesis treated by gastroelectrical stimulation reported a reduction in need for nutritional support from 44% (96/216) of patients at baseline to 11% (21/184) at follow-up (odds ratio 5.5 [95% CI 2.8 to 11.1]; p<0.00001)
 - c. Safety
 - Death (within 30 days) was reported in 3% (2/72) of patients treated by gastroelectrical stimulation, due to small bowel infarction and heart failure, and 3% (1/31) of patients treated by gastrectomy, due to myocardial infarction, in a comparative case series of 103 patients
 - ii. Gastric perforation related to an episode of vomiting (2 months after the procedure) was reported in 1 patient in a case series of 17 patients. The device was removed and the perforation was repaired.
 - iii. Device removal was reported in 11% (24/221) of patients in a case series of 221 patients (timing ranged from 1–43 months after the procedure). Reasons were infection at the pulse generator or electrode sites (13 patients), lack of symptom improvement (6 patients), lead dislodgements (2 patients), small bowel obstruction caused by wires (1 patient), penetration of electrode into the lumen of the stomach (1 patient) and 'associated with peptic ulcer disease' (1 patient)
- 2) **Ducrotte 2020**, randomized crossover study of gastric electrical stimulation for reducing refractory vomiting
 - a. N=172 patients (133 with gastroparesis) with chronic vomiting related to diabetes or post-surgical
 - b. All patients has GES inserted, half were randomized to have immediate activation and half randomized to a 4 month delay in activation
 - vomiting was measured on a 5 point scale from 0 (several vomiting episodes a day) to 4 (no vomiting)

- d. vomiting scores were higher (improved) in the group with the device on (median score, 2) than the control group (median score, 1; P < .001), in diabetic and nondiabetic patients. Vomiting scores increased (improved) significantly when the device was ON in patients with delayed (P < .01) or normal gastric emptying (P = .05). Gastric emptying was not accelerated during the ON period compared with the OFF period. Having the GES turned on was not associated with increased quality of life</p>
- e. A total of 101 adverse events were reported in the study, with 45 therapy or device related events: abdominal wall pain at the implantation site (n = 28), infections at the abdominal pouch level (n = 16), hematoma (n = 1). In 3 cases, the device-related adverse events were serious enough to prompt device removal.

Expert guidelines

- American Gastroenterology Association 2022, practice update on the management of medically refractory gastroparesis
 - a. Clinicians can consider gastric electrical stimulation for gastroparesis patients with refractory/intractable nausea and vomiting who have failed standard therapy and are not on opioids and do not have abdominal pain as the predominant symptom
 - GES does improve refractory nausea and vomiting in some patients with gastroparesis and may improve glycemic control, nutritional status, and quality of life, while reducing hospitalizations and medication use

Other payer policies

1) Premera BCBS 2021

- a. Gastric electrical stimulation may be considered medically necessary in the treatment of chronic, intractable nausea and vomiting secondary to gastroparesis of diabetic or idiopathic etiology when ALL of the following criteria are met:
 - Significantly delayed gastric emptying as documented by standard scintigraphic imaging of solid food AND
 - ii. Patient is refractory or intolerant of prokinetic medications and antiemetic medications AND
 - iii. Patient's nutritional status is sufficiently low that total parenteral nutrition is likely to become medically necessary
- b. Gastric electrical stimulation is investigational for the treatment of obesity and all other indications

2) Cigna 2021

- a. Permanent gastric electrical stimulation (GES) or gastric pacing (e.g., Enterra™ Therapy) is considered medically necessary when provided in accordance with the Humanitarian Device Exemption (HDE) specifications of the U.S. Food and Drug Administration (FDA) for intractable nausea and vomiting secondary to gastroparesis with failure, contraindication, or intolerance of pharmaceutical therapy.
- b. Gastric electrical stimulation (GES) or gastric pacing for any other indication is considered experimental, investigational or unproven

HERC staff summary

Gastric electrical stimulation has a limited literature, and newer studies tend to find that GES reduces vomiting but does not improve quality of life. One highly trusted source (NICE) found the evidence sufficient for coverage. Most private insurance covers these devices for patients with diabetic or idiopathic (not post-surgical) gastroparesis who fail or are not able to tolerate medications.

Currently, idiopathic gastroparesis is on an uncovered line while diabetic gastroparesis is on the covered type 1 and type 2 diabetes lines. Addition of GES procedure codes and a new guideline would allow standardization of review in the exceptions process for CCOs for the uncovered diagnosis.

HERC staff recommendations:

- Add the following codes to lines 8 TYPE 1 DIABETES MELLITUS, 27 TYPE 2 DIABETES MELLITUS and 529 DISORDERS OF FUNCTION OF STOMACH AND OTHER FUNCTIONAL DIGESTIVE DISORDERS
 - a. CPT 43647 Laparoscopy, surgical; implantation or replacement of gastric neurostimulator electrodes, antrum
 - b. CPT 43648 Laparoscopy, surgical; revision or removal of gastric neurostimulator electrodes, antrum
 - c. CPT 43881 Implantation or replacement of gastric neurostimulator electrodes, antrum, open
 - d. CPT 43882 Revision or removal of gastric neurostimulator electrodes, antrum, open
 - e. HCPCS E0765 FDA approved nerve stimulator, with replaceable batteries, for treatment of nausea
- 2) Advise HSD to remove the above codes from the Excluded File
- 3) Adopt a new guideline for lines 8, 27, and 529 as shown below

GUIDELINE NOTE XXX GASTRIC ELECTRICAL STIMULATION

Line 8, 27,529

Gastric electrical stimulation (CPT 43657, 43648, 43881, 43882) is included on these lines only for pairing with diabetic gastroparesis (ICD-10-CM E10.43, E11.43) or idiopathic gastroparesis (ICD-10-CM K31.84) and only when ALL of the following criteria are met:

- The patient has intractable nausea and vomiting secondary to gastroparesis of diabetic or idiopathic etiology; AND
- 2) The patient is refractory or intolerant of prokinetic medications and antiemetic medications;
- 3) The patient is not on opioid medications; AND
- 4) The patient does not have abdominal pain as the predominant symptom.

Plain Language Summary:

<u>Background:</u> Coronary CT angiography (CCTA) is an imaging test that looks at the blood vessels called arteries that supply blood to the heart. The test looks for coronary artery disease (CAD) which can lead to heart attacks.

<u>Should OHP cover coronary CT angiography?</u> Staff recommends OHP cover this treatment because it is equally effective in detecting CAD as other tests.

Question: Should coverage be added for coronary CT angiography?

Question source: various CCO medical directors

<u>Issue</u>: Cardiac CT angiography (CCTA) is a test for evaluation of coronary artery disease (CAD). Noninvasive anatomic tests provide information on location and extent of blockage and include coronary CT angiography (CCTA) and cardiac magnetic resonance imaging (CMRI). Functional tests allow assessment of whether symptoms are correlated with narrowing leading to ischemic areas and generally include exercise (treadmill) electrocardiography (ECG), exercise/pharmacologic stress echocardiography, exercise/pharmacologic cardiac nuclear imaging with single-photon emission computed tomography (SPECT) or pharmacologic stress with positron emission tomography (PET).

Currently, CCTA is non-covered based on a 2013 coverage guidance. CCO medical directors have received requests for this procedure and would like an updated review and decision on coverage.

CMRI, treadmill stress test, pharmacologic stress echocardiography and SPECT are all currently covered as diagnostic tests for suspected CAD.

Summary of 2013 Coverage Guidance on CCTA

Coronary computed tomographic angiography may be useful to "rule out" obstructive CAD in ED patients with acute chest pain and normal ECGs and initial cardiac enzymes, and in outpatients with stable chest pain in a population with low to intermediate probability of obstructive CAD. Cost-effectiveness analyses show either that CCTA is comparable or less costly than other diagnostic strategies, although for the most part, they did not consider the economic consequences of the harms of radiation or further evaluation of incidental findings. However, understanding how CCTA would be used in a clinical practice setting, and whether the cost-effectiveness assumptions are applicable as it would be used in clinical practice, is unclear. Use in other patient populations is not recommended due to unacceptable false positive or false negative results. Use in asymptomatic patients has not been evaluated

"Blue box": Coronary Computed Tomography Angiography (CCTA) is not recommended for coverage.

HSD has also requested advice on the coverage of CPT 0501T-0504T (Noninvasive estimated coronary fractional flow reserve (FFR) derived from coronary computed tomography angiography data using computation fluid dynamics physiologic simulation software analysis of functional data to assess the severity of coronary artery disease). All private insurers cover these codes with CCTA.

Of note, coronary artery calcium scoring (CPT 75571) is a different procedure and is not included in this review.

Current Prioritized List status

CPT 75571 (Computed tomography, heart, without contrast material, with quantitative evaluation of coronary calcium) is on line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS:

CPT 75573 (Computed tomography, heart, with contrast material, for evaluation of cardiac structure and morphology in the setting of congenital heart disease (including 3D image postprocessing, assessment of left ventricular [LV] cardiac function, right ventricular [RV] structure and function and evaluation of vascular structures, if performed)) is on 20+ congenital heart disease lines

GUIDELINE NOTE 173, INTERVENTIONS THAT ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS FOR CERTAIN CONDITIONS

Line 662

The following Interventions are prioritized on Line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS:

Procedure	Intervention Description	Rationale	Last Review
Code			
75572	Computed tomography, heart,	Insufficient evidence of	December,
	with contrast material, for	effectiveness	<u>2009</u>
	evaluation of cardiac structure		
	and morphology		
75574	Computed tomography, heart	Insufficient evidence of	August, 2013
		benefit, unclear harms of	<u>Coverage</u>
		radiation exposure	<u>guidance</u>

Evidence

- 1) **Washington HTA 2021**, Noninvasive Cardiac Imaging for Coronary Artery Disease Re-review https://www.hca.wa.gov/assets/program/noninvasive-cardiac-imaging-final-report-2021019.pdf
 - a. Effectiveness
 - N=17 RCTs compared CCT with functional testing (stress nuclear or stress ECHO) for evaluation of suspected ACS in the emergency department or similar setting
 - 1. There is no clear difference in the frequency of later MI between CCTA and functional imaging tests (SOE Moderate).
 - 2. There was no association between CCTA and reduction in all-cause mortality compared with functional testing (SOE Moderate)
 - ii. N=19 RCTs (22,335 patients) comparing CCTA to functional testing for referral for invasive cardiac testing (ICA):
 - CCTA was associated with more frequent invasive coronary angiography (ICA) referral compared with functional testing (19 RCTs, 14.4 vs. 12 per 100 patients, pooled RR 1.25, 95% CI 1.09 to 1.47, I 2 =67%, RD 2.7, 95% CI 1 to 4 per 100, I 2 =59%) (SOE Moderate)
 - Referral for any additional noninvasive testing was not different between CCTA and functional testing groups across populations (17 RCTs, 7.2 versus 7.6 per 100, pooled RR 0.82, 95% CI 0.53 to 1.28, I 2 =83%) (SOE Low) or when populations were considered individually
 - iii. N=17 CTS (11,595 patients) comparing CCTA to functional testing for referral for any revascularization
 - CCTA was associated with more frequent revascularization (9.5 per 100 patients) compared with functional testing (7.1 per 100 patients): 19 RCTs, pooled RR 1.52, 95% CI 1.26 to 1.90, I 2 =66%, RD 2.4, 95% CI 1.4 to 3.3 per 100) (SOE Moderate)
 - CCTA was associate with higher frequency of PCI as a revascularization procedure compared with functional testing (12 RCTs, 8.2 vs. 6.0 per 100 patients, pooled RR 1.63, 95% CI 1.22 to 2.35, I 2 =74%, RD 2.4, 95% CI 1.3 to 3.6 per 100 patients) across populations (SOE Moderate).
 - iv. For hospitalization in stable outpatients there was no difference in hospitalization between CCTA and functional testing across four RCTs (SOE Moderate)
 - v. Subsequent ED visits: In patients with suspected ACS there was no difference in emergency department visits after index testing between CCTA and functional testing across studies, regardless of comparator, at 1 to 6.5 months (7 RCTs, pooled RR 0.84, 95% CI 0.66 to 1.06, I 2 =0%) or at ≥12 months (5 RCTs, pooled RR 1.06, 95% CI 0.93 to 1.56, I 2 =16%) (SOE High for both time frames)
 - vi. Medication: CCTA was not consistently associated with initiation of, discontinuation of or changes in medications and results for many mediations were mixed. Evidence is insufficient to draw firm conclusions about the impact of testing on medication use (SOE Insufficient)
 - b. Safety
 - i. Major or serious test-related adverse events/harms are rare for all modalities
 - ii. No major complications were observed across 3 RCTS at time of index test or within 24 hours. (SOE Low) The largest RCT in stable outpatients also reported no test-related hospitalization in the CCTA arm and 0.1% (5/4837) in the functional testing arm (SOE low).

- iii. Contrast-related events related to CCTA occurred in ≤3% of patients at time of index testing as reported in six RCTs and one case series. Transient creatinine elevation not requiring dialysis were reported in two trials as 0.2% and 1% (SOE Low) and a third RCT18 reported that no contrast-induced nephropathy occurred (SOE Insufficient). Mild contrast reaction occurred in 0.5% to 2.1% of patients across six RCTs (SOE Low)
- iv. Radiation from index tests: Across six RCTs comparing CCTA specifically with SPECT radiation exposure at index tended to be lower with CCTA (SOE Low). Five RCTs reported that CCTA was associated with a lower effective radiation dose for the index test; the sixth trial reported that CCTA was associated with slightly higher radiation (estimated difference 1.8 mSv). Rough estimates of difference between tests ranged from approximately 1.30 mSv to 11.9mSv. Stress echocardiography and ETT do not involve ionizing radiation (SOE Low).

c. Conclusions

- i. There is no clear difference between CCTA and functional imaging tests (i.e., stress nuclear testing, stress echocardiography) regarding impact on improving clinical outcomes (MI, all-cause mortality) in stable outpatients with suspected CAD or in patients with suspected ACS presenting to the ED or similar settings.
- ii. CCTA was associated with higher frequency of ICA referral and use of PCI for revascularization compared with functional imaging
- iii. While radiation exposure at time of index testing tended to be lower in CCTA recipients compared with SPECT (stress echo does not involve ionizing radiation), the evidence suggests that cumulative radiation may be higher with CCTA as an initial test
- iv. Incidental findings requiring follow-up in patients receiving CCTA are common and require additional resources.
- v. Definitive conclusions regarding cost-effectiveness of any of the noninvasive imaging tests are not possible in large part due to substantial heterogeneity across economic studies regarding testing strategies and test sequencing

Expert guidelines

- 1) AHA/ACC 2021 guideline for evaluation and diagnosis of chest pain
 - a. Intermediate or high pretest likelihood of CAD in patients younger than 65 years of age OR less obstructive CAD suspected: CTTA is the favored test (vs stress testing)
 - b. CCTA also recommended for use when prior functional study was inconclusive
 - c. CCTA recommended when there are anomalous coronary arteries or when the patient requires evaluation of the aorta or pulmonary arteries
 - d. CCTA is contraindicated when there is a contrast allergy, clinical instability, renal impairment, arrhythmia or contraindication to nitroglycerin

Other payer policies

1) Washington HTA 2021

- a. CCTA is a covered benefit with conditions for:
 - i. Symptomatic adult patients (≥18 years of age) at intermediate or high risk of CAD, or
 - ii. Adult patients with known CAD who have new or worsening symptoms.

2) Aetna 2022

- a. Aetna considers cardiac computed tomography (CT) angiography of the coronary arteries using 64-slice or greater medically necessary for the following indications:
 - Rule out obstructive coronary stenosis in symptomatic persons with a low or intermediate pre-test probability of coronary artery disease or atherosclerotic cardiovascular disease by Framingham risk scoring, Pooled Cohort Equations, or by American College of Cardiology (ACC) criteria,
 - ii. Rule out obstructive coronary stenosis in persons with a low or intermediate pre-test probability of coronary artery disease or atherosclerotic cardiovascular disease by Framingham risk scoring, Pooled Cohort Equations, or by American College of Cardiology (ACC) criteria (see Appendix) with a positive (i.e., greater than or equal to 1 mm ST segment depression) stress test.
 - iii. Evaluation of asymptomatic persons at an intermediate pre-test probability of coronary heart disease or atherosclerotic cardiovascular disease by Framingham risk scoring or Pooled Cohort Equations (see Appendix) who have an equivocal or uninterpretable exercise or pharmacological stress test or have resting electrocardiogram (ECG) changes (such as left bundle branch block (LBBB), pathologic q-waves, or right bundle branch block (RBBB) with left anterior fascicular block (LAFB) in which coronary artery disease (CAD) is a possible etiology. **Note**: Current guidelines from the American Heart Association recommend against routine stress testing for screening asymptomatic adults.
 - iv. Pre-operative assessment of persons scheduled to undergo 'high-risk" non-cardiac surgery, where an imaging stress test or invasive coronary angiography is being deferred unless absolutely necessary. The ACC defines high-risk surgery as emergent operations, especially in the elderly, aortic and other major vascular surgeries, peripheral vascular surgeries, and anticipated prolonged surgical procedures with large fluid shifts and/or blood loss involving the abdomen and thorax.
 - v. Pre-operative assessment for planned non-coronary cardiac surgeries including valvular heart disease, congenital heart disease, and pericardial disease, in lieu of cardiac catheterilzation as the initial imaging study, in persons with low or intermediate pretest risk of obstructive CAD.
 - vi. Detection and delineation of suspected coronary anomalies in young persons (less than 30 years of age) with suggestive symptoms (e.g., angina, syncope, arrhythmia, and exertional dyspnea without other known etiology of these symptoms in children and adults; dyspnea, tachypnea, wheezing, periods of pallor, irritability (episodic crying), diaphoresis, poor feeding and failure to thrive in infants).
 - vii. Calculation of fractional flow reserve (HeartFlow FFR_{CT}) for persons who have a coronary CTA that has shown coronary artery disease of uncertain functional significance, or is non-diagnostic.

3) Cigna 2011

- a. CIGNA covers 64-slice or greater multidetector-row computed tomography angiography (CTA) as medically necessary as an adjunct to other testing for ANY of the following indications:
 - i. evaluation of chest pain in an individual with a very low, low, or intermediate pre-test probability of coronary artery disease1 (CAD) when the individual cannot perform or has a contraindication to exercise and chemical stress testing

- (i.e. exercise treadmill stress test, stress echo, and nuclear stress test [i.e., myocardial perfusion imaging])
- ii. exclusion of CAD in an individual with a low or very low pre-test probability of CAD when recent stress test results (i.e., exercise treadmill, stress echo, or nuclear stress test [i.e., myocardial perfusion imaging]) are uninterpretable, equivocal, or there is a suspicion that the results are falsely positive
- iii. exclusion of CAD in an individual with an intermediate pre-test probability of CAD when recent stress test results (i.e., exercise treadmill, stress echo, or nuclear stress test [i.e., myocardial perfusion imaging]) are uninterpretable or equivocal, AND CTA will be performed in lieu of an angiography.
- iv. exclusion of CAD in a symptomatic individual (e.g., acute chest pain in an emergency department setting), and the individual has an intermediate pre-test probability of CAD, and there are no changes noted on the ECG and serial enzymes are negative
- v. evaluation of suspected or known coronary artery anomalies associated with congenital conditions
- vi. for morphologic evaluation of the coronary arteries in an individual with dilated cardiomyopathy or new onset heart failure, when ischemia is the suspected etiology and cardiac catheterization and/or nuclear stress test (i.e., myocardial perfusion imaging) have not been performed
- vii. pre-operative assessment of coronary arteries in an individual undergoing repair of aortic dissection, aortic aneurysm repair or valvular surgery AND CTA will be performed in lieu of an angiography
- viii. post-coronary artery bypass grafting (CABG) when BOTH of the following criteria are met:
 - 1. repeat intervention is being considered
 - 2. recent coronary angiography has been completed but additional information is needed before a treatment decision can be made
- b. CIGNA does not cover multidetector-row computed tomography angiography (CTA) for any other indication, including but not limited to those listed below, because it is considered experimental, investigational or unproven:
 - i. evaluation of chest pain in an intermediate or high pre-test probability of CAD individual when recent stress test result (i.e., exercise treadmill, stress echo, or nuclear stress test [i.e., myocardial perfusion imaging]) are either clearly positive or unequivocally negative
 - ii. screening for CAD in an asymptomatic individual
 - iii. post-revascularization procedure (e.g., percutaneous coronary intervention, coronary artery bypass grafting surgery), including evaluation of bypass grafts, coronary anatomy or evaluation for in-stent restenosis except when an individual is post-coronary artery bypass grafting (CABG), repeat intervention is being considered but additional information is required following completion of recent coronary angiography

Coronary CT Angiography (CCTA)

Expert input:

Dr. Abigail Khan, OHSU cardiology

I agree with your assessment that CCTA has a strong body of evidence supporting that it is not inferior to stress echo or SPECT and its diagnostic use is now supported by national guidelines. I do think we should cover CCTA given its use is quickly becoming standard of care in certain settings and there are situations in which it is the preferred test over stress testing. Importantly, it can also allow for more expedited assessment of symptomatic patients than stress testing, which is more labor intensive and has more scheduling complexities in real world practice.

...after consulting with a CT expert my feelings are that we do not need additional guidelines for CCTA if they are not in place for stress testing. There is actually a stronger body of literature supporting CCTA than stress testing at this point. CCTA is unlikely to be overutilized any more than stress testing (and is actually a cheaper test). The complexity is patients who have had prior stents in which the use of CCTA is not straightforward. That said I'm not sure if an OHP coverage guideline is the right way to go about addressing this issue. I suspect that providers will mostly send these patients for other testing (stress or cath) anyways, so we are not talking about a huge population of affected patients.

Dr. David Saenger, Cardiologist

Coronary CTA is becoming the standard first best test to evaluate patients for chest pain. This is both in the acute setting and in outpatient clinic. There are, of course, limitations and contraindications to CCTA. We have an algorithm that we wrote at OHVI that I can share. Bottom line is that relative contraindications are obesity (BMI > 35), tachycardia (especially a fib with RVR), prior PCI (except stents in large vessels like the left main are ok), and renal failure. There are contraindications to stress testing too, of course. So I don't think we need to have strict rules for ordering the test.

Coronary CT Angiography (CCTA)

HERC staff summary

CCTA is equally effective on improving clinical outcomes (MI and all-cause mortality) compared to stress ECHO or SPECT in patients suspected of having coronary artery disease (CAD). CCTA is associated with higher frequency of referral for invasive coronary artery catheterization and has a higher rate of incidental findings than other tests. The amount of radiation exposure from CCTA does not appear to be significant with newer protocols. The AHA/ACA guideline recommends CCTA as the test of choice for evaluation of patients under age 65 with intermediate or high suspicion of CAD or who have an indeterminate prior functional study. All private insurers surveyed cover this test, with different criteria. Experts consulted did not feel that there was a need for a guideline for this procedure.

HERC staff recommendations:

- 1) Remove CPT 75572 and 75574 (CT heart) from line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS
 - a. Advise HSD to add CPT 75572 and 75574 to the Diagnostic Procedures File
 - b. Remove the entries for CPT 75572 and 75574 from GN173
 - c. Advise HSD to add CPT 0501T-0504T (Noninvasive estimated coronary fractional flow reserve (FFR) derived from coronary computed tomography angiography data using computation fluid dynamics physiologic simulation software analysis of functional data to assess the severity of coronary artery disease) to the Diagnostic Procedures File

GUIDELINE NOTE 173, INTERVENTIONS THAT ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS FOR CERTAIN CONDITIONS

Line 662

The following Interventions are prioritized on Line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS:

Procedure Code	Intervention Description	Rationale	Last Review
75572	Computed tomography, heart, with contrast material, for evaluation of cardiac structure and morphology	Insufficient evidence of effectiveness	December, 2009
75574	Computed tomography, heart	Insufficient evidence of benefit, unclear harms of radiation exposure	August, 2013 Coverage guidance

Rhinophyma Treatment

Plain Language Summary:

<u>Background:</u> Rhinophyma shaving is removing thickened skin from the nose due to a skin condition. Rarely it can cause problems with function such as breathing or infections. Most cases are considered cosmetic, which would only improve a person's appearance, not function. If a case was severe, it would be coded as a sinus condition and covered.

<u>Should OHP cover rhinophyma shaving?</u> Staff recommends OHP not cover this treatment because most cases are considered cosmetic.

Question: Should the current placement of surgical planing for rhinophyma be changed?

Question source: HSD Medical Management Committee

<u>Issue</u>: Rhinophyma is soft tissue and sebaceous hyperplasia of the nose which is caused by severe rosacea. Rhinophyma responds to electrosurgery, laser excision, and surgical debulking. Because telangiectasias and rhinophyma do not cause functional limitations, their treatment is considered cosmetic by most insurance carriers.

Currently, surgical planing for rhinophyma is on the covered line for chronic sinusitis and on two uncovered lines for disorders of the nasal cavity. It is not on the rosacea line. MMC recently received a request for surgical planing for a patient which sinus issues and requested that the HERC review the current coverage of this procedure. If the rhinophyma was severe enough to cause nasal obstruction, the condition could be coded with ICD-10 J34.89 (Other specified disorders of nose and nasal sinuses) which is on line 506 NASAL POLYPS, OTHER DISORDERS OF NASAL CAVITY AND SINUSES

Current Prioritized List status

CPT 30120 (Excision or surgical planing of skin of nose for rhinophyma) is on lines 466 CHRONIC SINUSITIS 506 NASAL POLYPS, OTHER DISORDERS OF NASAL CAVITY AND SINUSES 525 BENIGN NEOPLASM OF NASAL CAVITIES, MIDDLE EAR AND ACCESSORY SINUSES

Rosacea is on line 522 ROSACEA; MILD/MODERATE ACNE

Other payer policies

- 1) Aetna 2021
 - a. Aetna considers excision or shaving of rhinophyma medically necessary for the treatment of bleeding or infection refractory to medical therapy (i.e., the need for repeated cautery of bleeding telangiectasias or frequent courses of antibiotics for pustular eruptions)
- 2) Anthem BCBS 2021
 - a. Excision or shaving of the rhinophyma is considered **medically necessary** when **both** of the following criteria are met:
 - the medical record documentation includes evidence of bleeding or infection; and
 - ii. the procedure can be reasonably expected to improve functional impairment as a result of bleeding or infection.

Rhinophyma Treatment

b. Excision or shaving of the rhinophyma is considered **cosmetic and not medically necessary** when the medically necessary criteria in this section are not met.

Rhinophyma Treatment

HERC staff recommendations

- 1) Remove CPT 30120 (Excision or surgical planing of skin of nose for rhinophyma) from the following lines
 - a. 466 CHRONIC SINUSITIS
 - b. 506 NASAL POLYPS, OTHER DISORDERS OF NASAL CAVITY AND SINUSES
 - c. 525 BENIGN NEOPLASM OF NASAL CAVITIES, MIDDLE EAR AND ACCESSORY SINUSES
- 2) Add CPT 30120 to line 522 ROSACEA; MILD/MODERATE ACNE

Spinal Cord Stimulation for Diabetic Neuropathy

Plain Language Summary:

<u>Background:</u> Spinal cord stimulation is a treatment for lower body nerve issues for people with nerve problems. The studies showed that this device might help patients with nerve problems caused by diabetes, but there was a high rate of problems such as infections and equipment failure.

<u>Should OHP cover this treatment?</u> Staff recommends OHP not cover this treatment because there is limited evidence which shows harms outweigh benefits.

Question: Should spinal cord stimulation be added as a treatment for diabetic neuropathy?

Question source: Medtronic

<u>Issue</u>: On January 21, 2022, Medtronic received U.S. Food and Drug Administration approval of the Intellis™ rechargeable neurostimulator and Vanta™ recharge-free neurostimulator for the treatment of chronic pain associated with diabetic peripheral neuropathy (DPN) of the lower extremities. Medtronic requested review of neurostimulators for pairing with diabetic peripheral neuropathy.

Current Prioritized List Status

ICD-10-CM E10.4 family (Type 1 diabetes mellitus with diabetic neuropathy) is on line 8 TYPE 1 DIABETES MELLITUS

ICD-10-CM E11.4 family (Type 2 diabetes mellitus with diabetic neuropathy) is on line 2 TYPE 2 DIABETES MELLITUS

CPT 63650 (Percutaneous implantation of neurostimulator electrode array, epidural), 63655 (Laminectomy for implantation of neurostimulator electrodes, plate/paddle, epidural) and 63663-63688 (Revision or replacement of spinal neurostimulator equipment) are on lines 292 NEUROLOGICAL DYSFUNCTION IN POSTURE AND MOVEMENT CAUSED BY CHRONIC CONDITIONS, 346 CONDITIONS OF THE BACK AND SPINE WITH URGENT SURGICAL INDICATIONS and 530 CONDITIONS OF THE BACK AND SPINE WITHOUT URGENT SURGICAL INDICATIONS

Evidence

- 1) Strand 2022, systematic review of spinal cord stimulation for diabetic peripheral neuropathy
 - a. Both authors with conflicts of interest
 - b. Reported complications:
 - i. Lead migration in 0-30%
 - ii. Revision of leads in 0-30%
 - iii. Infection in 0-20%
 - iv. Explant in 0-17%
 - c. RCTs of low frequency spinal cord stimulation
 - i. De Vos et al
 - 1. N=60 patients (randomized 2:1 to best medical treatment or spinal cord stimulation)
 - a. N=37 with spinal cord simulation
 - 2. Outcomes at six months for 36 participant receiving SCS and 18 control participant showed that pain scores decreased by 55% with LF-SCS, while there was no decrease in mean pain among control participant.

Spinal Cord Stimulation for Diabetic Neuropathy

The responder rate was 69% (25 of 36) among participant treated with SCS, while 6% (1 of 18) of controls were responders after six months of treatment

- ii. Slangen et al
 - 1. N=36 patients (N=22 randomized to spinal cord stimulation)
 - 2. Daytime mean pain NRS scores decreased by 3.1 points after six months of stimulation in the SCS group but did not change in the BMT group (P < .001), and corresponding nighttime pain scores declined 2.4 points in the SCS treatment arm and 0.9 points in the BMT arm (P < .003). The sixmonth daytime responder rate (≥50% pain relief) was 9 out of 16 (56%) in the per protocol (PP) population of the SCS arm and 0 out of 14 (0%) in the BMT arm (P < .001); while nighttime responder rates were 8 out of 16 (50%) in the SCS group and 1 out of 14 (7%) in the BMT group (P < .05). At the last follow-up, 12 of 16 participant (75%) treated with SCS and followed for six months reported a PGIC for pain score ≥6, demonstrating "much" or "very much" improvement, while none of the participant receiving BMT did (P < .001)</p>
- d. Conclusions: There is currently a substantial unmet need for safe and effective treatments for PDN. Many patients with PDN do not benefit from pharmacotherapies in current use and are candidates for treatment with neuromodulation. Conventional LF-SCS...are supported by high-quality evidence from RCTs and prospective studies
- 2) Henson 2021, systematic review of spinal cord stimulation for diabetic peripheral neuropathy
 - a. N=14 prospective studies
 - b. N=2 RCTs
 - i. De Vos et al
 - ii. Slangen et al
 - c. N=11 prospective cohort studies
 - d. Conclusion: Based on our analysis of the available evidence, there is moderate-quality evidence for the safety and efficacy of spinal cord stimulation for painful diabetic neuropathy. However, further high-quality research, including a large-scale randomized controlled trial is warranted
- 3) **Peterson 2021**: RCT of high frequency spinal cord stimulation for patients with painful diabetic neuropathy
 - a. SENZA trial
 - b. All authors reported conflicts of interest; study funded by industry
 - c. N=216 patients with peripheral neuropathy refractory to at least 2 classes of medications and lower limb pain with an initial VAS score of 5 or higher
 - i. N=103 assigned to conventional medical management (CMM)
 - 1. Data reported for 93 by 6 month follow up
 - ii. N=113 assigned to CMM plus spinal cord stimulator (SCC plus CMM)
 - 1. Data reported for 87 by the 6 month follow up
 - d. In the CMM group, 5 of 94 patients (5%) met the composite primary end point of 50% or more pain relief using the VAS without observed deterioration on neurological examination compared with 75 of 95 in the 10-kHz SCS plus CMM group (79%; difference, 73.6%; 95% CI, 64.2-83.0; P < .001).
 - e. At 6-month follow-up, there was no change in mean pain VAS scores for the CMM group, with a baseline mean of 7.0 cm (95% CI, 6.7-7.3) and a 6-month mean of 6.9 cm

Spinal Cord Stimulation for Diabetic Neuropathy

- (95% CI, 6.5-7.3); however, lower limb pain VAS scores decreased by a mean of 76.3% (95% CI, 70.8-81.8) for the implanted group.
- f. At 6 month follow up, there was no change for [health related quality of life score] in the CMM group but a mean 16-point (95% CI, 11.3-20.5) improvement for those in the 10-kHz SCS plus CMM group (P < .001)
- g. Adverse events
 - i. None in CMM group
 - ii. there were 18 AEs reported among 14 patients in the 10-kHz SCS plus CMM group: 3 study-related AEs for infection, 2 for wound dehiscence, and 1 for impaired healing among 5 of 90 patients (6%). Of 90 total implanted patients, 2 (2%) required explant.
- h. Conclusion: Patients with PDN refractory to best available treatments can be safely and effectively treated with high-frequency (10- kHz) SCS.

Other payer policies:

- 1) Aetna 2022 does not consider spinal cord stimulation medically necessary for diabetic peripheral neuropathy
- 2) Cigna 2020 did not list diabetic peripheral neuropathy as a covered indication for spinal cord stimulation

HERC staff summary

The evidence on spinal cord stimulation for diabetic peripheral neuropathy consists of 3 RCTs with a total of 146 patients in the SCS groups. These RCTs showed consistent improvement in pain relief for SCS vs medical management alone. However, there was a significant rate of adverse events, including infection and equipment breakage/failure. Diabetic neuropathy is a very common condition and RCTs with large populations are feasible.

HERC staff recommendation:

1) Make no change in non-pairing of spinal cord stimulators with diabetic peripheral neuropathy.

Subacromial Spacers

Plain Language Summary:

<u>Background:</u> Where should shoulder surgery with balloon implants to treat conditions such as severely torn rotator cuffs be prioritized? This is a new treatment and is not well-studied. No other insurer appears to cover this treatment.

<u>Should OHP cover this treatment?</u> Staff recommends OHP not cover this treatment because there isn't good evidence to cover it.

Question: Where should arthroscopy with implantation of a subacromial spacer be prioritized?

Question source: HERC staff

<u>Issue</u>: CMS released a new HCPCS code effective 4/4/22 regarding placement of subacromial spacers. This HCPCS code was adopted "to describe the implantation of a saline-filled balloon for the shoulder to treat irreparably torn rotator cuff tendons." There is no similar HCPCS or CPT code currently available.

<u>Code</u>: HCPCS C9781 (Arthroscopy, shoulder, surgical; with implantation of subacromial spacer (e.g., balloon), includes debridement (e.g., limited or extensive), subacromial decompression, acromioplasty, and biceps tenodesis when performed)

Evidence:

- 1) **Johns 2020**, systematic review of implantable subacromial balloon spacers for patients with massive irreparable rotator cuff tears
 - a. N=19 studies. 13 studies described as non-randomized clinical trials
 - i. No comparator groups in any trial
 - ii. N=337 patients
 - iii. All studies included patients with persistent symptoms for a minimum of 3 to 6 months with failure of conservative treatment, including activity modification, treatment with non-steroidal anti-inflammatory medications, intra-articular corticosteroid injections, and/or physical therapy
 - b. Patients treated with a subacromial balloon spacer demonstrated overall improvement in postoperative TCS compared with preoperative TCS based on the data from 11 studies (preoperative range: 22.5-41.8; postoperative range: 51.4-72.3). All of these studies reported statistically significant improvement in TCS from preoperative to postoperative measurements at all short-term and long-term follow-up timepoints
 - c. There was significant improvement of shoulder abduction (preoperative range: 70-113 degrees; postoperative range: 110-165 degrees), shoulder flexion (preoperative range: 80-130 degrees; postoperative range: 106.5-161 degrees), and external rotation (preoperative range: 25-44.5 degrees; postoperative range: 35-63.7 degrees) from preoperative to postoperative ROM after placement of the subacromial balloon spacer
 - d. Complications: 1 of 350 (0.29%) patients experienced a transient forearm dysesthesia in the lateral cutaneous nerve of the forearm after implantation of the subacromial balloon spacer. A total of 1 of 350 (0.29%) procedures was complicated by superficial wound infection at the surgical site, which resolved after a course of antibiotics per orem, and 1 of 350 (0.29%) procedures was complicated by a deep wound infection, which was culture-negative and treated with 1 week of intravenous antibiotics followed by 2 weeks of per orem antibiotics. One patient with an increasingly painful shoulder

Subacromial Spacers

was found to have remnants of a deflated InSpace Balloon with transformation to scar tissue in the subacromial space on MRI. In total, 11 of 350 (3.14%) of procedures required reoperation, including 5 (1.42%) for InSpace Balloon migration, 1 (0.29%) for synovitis, and 6 (1.71%) underwent reverse total shoulder arthroplasty due to absence of clinical improvement or worsening of symptoms at various postoperative follow-up durations ranging from 6 weeks to 16 months. A total of 4 patients were found to have synovitis on MRI at 3 years post-implantation, and there was 1 shoulder dislocation at 6 weeks postoperative secondary to an acute trauma

- e. Conclusions: Existing literature of subacromial balloon spacers has high risk of bias, lack of appropriate control, and low level of evidence. A qualitative synthesis indicates that subacromial balloon spacer implantation in patients with massive irreparable rotator cuff tears is cost-effective and leads to improved function (TCS and OSS) and ROM
- 2) **Stewart 2019**, systematic review of subacromial balloon spacer implantation for massive and irreparable rotator cuff tears
 - a. N=12 studies (284 patients)
 - i. 10 case series, 1 prospective cohort study, and 1 retrospective cohort study
 - b. In 2 studies, strength was not statistically significant. In 1 study, statistical significance was not reported for any subscale value (pain, ADL, ROM, strength)
 - c. 4 studies observed increases in active abduction; however, results from 1 of these studies were not statistically significant.
 - d. Of the 3 studies that reported active anterior elevation, all observed increases; however, only the results from 1 study were statistically significant (P = .00000001)
 - e. Complications occurred in 2.1% of patients. These complications included transient neurapraxia of the lateral antebrachial cutaneous nerve in 1 patient, superficial wound infection in 1 patient, deep wound infection in 1 patient, and balloon migration in 3 patients
 - f. This systematic review of the existing literature suggests that subacromial balloon spacer placement is a minimally invasive, technically simple procedure with low rates of perioperative complications and favorable patient reported outcomes at limited short-term follow up
 - g. Further prospective randomized or comparative studies are warranted to ascertain clinical outcomes of subacromial balloon spacer in the management of massive and irreparable RCTs

Other payer policies: None found

Subacromial Spacers

HERC staff summary

Arthroplasty with subacromial balloon spacer placement is an emerging technology with very limited evidence of effectiveness. Of note, Metcalfe (2021) has published a protocol for an RCT of this technology (START:REACTS study).

HERC staff recommendation:

1) Place HCPCS C9781 Arthroscopy, shoulder, surgical; with implantation of subacromial spacer (e.g., balloon), includes debridement (e.g., limited or extensive), subacromial decompression, acromioplasty, and biceps tenodesis when performed) on line 662/GN173 as shown below

GUIDELINE NOTE 173, INTERVENTIONS THAT ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS FOR CERTAIN CONDITIONS

The following Interventions are prioritized on Line 662 CONDITIONS FOR WHICH CERTAIN INTERVENTIONS ARE UNPROVEN, HAVE NO CLINICALLY IMPORTANT BENEFIT OR HAVE HARMS THAT OUTWEIGH BENEFITS:

Procedure Code	Intervention Description	Rationale	Last Review
<u>C9781</u>	Arthroscopy, shoulder, surgical; with implantation of subacromial	Insufficient evidence of effectiveness	May 2022
	spacer (e.g., balloon)	<u>Grenedaess</u>	

Plain Language Summary:

<u>Background:</u> Erythropoietin is a drug that is used to treat low blood count caused by kidney disease. Currently, it is covered only for the worst stage of chronic kidney failure with certain conditions.

<u>Should OHP cover this drug for lower stages of chronic kidney disease?</u> Staff recommends OHP expand coverage of this drug because the Oregon Health Authority's Pharmacy & Therapeutics Committee and the Food and Drug Administration (FDA) allow use of this drug to treat kidney disease of all stages. Other insurance companies also cover this drug for all stages of kidney disease when needed.

<u>Question:</u> Should several changes be made to the Prioritized List to add coverage for erythropoietin for non-end stage renal disease?

Question source: Jennifer Smith, PharmD, Pharmacy Manager, Providence Health Plan

<u>Issue</u>: Chronic renal failure with a hemoglobin level <10 was added as an indication for erythropoiesis-stimulating agents in Guideline Note 7 in 2012. The current GN 7 only applies to Line 59 END STAGE RENAL DISEASE. Line 59 only includes ICD-10-CM N18.5 (Chronic kidney disease, stage 5) and N18.6 (End stage renal disease). Earlier stages of chronic kidney disease (ICD-10-CM N18.1-N18.4 and N18.9) are on line 339 CHRONIC KIDNEY DISEASE, which is not referenced in the guideline.

The FDA has approved erythropoietin for all stages of chronic kidney disease with a low hemoglobin level. According to P&T, all ICD-10 codes above the funding line (i.e., all N18 series codes) are being funded for erythropoietin currently in their PA process. However, P&T staff report that as a physician-administered drug, there are very few PA requirements.

Additionally, Dr. Smith requested that ICD-10-CM D63.1 (Anemia in chronic kidney disease) be a code allowable to pair with erythropoietin. However, on further research, HERC staff has determined that D63.1 has only one sub-diagnosis and that is "Erythropoietin resistant anemia" and therefore would not be appropriate to use with erythropoietin.

FDA Epogen labeling 7/2018:

1 INDICATIONS AND USAGE

1.1 Anemia Due to Chronic Kidney Disease Epogen is indicated for the treatment of anemia due to chronic kidney disease (CKD), including patients on dialysis and not on dialysis to decrease the need for red blood cell (RBC) transfusion.

For adult patients with CKD on dialysis:

- Initiate Epogen treatment when the hemoglobin level is less than 10 g/dL.
- If the hemoglobin level approaches or exceeds 11 g/dL, reduce or interrupt the dose of Epogen.
- The recommended starting dose for adult patients is 50 to 100 Units/kg 3 times weekly intravenously or subcutaneously. The intravenous route is recommended for patients on hemodialysis.

For adult patients with CKD not on dialysis:

- Consider initiating Epogen treatment only when the hemoglobin level is less than 10 g/dL and the following considerations apply:
 - o The rate of hemoglobin decline indicates the likelihood of requiring a RBC transfusion and,

- o Reducing the risk of alloimmunization and/or other RBC transfusion-related risks is a goal
- If the hemoglobin level exceeds 10 g/dL, reduce or interrupt the dose of Epogen, and use the lowest dose of Epogen sufficient to reduce the need for RBC transfusions.
- The recommended starting dose for adult patients is 50 to 100 Units/kg 3 times weekly intravenously or subcutaneously.

For pediatric patients with CKD:

- Initiate Epogen treatment only when the hemoglobin level is less than 10 g/dL.
- If the hemoglobin level approaches or exceeds 12 g/dL, reduce or interrupt the dose of Epogen.
- The recommended starting dose for pediatric patients (ages 1 month or older) is 50 Units/kg 3 times weekly intravenously or subcutaneously.

Evidence

- 1) Stauffer 2014, prevalence of anemia is CKD in the US
 - a. NHANES survey of 12,077 adults
 - b. Treatments for anemia include iron supplementation and erythropoietin stimulating agents
 - c. The prevalence of anemia increased with stage of CKD
 - i. 8.4% at stage 1
 - ii. 12.2% at stage 2
 - iii. 17.4% at stage 3
 - iv. 50.3% at stage 4
 - v. 53.4% at stage 5.
 - d. A total of 22.8% of CKD patients with anemia reported being treated for anemia within the previous 3 months: 14.6% of patients at CKD stages 1–2 and 26.4% of patients at stages 3–4. 430.5 of stage 5 patients reported treatment

Other payer policies

- 1) Aetna 2022
 - a. Allows all levels of chronic kidney disease (N18.1-N18.9) for use with epo administration
- 2) Cigna 2020
 - a. Chronic Kidney Disease Anemia and EITHER of the following:
 - i. Individual is on dialysis.
 - ii. Individual is not on dialysis and EITHER of the following:
 - 1. Initial treatment and ONE of the following:
 - a. The patient is ≥ 18 years of age with a hemoglobin < 10.0 g/dL OR
 - b. The patient is < 18 years of age with a hemoglobin \leq 11.0 g/dL
 - b. Established treatment and ONE of the following:
 - i. The patient is ≥ 18 years of age with a hemoglobin < 11.5 g/dL; OR
 - ii. The patient is < 18 years of age with a hemoglobin ≤ 12.0 g/dL
 - c. Note: no coding was included in their coverage document

Current utilization

Per P&T staff, the following codes were used in the past 5 months for Epo administration

Patients	Claims	CodeFFSPHP	${\sf CodeDiagCondMedIDet}$	Diagnosis description
1108	4641	CCO	N18.6	End stage renal disease
86	298	CCO	N18.4	CKD stage 4
23	87	CCO	N18.5	CKD stage 5
25	77	CCO	N18.30	CKD stage 3 unspecified
29	76	CCO	N18.32	CKD stage 3b
15	36	CCO	N18.31	CKD stage 3a
9	20	CCO	N18.9	CKD unspecified
3	3	CCO	N18.2	CKD stage 2
84	260	FFS	N18.6	End stage renal disease
17	62	FFS	N18.4	CKD stage 4
5	13	FFS	N18.5	CKD stage 5
7	13	FFS	N18.32	CKD stage 3b
5	12	FFS	N18.31	CKD stage 3a
3	9	FFS	N18.9	CKD unspecified
1	4	FFS	N18.30	CKD stage 3 unspecified

HERC staff summary

P&T criteria and FDA guidelines allow treatment with erythropoietin at all stages of CKD. Currently, only stage 5 (end stage) renal disease is paired with erythropoietin in GN7. Private payers are reimbursing for lower stages of CKD to be treated with erythropoietin if the patient meets anemia requirements. As P&T is allowing all N18 codes for erythropoietin use, HERC staff recommends that GN7 be added to line 339 CHRONIC KIDNEY DISEASE so that the anemia criteria in the guideline will apply to all the N18 codes, as these codes appear to be widely used for erythropoietin administration. This will also allow erythropoietin use in less severe stages of renal disease per FDA criteria.

HERC staff recommendations:

- 1) Add Guideline Note 7 to Line 339 CHRONIC KIDNEY DISEASE
 - a. Will ensure that the N18 code series is included in and regulated by this guideline
- 2) Modify GN7 as shown below

GUIDELINE NOTE 7, ERYTHROPOIESIS-STIMULATING AGENT (ESA) GUIDELINE

Lines 12,59,92,94,111-115,125,133,135,157,158,161,163,179,191,199,200,208,210,214,215,217, 229,234,237,238,258-262,271,276,286-288,294,295,314-316,329,339,396,397,401,419,435,559,593

- A) Indicated for anemia (Hgb < 10gm/dl or Hct < 30%) induced by cancer chemotherapy given within the previous 8 weeks or in the setting of myelodysplasia.
 - 1) Reassessment should be made after 8 weeks of treatment. If no response, treatment should be discontinued. If response is demonstrated, ESAs should be discontinued once the hemoglobin level reaches 10, unless a lower hemoglobin level is sufficient to avoid the need for red blood cell (RBC) transfusion.
- B) Indicated for anemia (Hgb < 10gm/dl or HCT < 30%) associated with HIV/AIDS.
 - 1) An endogenous erythropoietin level < 500 IU/L is required for treatment, and patient may not be receiving zidovudine (AZT) > 4200 mg/week.
 - 2) Reassessment should be made after 8 weeks. If no response, treatment should be discontinued. If response is demonstrated, the lowest ESA dose sufficient to reduce the need for RBC transfusions should be used, and the Hgb should not exceed 11gm/dl.
- c) Indicated for anemia (Hgb < 10 gm/dl or HCT <30%) associated with chronic renal <u>disease</u> failure, with or without dialysis.
 - 1) Reassessment should be made after 12 weeks. If no response, treatment should be discontinued. If response is demonstrated, the lowest ESA dose sufficient to reduce the need for RBC transfusions should be used, and the Hgb should not exceed 11gm/dl. In those not on dialysis, the Hgb level should not exceed 10gm/dl.

Orthodontia Guideline Update

<u>Issue</u>: OHA Dental Director, Dr. Kaz Rafia, Dr. Rafia requested addition of a requirement to be clear of active dental decay and periodontal disease, as these conditions increase the medical risk of orthodontics. Dr. Rafia also requested the removal of #3-6 as they are redundant to #7 in the guideline.

HERC staff recommendation:

1) Modify Guideline Note 196 as shown below

GUIDELINE NOTE 169, ORTHODONTICS FOR CRANIOFACIAL ANOMALIES AND HANDICAPPING MALOCCLUSION

Line 256

Orthodontic treatment is included on this line for persons under the age of 21 with

- 1) Cleft lip and palate, cleft palate or cleft lip with alveolar process involvement, OR
- 2) Other craniofacial anomalies resulting in significant malocclusion expected to result in difficulty with mastication, speech, or other oral function, OR
- 3) Deep impinging overbite when lower incisors are destroying the soft tissue of the palate, tissue laceration and/or clinical attachment must be present, OR
- 4) Crossbite of individual anterior teeth when clinical attachment loss and recession of the gingival margin are present, OR
- 5)—Severe traumatic deviation, OR
- 6) Overjet greater than 9mm with incompetent lips or mandibular protrusion (reverse overjet) greater than 3.5mm with masticatory and speech difficulties; OR
- 7) Severe malocclusions with a Handicapping Labiolingual Deviation Index California Modification score of 26 or higher; AND
- 8) Free and clear of active decay and periodontal disease, verified by a dental exam in past 6 months

Advanced dental imaging is included on this line only when required for surgical planning for repair of craniofacial anomalies.

Below the Line Review 2022 Benign Carcinoid Gastrointestinal Tumors

<u>Issue</u>: Benign carcinoid tumors of the gut other than colon (ICD-10-CM D3A.0 family) are currently on line 638 BENIGN NEOPLASMS OF DIGESTIVE SYSTEM. Carcinoid tumors of the colon and appendix are on line 166 ANAL, RECTAL AND COLONIC POLYPS. Benign carcinoid tumors of other organs (lung, thymus) are on covered lines. Benign carcinoid tumors of the kidney (ICD-10CM D3A.093) are on line 511 ENIGN NEOPLASM OF KIDNEY AND OTHER URINARY ORGANS.

Malignant carcinoid tumors (ICD-10-CM C7A.0 family) are all on covered lines. Malignant GI carcinoid tumors are on line 157 CANCER OF COLON, RECTUM, SMALL INTESTINE AND ANUS.

Carcinoid tumor is a type of neuroendocrine tumor that grows from neuroendocrine cells and can occur in many parts of the body. Carcinoid tumors often grow very slowly. In children and young adults, carcinoid tumors are most often found in the appendix or in the lungs. In adults, carcinoid tumors are most often found in the digestive tract. Many patients have no symptoms from carcinoid tumors. Others have pain in the abdomen, nausea, diarrhea, or carcinoid syndrome (feeling flushed, nausea, diarrhea).

At the time of the creation of the Prioritized List, malignant and benign carcinoid tumors were thought to be distinct entities. Current understanding is that all carcinoid tumors are malignant and capable of metastasizing.

Per the NIH [https://www.cancer.gov/types/gi-carcinoid-tumors/patient/gi-carcinoid-treatment-pdq], carcinoid tumors are treated with surgery if resectable, radiation, chemotherapy, and/or hormone therapy.

HERC staff recommendations:

- Add the ICD-10-CM D3A.0 family (benign GI carcinoid tumors) to line 157 CANCER OF COLON, RECTUM, SMALL INTESTINE AND ANUS and remove from line 638 BENIGN NEOPLASMS OF DIGESTIVE SYSTEM
- 2) Add ICD-10CM D3A.093 (Benign carcinoid tumor of the kidney) to line 214 CANCER OF KIDNEY AND OTHER URINARY ORGANS and remove from line 638 BENIGN NEOPLASMS OF DIGESTIVE SYSTEM

Section 3.0 Coverage Guidances

SCOPE STATEMENT FOR HERC COVERAGE GUIDANCE

BARIATRIC PROCEDURE COVERAGE

Population	Adults and adolescents with obesity (BMI ≥30) who are being considered for
description	bariatric procedures
	Population scoping notes: Exclude overweight (BMI <30)
Intervention(s)	Bariatric procedures (for example adjustable gastric banding, Roux-en-y gastric bypass, biliopancreatic diversion, duodenal switch (BPD), vertical sleeve gastrectomy, single anastomosis duodenal-ileal bypass with sleeve gastrectomy (SADI-S), intragastric balloons) Intervention exclusions: bariatric devices that are not FDA approved or not available in the United States.
Comparator(s)	Nonsurgical treatment (for example, medical management, pharmacotherapy, intensive multicomponent behavioral interventions, behavioral counseling, structured weight management programs, other devices or procedures, or combinations of these therapies)
Outcome(s) (up to five)	Critical: All-cause mortality Important: Clinically significant improvement or resolution of chronic disease, weight change, quality of life, harms Considered but not selected for GRADE Table: Specific chronic diseases (for example, arthritis, sleep apnea), changes in healthcare utilization
Study Design(s)	Effectiveness: Randomized controlled trials (RCTs) only, comparative cohort studies for adolescent outcomes only Harms: RCTs; other observational studies (e.g., large registry studies)
Follow-up	Effectiveness: Minimum of 1 year Harms: Any time period

Key questions	1.	What is the effectiveness of bariatric procedures for the treatment of obesity in adults and adolescents as compared to other treatments?		
	2.	What are the harms of bariatric procedures for the treatment of obesity in idults and adolescents?		
	3.	Is there evidence of differential effectiveness or harms for bariatric procedures by:		
		a. Age		
		b. Sex		
		c. Race/ethnicity		
		d. BMI category		
		e. Comparator		
		f. Whether the patient has received prior bariatric surgery		
		g. Comorbidities (e.g., medical, behavioral health, other disabilities)		
		 Site of procedure (inpatient vs outpatient surgical center, centers of excellence vs not) 		
		i. Time since procedure		
Contextual	1.	What kinds of accreditation standards and center of excellence designations		
questions		exist in the United States and what are the requirements of each?		
	2.	What is the appropriate minimum age or developmental stage for bariatric surgery?		

CHANGE LOG

Date	Change	Rationale
3/16/2022	Added "comparator" to list of factors for key question 3.	Leadership discussion that some comparators are more effective
		than others.
4/7/2022	Deleted "laparascopic" qualifier for SADI-S, add	EbGS discussion
	comparative cohort studies for adolescent	
	outcomes. Added prior bariatric surgery to the list	
	of factors for key question 3.	

Coverage Guidance – PANDAS/PANS

<u>Question</u>: How should the Coverage Guidance *Pediatric Autoimmune Neuropsychiatric Disorders* Associated with Streptococcal Infections (PANDAS) and Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS) be applied to the Prioritized List?

Question source: EbGS

<u>Issue</u>: EbGS approved a coverage guidance regarding PANDAS/PANS at their April 2022 meeting. The "blue box" wording is shown below:

HERC Coverage Guidance

Tonsillectomy, adenoidectomy, adenotonsillectomy, and prophylactic antibiotic therapy are not recommended for coverage to treat pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) and pediatric acute-onset neuropsychiatric syndrome (PANS) (weak recommendation).

Up to 3 monthly immunomodulatory courses of intravenous immunoglobulin (IVIG) therapy or therapeutic plasma exchange are recommended for coverage to treat PANDAS and PANS (*weak recommendation*) when both of the following are met:

- a) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- b) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of IVIG or plasma exchange. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.

Coverage Guidance – PANDAS/PANS

Current Prioritized List status

CODES	DESCRIPTION	
ICD-10-0	CM Codes	242 DICORDERC INVOLVING THE
D89.89	Other specified disorders involving the immune	313 DISORDERS INVOLVING THE
	mechanism, not elsewhere classified	IMMUNE SYSTEM UNDEFINED CONDITIONS
D89.9	Disorder involving the immune mechanism,	UNDEFINED CONDITIONS
	unspecified	F36 VIDAL SELE LIMITING
		536 VIRAL, SELF-LIMITING ENCEPHALITIS, MYELITIS AND
G04.81	Other encephalitis and encephalomyelitis	ENCEPHALITIS, INVELITIS AND
		dysfunction lines
CPT Cod	los.	dysturiction lines
CFICOU		
90832	Behavioral therapy	Multiple lines but not 212
90652	Psychotherapy, 30 minutes with patient Psychotherapy, 30 minutes with patient when	Multiple lines but not 313 See above
	performed with an evaluation and management	See above
90833	service (List separately in addition to the code for	
	primary procedure)	
90834	Psychotherapy, 45 minutes with patient	See above
30034	Psychotherapy, 45 minutes with patient when	See above
	performed with an evaluation and management	See above
90836	service (List separately in addition to the code for	
	primary procedure)	
90837	Psychotherapy, 60 minutes with patient	See above
30037	Psychotherapy, 60 minutes with patient when	See above
	performed with an evaluation and management	See above
90838	service (List separately in addition to the code for	
	primary procedure)	
90839	Psychotherapy for crisis; first 60 minutes	See above
	Intravenous immunoglobulin therapy	
	Immune globulin (IVIG), human, for intravenous	Ancillary
90283	use	7
	Intravenous infusion, for therapy, prophylaxis, or	Ancillary
96365	diagnosis (specify substance or drug); initial, up	,
	to 1 hour	
	Intravenous infusion, for therapy, prophylaxis, or	Ancillary
	diagnosis (specify substance or drug); each	,
96366	additional hour (List separately in addition to	
	code for primary procedure)	
00004	Home infusion/specialty drug administration, per	Ancillary
99601	visit (up to 2 hours)	, ·
	Plasma exchange	
36514	Therapeutic apheresis; for plasma pheresis	Multiple lines including 313

Coverage Guidance – PANDAS/PANS

HERC staff recommendations:

- 1) Add ICD-10-CM D89.9 (Disorder involving the immune mechanism, unspecified) to line 313
- 2) Add a new guideline to line 313 DISORDERS INVOLVING THE IMMUNE SYSTEM as shown below
 - a. There are no CPT codes on line 313 for tonsillectomy, adenoidectomy, or adenotonsillectomy; therefore no need to call out lack of coverage in the guideline
 - b. CPT 36514 (Therapeutic apheresis) which is used for plasma pheresis is already on line 313.

GUIDELINE NOTE XXX PANDAS AND PANS

Line 313

Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) is included on this line when coded with ICD-10-CM D89.89 (Other specified disorders involving the immune mechanism, not elsewhere classified). Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS) is included on this line when coded with ICD-10-CM D89.9 (Disorder involving the immune mechanism, unspecified).

Up to 3 monthly immunomodulatory courses of intravenous immunoglobulin (IVIG) therapy or therapeutic plasma exchange are included on this line to treat PANDAS and PANS when both of the following are met:

- a) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- b) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of IVIG or plasma exchange. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.

Long term antibiotic therapy is not included on this line for treatment of PANDAS/PANS.

Health Evidence Review Commission (HERC)

Coverage Guidance: Pediatric Autoimmune Neuropsychiatric Disorders Associated with Streptococcal Infections (PANDAS) and Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS)

DRAFT for HERC meeting 5/19/2022

HERC Coverage Guidance

Tonsillectomy, adenoidectomy, adenotonsillectomy, and prophylactic antibiotic therapy are not recommended for coverage to treat pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) and pediatric acute-onset neuropsychiatric syndrome (PANS) (weak recommendation).

Up to 3 monthly immunomodulatory courses of intravenous immunoglobulin (IVIG) therapy or therapeutic plasma exchange are recommended for coverage to treat PANDAS and PANS (*weak recommendation*) when both of the following are met:

- A) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- B) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of IVIG or plasma exchange. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.

Note: Other treatments (corticosteroids, SSRIs, NSAIDs, short-course antibiotics, and behavioral therapies) were included in an initial version of this report. These therapies were determined to be beyond the scope of a HERC coverage guidance, as these therapies are commonly used for many indications and are not typically subject to utilization control. Only treatments subject to coverage criteria were retained for the final version of this report.

Note. Definitions for strength of recommendation are in Appendix A, GRADE Table Element Descriptions. Rationales for each recommendation appear below in the GRADE tables.



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Rationale for development of coverage guidances and multisector intervention reports

Coverage guidances are developed to inform coverage recommendations for public and private health plans in Oregon as plan administrators seek to improve patients' experience of care, population health, and the cost-effectiveness of health care. In the era of public and private sector health system transformation, reaching these goals requires a focus on maximizing the benefits and minimizing the harms and costs of health interventions.

HERC uses the following principles in selecting topics for its reports to guide public and private payers:

- Represents a significant burden of disease or health problem
- Represents important uncertainty with regard to effectiveness or harms
- Represents important variation or controversy in implementation or practice
- Represents high costs or significant economic impact
- Topic is of high public interest

HERC bases its reports on a review of the best available research applicable to the intervention(s) in question. For coverage guidances, which focus on diagnostic and clinical interventions, evidence is evaluated using an adaptation of the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) methodology. For more information on coverage guidance methodology, see Appendix A.

Multisector interventions can be effective ways to prevent, treat, or manage disease at a population level. In some cases, HERC has reviewed evidence and identified effective interventions, but has not made formal coverage recommendations when these policies are implemented in settings other than traditional health care delivery systems because effectiveness could depend on the environment in which the intervention is implemented.

GRADE Tables

HERC develops recommendations by using the concepts of the GRADE system. GRADE is a transparent and structured process for developing and presenting evidence and for performing the steps involved in developing recommendations. The table below lists the elements that determine the strength of a recommendation. HERC reviews the evidence and assesses each element, which in turn is used to develop the recommendations presented in the coverage guidance box. Estimates of effect are derived from the evidence presented in this document. Assessments of confidence are from the published systematic reviews and meta-analyses, where available and judged to be reliable. The level of confidence in the estimate is determined by HERC based on the assessment of 2 independent reviewers from the Center for Evidence-based Policy.

In some cases, no systematic reviews or meta-analyses encompass the most current literature. In those cases, HERC may describe the additional evidence or alter the assessments of confidence in light of all available information. Such assessments are informed by clinical epidemiologists from the Center for Evidence-based Policy. Unless otherwise noted, statements regarding resource allocation, values and preferences, and other considerations are the assessments of HERC, as informed by the evidence reviewed, public testimony, and subcommittee discussion.

Recommendations for coverage are based on the balance of benefit and harms, resource allocation, values and preferences and other considerations. See Appendix A for more details about the factors that constitute the GRADE table.



GRADE Tables

Should prophylactic antibiotics be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome	Resource	Values and	Other
Outcomes	Confidence in Estimate	Allocation	Preferences	Considerations
Change in psychiatric symptom scores (Critical outcome) Hospitalizations (Critical outcome)	In a single RCT (N = 37) comparing penicillin to placebo for 4 months, there was no significant difference in neuropsychiatric symptoms between children when they received penicillin or placebo. The cross-over design meant that each participant was in both the intervention and control group for the analysis. In a second RCT comparing prophylactic azithromycin (N = 17) to placebo (N = 14) for 4 weeks, the azithromycin group had a significantly greater reduction OCD severity, but no significant difference in number of obsessions and compulsions. Significantly more children were classified as responding at a clinically significant level in the azithromycin group (7/17) than in the placebo group (1/14). In a third RCT that tested prophylactic antibiotic therapy for 1 year, significantly more (5/11) children who received penicillin demonstrated significant reduction in symptoms compared to (1/12) children who received azithromycin. Compared to the year before baseline, children in both groups had fewer exacerbations during the trial year. • • (very low confidence, based on 3 RCTs, n = 91) No evidence identified.	Antibiotics are inexpensive and readily available. Treatment of complications of long-term or frequent antibiotic use would add cost.	Some parents would want any treatment that might help their child's symptoms. However, other parents would have concerns about the risks and side effects of long-term or frequent antibiotic use.	Long-term or frequent antibiotic use is associated with a range of negative consequences, including but not limited to <i>C. difficile</i> infection, gut flora disruption, diarrhea, and increased antibiotic resistance leading to reduced ability to treat other infections with antibiotics. Most health plan cover short-term antibiotics without prior authorization criteria but may scrutinize or not cover long-term prescriptions.

Should prophylactic antibiotics be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome Confidence in Estimate	Resource Allocation	Values and Preferences	Other Considerations
Harms (Important outcome)	The few harms that were reported included heart rate irregularity (9/12) for children who received azithromycin, and loose stool (no statistics reported). • • (very low confidence, based on 1 RCTs, n = 23)			
Function or quality of life for patient (Important outcome)	No evidence identified.			
Function or quality of life for patient (Important outcome)	No evidence identified.			

Balance of benefits and harms: We have very low confidence that prophylactic antibiotic use is helpful in PANDAS/PANS, given the small sample sizes in the included studies. There are concerning known harms of frequent or long-term antibiotic use.

Rationale: Prophylactic antibiotic therapy is not recommended for coverage for PANDAS/PANS because of insufficient comparative evidence that prophylactic antibiotic use leads to any measurable benefit for these conditions. The known risks of prophylactic or long-term antibiotic use outweigh potential benefits in these conditions. The recommendation is weak because of the very low quality of the evidence.

Recommendation: Prophylactic antibiotic therapy is not recommended for coverage for PANDAS/PANS (weak recommendation).

Note. GRADE table elements are described in Appendix A. A GRADE Evidence Profile is in Appendix B.

Abbreviations. OCD: obsessive-compulsive disorder; PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome; RCT: randomized controlled trial.

Should tonsillectomy and/or adenoidectomy be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome/ Confidence in Estimate	Resource Allocation	Values and Preferences	Other Considerations
Change in psychiatric symptom scores (Critical outcome)	No difference in neuropsychiatric symptoms between surgery and non-surgery groups among children diagnosed with PANDAS. ••○ (low confidence, based on 2 comparative cohort studies, n = 232)	Tonsillectomy and adenoidectomy are invasive procedures requiring general anesthesia and specialty surgical care.	Parents would not value an invasive surgery with risks as well as the risks of general anesthesia	ive and/or adenoidectomy frequently have coverage limitations, such as multiple streptococcal infections in one
Hospitalizations (Critical outcome) Harms (Important outcome)	No evidence identified. No evidence identified.		for a procedure that has no evidence of benefits.	
Function or quality of life for patient (Important outcome)	No evidence identified.			year. This procedure has historically been overused.
Function or quality of life for patient (Important outcome)	No evidence identified.			

Balance of benefits and harms: We have low confidence that that there is no benefit from tonsillectomy and/or adenoidectomy for PANDAS/PANS, and this procedure has known harms.

Rationale: Tonsillectomy and/or adenoidectomy are not recommended for coverage for treatment of PANDAS/PANS because these procedures have known harms and because evidence shows that these procedures do not improve the outcomes in this condition. The recommendation is weak because of the low quality of the evidence.

Recommendation: Tonsillectomy and/or adenoidectomy are not recommended for coverage (*weak recommendation*) for treatment of PANDAS/PANS.

Note. GRADE table elements are described in Appendix A. A GRADE Evidence Profile is in Appendix B.

Abbreviations. PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome.

Should IVIG be recommended for coverage for PANDAS/PANS?

Outcome	Estimate of Effect for Outcome/ Confidence in Estimate	Deserves Allegation	Values and	Other
Outcomes		Resource Allocation	Preferences	Considerations
Change in	Compared to Saline Placebo	IVIG is expensive and	Parents would	IVIG is a blood
psychiatric	Among children meeting the criteria for PANDAS and	requires the cost of an	value any	product with the
symptom scores	OCD in an RCT, 7/18 had a significant decrease in	infusion center,	treatment that	inherent risks that
(Critical outcome)	symptoms 6 weeks after receiving 2 consecutive days	nursing care, and	would improve	accompany
	of IVIG infusions, and 4/17 children in the placebo	possible	their child's	accepting any form
	group had a significant decrease in symptoms. When	hospitalization.	symptoms.	of blood product.
	comparing the IVIG group and placebo group, there	Treatment for side	However, many	
	were no statistically significant differences. During an	effects of IVIG would	parents would	IVIG therapy has a
	open-label phase of this same trial, 17/24 children	add cost.	value avoiding a	significant rate of
	meeting the criteria for PANDAS and OCD had a		treatment with	mild side effects
	significant decrease in symptoms 12 to 18 weeks	IVIG is a scarce	known side effects	including fever,
	after receiving 2 consecutive days of IVIG infusions	resource and	that has little	body aches, nausea,
	on 1 or 2 occasions.	shortages have been	evidence of	rash, and fatigue.
	Another RCT compared children who received IVIG (N	reported in the past.	effectiveness.	
	= 9) to children who received saline placebo (N = 10)			Severe side effects
	1 month after treatment reported that the IVIG			include thrombosis,
	group improved significantly more on most measures			renal dysfunction,
	compared to the placebo group. One year after			and acute renal
	treatment, the improvements in the IVIG group were			failure, and life-
	maintained, but the placebo group was not followed			threatening allergic
	to determine whether the IVIG group's symptoms			reaction.
	remained significantly better than the placebo			
	group's symptoms.			IVIG can interfere
				with vaccine
	Compared to plasma exchange			effectiveness for
	No significant difference 1 month or 1 year after			vaccines given
	treatment between children receiving IVIG (N = 9) or			
	plasma exchange (N = 10); both groups had			

Should IVIG be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome/ Confidence in Estimate	Resource Allocation	Values and Preferences	Other Considerations
	significant improvement in symptoms compared to baseline at both 1-month and 1-year follow-ups			within several months of IVIG.
Hospitalizations (Critical outcome)	• (very low confidence, based on 2 RCTs, N = 54) No evidence identified.			Several products on the market are FDA- approved for people
Harms (Important outcome)	1/33 children who received IVIG infusions had an allergic reaction to the IVIG infusion that resolved without complication. 31/33 children reported mild or moderate adverse events such as nausea, vomiting, headache, fever, joint pain, tiredness, stomach pain, or decreased appetite. • • (very low confidence, based on 2 RCTs, N = 64)			under the age of 19.
Function or quality of life for patient (Important outcome)	No evidence identified.			
Function or quality of life for patient (Important outcome)	No evidence identified.			

Balance of benefits and harms: There were mixed results from 2 very small trials regarding the clinical effectiveness of IVIG. Outside of PANDAS, no evidence met inclusion criteria for PANS. IVIG has a significant rate of known harms.

Rationale: Expert opinion and lower-quality observational data indicate there may be benefit for some patients with these conditions, but recommended treatment protocols and criteria for treatment vary widely. Due to the severe impact of symptoms associated with PANDAS/PANS on child health, growth and development, and the lack of known effective treatments, coverage of IVIG is recommended when recommended by the patient's PCP and a pediatric subspecialist and after less-intensive therapies were not effective, were not tolerated or did not result in sustained improvement in symptoms. The recommendation is weak because of the very low quality of the evidence.

Should IVIG be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome/	Resource Allocation	Values and	Other
	Confidence in Estimate		Preferences	Considerations

Recommendation:

Up to 3 monthly immunomodulatory courses of intravenous immunoglobulin (IVIG) therapy are recommended for coverage to treat PANDAS and PANS (*weak recommendation*) when both of the following are met:

- a) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- b) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of IVIG. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.

Note. GRADE table elements are described in Appendix A. A GRADE Evidence Profile is in Appendix B.

Abbreviations. IVIG: intravenous immunoglobulin; OCD: obsessive-compulsive disorder; PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome; RCT: randomized controlled trial.

Should therapeutic plasma exchange be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome/	Resource Allocation	Values and	Other
Outcomes	Confidence in Estimate		Preferences	Considerations
Change in psychiatric symptom scores (Critical outcome)	Compared to saline placebo In the same RCT that is described in the IVIG table, the plasma exchange group (N = 10) was compared to the same placebo group (N = 10) 1 month after treatment. The plasma exchange group improved significantly more on most measures compared to the placebo group. One year after treatment, the improvements in the plasma exchange were maintained, but the placebo group was not followed to determine whether the plasma exchange group's symptoms remained significantly better than the placebo group's symptoms.	Plasma exchange is an expensive therapy which requires a monitored infusion in a clinical setting. Children in the studies included in this review required multiple treatment sessions.		
	Compared to intravenous immunoglobulin No significant difference 1 month or 1 year after treatment between children receiving IVIG (N = 9) or plasma exchange (N = 10); both groups had significant improvement in symptoms compared to baseline at both 1-month and 1-year follow-ups • • (very low confidence, based on 1 RCT, N = 29)			blood pressure, nausea, vomiting, itchy skin, hives, low calcium levels in the blood, venous access malfunction, infections, thrombosis, and anaphylactic shock.
Hospitalizations (Critical outcome)	No evidence found.			
Harms (Important outcome)	All children who received plasma exchange (10/10) experienced mild side effects such as nausea, vomiting, anxiety, or fever. • • (very low confidence, based on 1 RCT, N = 29)			

Should therapeutic plasma exchange be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome/ Confidence in Estimate	Resource Allocation	Values and Preferences	Other Considerations
Function or	No evidence found.			
quality of life for				
patient (Important				
outcome)				
Function or	No evidence found.			
quality of life for				
patient (Important				
outcome)				

Balance of benefits and harms: The comparative evidence that plasma exchange is effective for treating PANDAS is very limited and shows no clear benefit. The harms of this treatment are generally mild but serious complications can occur.

Rationale: Expert opinion and lower-quality observational data indicate there may be benefit for some patients with these conditions, but recommended treatment protocols and criteria for treatment vary widely. Due to the severe impact of symptoms associated with PANDAS/PANS on child health, growth and development, and the lack of known effective treatments, coverage of plasma exchange is recommended when recommended by the patient's PCP and a pediatric subspecialist and after less-intensive therapies were not effective, were not tolerated or did not result in sustained improvement in symptoms.

Should therapeutic plasma exchange be recommended for coverage for PANDAS/PANS?

Outcomes	Estimate of Effect for Outcome/	Resource Allocation	Values and	Other
	Confidence in Estimate		Preferences	Considerations

Recommendation:

Up to 3 monthly immunomodulatory courses of therapeutic plasma exchange are recommended for coverage to treat PANDAS and PANS (*weak recommendation*) when both of the following are met:

- a) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- b) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of plasma exchange. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.

Note. GRADE table elements are described in Appendix A. A GRADE Evidence Profile is in Appendix B.

Abbreviations. IVIG: intravenous immunoglobulin; PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome; RCT: randomized controlled trial.

Background

Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) and pediatric acute-onset neuropsychiatric syndrome (PANS are conditions associated with a sudden onset of changes or regression in behaviors and experiences prior to puberty in multiple domains, such as motor, neurological, psychiatric, and biological systems.¹⁻³ Care providers and researchers from multiple disciplines (including microbiology, neurology, neuroimmunology, immunology, child psychiatry, infectious diseases, rheumatology, and pediatrics) have contributed to publications seeking to define these conditions.³ These conditions have an abrupt onset of symptoms and may include exacerbations, sudden worsening of symptoms in short bursts, in a sawtooth-like pattern.¹⁻³

In PANDAS, the triggering mechanism for these changes is hypothesized to be a beta-hemolytic streptococcal infection within 6 months of symptom onset, and is characterized by sudden onset of obsessive-compulsive disorder (OCD), along with verbal or motor tics.^{2,4} This hypothesized disease pathway aligns with large epidemiological cohort studies of children in Europe⁵ and Asia⁶ that observed an 18% to 22% increased risk of any neuropsychiatric disorders in children who had streptococcal infections as compared with children without streptococcal infections, including a higher risk for obsessive-compulsive and tic disorders.^{5,6} However, some researchers suggest that tying the diagnosis to streptococcus infection to the exclusion of other etiologies has limited the exploration of other disease pathways that could inform diagnosis and treatment of symptoms.^{7,8}The prevalence of PANDAS is not known, but some studies suggest that males are more likely than females to be diagnosed with PANDAS.⁹

PANS is characterized by sudden onset of OCD, with or without severe eating restrictions, and 2 or more other symptoms in neurological, behavioral, or cognitive domains.³ PANDAS can be considered a subset of PANS. These symptoms could result from multiple disease pathways or other disorders, including but not limited to streptococcus, varicella, or bacterial pneumonia infections.^{3,10} The prevalence of PANS is not known.

Two other conditions with similar symptoms are pediatric infection-triggered autoimmune neuropsychiatric disorders (PITAND) and childhood acute neuropsychiatric syndromes (CANS).^{10,11}

The natural histories of PANDAS and PANS are still being studied, but early signals suggest that 60% to 80% of pediatric patients have a significant reduction in symptoms over time, similar to childhood-onset OCD. The American Academy of Child and Adolescent Psychiatry published a practice parameter for assessing and treating childhood-onset OCD; they noted some clinical experts believe a small subset of children that have been diagnosed with OCD or Tourette disorder might have clinical exacerbations linked to streptococcal infection. The authors report that more males than females are diagnosed with pediatric OCD, typically diagnosed between the ages of 7 and 12 years; earlier onset is associated with comorbid psychiatric diagnoses (e.g., mood disorders, attention deficit disorder, anxiety disorders, phobias). Since the property of t

There is some discussion about whether PANDAS and PANS is related to pediatric autoimmune encephalitis, which is also characterized by abrupt onset of similar abnormal behavioral symptoms and disruptions in multiple biological systems (e.g., gastrointestinal, nervous). 1,14-16 Autoimmune encephalitis in children is characterized by a sudden onset of symptoms including seizures, irritability, aggression, and abnormal movements, and could be associated with an acute infection or presence of a tumor. 1,14,17

The prevalence of pediatric autoimmune encephalitis is not known, but a population study of adults and children suggested the incidence rate of autoimmune encephalitis was 0.8 per 100,000, and that males had more than twice the prevalence of females. Autoimmune encephalitis is a life-threatening condition usually treated in a hospital setting. Because of the differences in diagnostic criteria and disease process between autoimmune encephalitis and PANDAS/PANS, the scope of this report excludes autoimmune encephalitis.

Diagnostic Criteria and Tests

Table 1 presents diagnostic criteria and tests by condition and includes information from publications summarized in the Evidence Review and Clinical Practice Guidelines sections of this coverage guidance. 3,11,14-17,19-32

Table 1. Proposed Diagnostic Criteria, Tests and Processes

Proposed Diagnostic Tests and Processes
In patients with OCD, complete blood count, erythrocyte sedimentation rate, C-reactive protein, metabolic panel, urine analysis, pharyngeal swab and anti-streptococcal antibodies. Positive results from the pharyngeal swab and anti-streptococcal antibodies indicate exposure to the streptococcal infection do not differentiate between the state of carrier and acute infection. For children with neurological and psychiatric symptoms, physical or neurological examination require the analysis of the cerebrospinal fluid and neuroimaging exams.
Differential diagnosis.
Complete medical and psychiatric history, physical examination, laboratory testing of blood and possibly cerebrospinal fluid, and selected paraclinical evaluations, such as magnetic resonance imaging, electrocardiogram/ echocardiography, electroencephalography, and polysomnography.

Proposed Diagnostic Criteria	Proposed Diagnostic Tests and Processes
 Somatic signs and symptoms, including sleep disturbances, enuresis, or urinary frequency 	
Rule out Sydenham chorea, autoimmune encephalitis, neuropsychiatric lupus, central	Differential diagnosis.
nervous system vasculitis, and other conditions that better account for the symptoms	

Abbreviations. OCD: obsessive-compulsive disorder; PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome.

We identified 4 publications that specifically presented or summarized evidence for diagnostic criteria and tests related to PANDAS or PANS. 4,9,33,34 Nielsen and colleagues performed a systematic review and meta-analysis of studies on the association between streptococcal infections and exacerbations of neuropsychiatric symptoms. The authors concluded that although children diagnosed with PANDAS had more neuropsychiatric exacerbations than children with streptococcal infections without a follow-up diagnosis of PANDAS, these exacerbation were not significantly temporally associated with streptococcal infections. 34

Baj and colleagues reviewed published literature in search of distinguishing features of patients diagnosed with PANDAS and concluded that despite more than 20 years of research into this condition, it remains challenging to differentiate PANDAS from OCD or tic disorders. Their observations of characteristics that appear to be different for children diagnosed with PANDAS include:

- some alterations of cortico-basal ganglia circuitry, due to the effect of antibodies produced in response to the condition on various neuronal proteins, including tubulin, lysoganglioside, and dopamine receptors;
- deposits of antibodies which are also accumulated in the striatal interneurons;
- significantly enlarged volumes of corpus striatum, caudate, putamen, globus pallidus, and basal ganglia; and
- significant alterations to the gut microbiota.

Gamucci and colleagues described the clinical, neuropsychological, and biological characterization of PANDAS and PANS, and recommended 4 categories of tools to add in the diagnostic process.⁴ Proposed neuropsychological tests to assess motor and vocal tics, obsession and compulsion⁴:

- Children's Yale—Brown Obsessive Compulsive Scale for presence and severity of motor and vocal tics: and
- Yale Global Tic Severity Scale for presence and severity of child's obsession and compulsion.

Proposed neuropsychological tests to assess anxiety⁴:

• Multidimensional Anxiety Scale for Children (MASC) for the presence and types of child's anxiety symptoms for ages 8 to 19 years.

Proposed neuropsychological tests to assess short-term memory and attention⁴:

 Digit Span subtest Wechsler Intelligence Scale for Children for verbal short-term memory for ages 6 to 16 years;

- Coding subtest Wechsler Intelligence Scale for Children for visual-motor dexterity and nonverbal short-term memory for ages 6 to 16 years; and
- Symbol Search subtest Wechsler Intelligence Scale for Children for accuracy, attention and concentration for ages 6 to 16 years.

Proposed neuropsychological tests to assess processing speed⁴:

 Processing Speed Index Wechsler Intelligence Scale for Children (WISC III-IV) for speed of cognitive processes and response output on visual-motor tasks for ages 6 to 16 years

In addition to the scales proposed by Gamucci and colleagues above, Leibold and colleagues validated a Global Impairment Score scale to measure impairment in children and adolescents as part of the diagnostic process for PANS.³³ This scale was designed to be answered by a child's caregiver, and is scored on a scale of 0 to 100.³³

For additional measures proposed in guidelines, please refer to the Clinical Practice Guidelines section of this coverage guidance.

Treatments

Table 2 presents treatments by condition and includes information about treatments from the publications summarized in the evidence review and clinical practice guidelines sections of this coverage guidance. ^{3,11,14-17,19-32} Not all treatments in Table 2 have been evaluated in studies with prospective comparative designs; the evidence review portion of this coverage guidance will synthesize findings from comparative studies related to treatments and outcomes.

Table 2. Treatments Proposed for PANDAS and PANS

Treatments	PANDAS	PANS
Antibiotics		
Amoxicillin	X	Х
Aripiprazole		Χ
Azithromycin	X	
Penicillin	X	
Surgical Interventions		
Tonsillectomy	X	
Adenoidectomy	X	
Intravenous Immunoglobulin and Plasma Exchange		
Intravenous immunoglobulin	X	Χ
Plasma exchange	X	X
SSRIs		
Fluoxetine	X	
NSAIDs		
Naproxen sodium	X	
Antipsychotics		
Pimozide	X	
Risperidone		Χ
Corticosteroids		
Dexamethasone		Χ

Treatments	PANDAS	PANS
Prednisone	X	
Behavioral Interventions		
Cognitive behavioral therapy	X	

Abbreviations. NSAID: nonsteroidal anti-inflammatory drug; PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome; SSRI: selective serotonin reuptake inhibitor.

Evidence Review

We identified 2 systematic reviews, 5 RCTs with 6 publications, and 2 comparative cohort studies that reported interventions for children diagnosed with PANDAS or PANS. 11,24-32 Table 3 summarizes key characteristics of each included study. Given the varied study designs, treatments, and outcomes collected, neither of the systematic reviews included a meta-analysis section.

Sigra and colleagues included in their systematic review any report of any treatments for children with PANDAS, PANS, CANS, or PITAND published in English that also reported outcomes; this expansive inclusion criteria resulted in 5 RCTs, 7 observational survey study, and 65 case reports.²⁴ We rated this systematic review itself as having a low risk of bias, although it is important to note that the review authors concluded that there is not enough rigorous research about treatments for children with PANDAS, PANS, CANS, or PITAND, and the existing studies themselves have a high risk of bias. Sigra and colleagues concluded there was insufficient evidence to clearly recommend specific treatments for children with these diagnoses, but that psychiatric behavioral interventions, immunomodulatory therapies, and antibiotics likely have roles in the treatment of these disorders and should be more systematically investigated.²⁴

In addition to summarizing comparative evidence regarding antibiotics, tonsillectomy, IVIG, and therapeutic plasma exchange, Sigra and colleagues sumarized noncomparative evidence for behavioral therapy, corticosteroids, nonsteroidal anti-inflammatory drugs (NSAIDs), and selective serotonin reuptake inhibitors (SSRIs). The <u>first draft of this coverage guidance</u> included the latter interventions, although it was not possible to make a clear determination of effectiveness or harms of these interventions due to the lack of comparative evidence.

Farhood and colleagues included in their systematic review 13 studies testing treatments for PANDAS that also reported outcomes related to change in symptoms, and excluded case reports; 3 included studies were RCTs, and 10 had retrospective designs.²⁷ We rated this review as having a high risk of bias. This review included studies of adenotonsillectomy, antibiotic therapy, intravenous immunoglobulin (IVIG) therapy, and cognitive behavioral therapy.²⁷ The authors suggested that immunoglobulin therapy might be effective for certain populations, and that psychotherapy and antibiotic therapies were likely low-risk interventions.²⁷ However, the authors concluded that the study designs left results open to question due to inability to account for confounding factors, such as co-occurring treatments, and were unable to strongly recommend any specific course of treatment.²⁷ All of the studies included in Farhood and colleagues' systematic review were also included in Sigra and colleagues' systematic review. Given the later search and publication dates and the lower risk of bias for Sigra and colleagues' review, we restrict our summary of review findings to the Sigra review in the following sections.

The RCTs all had fewer than 40 participating children, so the number of children in each treatment and placebo group was also small during comparative stages of the trials. These RCTs compared antibiotics to placebo and had moderate to high risk of bias, ^{25,30,31} or compared IVIG to placebo or plasma exchange and had low to high risk of bias. ^{26,32} At the end of the trial phase, the investigators of 3 of the RCTs offered the active treatment under consideration to the children who had been in the group receiving a placebo, which makes the long-term follow-up of participants in these trials an open-label observation follow-up (range, 4 weeks to 57 months). ^{11,25,26,31}

The number of children included in the 2 comparative cohort studies was larger (more than 100), and both studies focused on surgical interventions for symptom relief for children diagnosed with PANDAS.^{28,31} We rated both studies as having a high risk of bias, primarily due to an inability to account for confounding factors.

The following sections organize findings from these studies by type of intervention. First, we summarize relevant RCTs and comparative cohort studies, and then we compare those findings with conclusions from the systematic reviews that included results from noncomparative study designs such as case reports.



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Table 3. Characteristics of Included Studies

First Author, Year Number Follow-up(s)	Inclusion Criteria	Exclusion Criteria	Intervention	Risk of Bias
Systematic Reviews				
Sigra et al., 2018 ²⁴ 5 RCTs, 7 observational survey studies, and 65 case reports Not applicable	Studies in which patients with PANDAS, PANS, CANS, or PITAND were given treatment, that presented outcome data, and were written in English	No exclusion criteria explicitly listed	Cognitive behavior therapy, antibiotics, tonsillectomy, corticosteroids, therapeutic plasma exchange, IVIG, rituximab, nonsteroidal anti-inflammatory drugs	Low
Farhood et al., 2016 ²⁷ 3 RCTs and 10 retrospective designs Not applicable RCTs	Studies testing treatments for PANDAS and reported outcomes, and were written in English or Spanish	Review articles, single case reports, and studies of natural history or diagnostic strategies	Tonsillectomy, adenoidectomy, antibiotics, IVIG, cognitive behavioral therapy, or SSRIs	High
Murphy et al., 2017 ²⁵ N = 31 2 and 4 weeks	Children with an acute onset or acute relapse within 6 months of evaluation (abrupt, dramatic overnight onset) of moderate or worse OCD symptoms and presence of a sudden and severe co-occurrence of at least 2 neuropsychiatric symptoms.	Children with a gradual onset or duration of OCD symptoms of more than 6 months; who were receiving extended-course antibiotics (i.e., not a typical treatment course of antibiotics for an infection, or prophylactic antibiotics) and/or other immune therapy for PANS; with a primary diagnosis of tics; who were receiving exposure-based cognitive behavioral therapy; who had a history of nonresponse to a prior antibiotic trial; or who had a diagnosis of moderate to severe autism spectrum disorder,	Azithromycin and probiotics versus placebo with probiotics for 4 weeks; after this all participants were offered azithromycin	Moderate

First Author, Year Number	Inclusion Criteria	Exclusion Criteria	Intervention	Risk of Bias
Follow-up(s)				
		intellectual disability, and/or chronic neurological disease.		
Williams et al., 2016 ²⁶ Leon et al., 2018 ¹¹ N = 35 3 and 6 months during the trial, and a 57-month observational follow- up	Children who were 4 to 13 years of age in first episode of PANDAS symptoms and documentation that symptoms first appeared within 6 to 8 weeks of streptococcal infection or exposure; who had a sudden onset or exacerbation of OCD (reaching peak severity and impairment within 24 to 48 hours); and had at least 3 neuropsychiatric symptoms (which meets criteria for PANS).	Children with a history of Sydenham chorea or acute rheumatic fever; who had symptoms consistent with autism spectrum disorder or schizophrenia; who had severe physical, behavioral, or psychiatric symptoms that would prevent study participation; or prior corticosteroid or immunomodulatory therapy for PANDAS	IVIG versus placebo for 6 weeks; participants in the placebo group were then given the opportunity to receive IVIG; 31 participants received at least 1 dose of IVIG over the course of the study	Low risk for original trial, and high risk for long- term follow- up
Snider et al., 2005 ³⁰ N = 23 12 months	Children with a tic disorder and/or OCD; who had a history of a sudden onset of symptoms or an episodic course with abrupt symptom exacerbations interspersed with periods of partial or complete remission; who had onset of neuropsychiatric symptoms prior to puberty; and who had documentation of a temporal association between a preceding streptococcal infection and the onset or exacerbation of neuropsychiatric symptoms.	No specific exclusion criteria listed.	Azithromycin versus penicillin for 12 months	High

First Author, Year Number Follow-up(s)	Inclusion Criteria	Exclusion Criteria	Intervention	Risk of Bias
Garvey et al., 1999 ³¹ N = 37 4 months	Children between 4 and 15 years of age with a tic disorder and/or OCD; who had history of a sudden onset of symptoms or an episodic course with abrupt symptom exacerbations interspersed with periods of partial or complete remission (a sawtooth, rather than a waxing and waning course); who had an onset of symptoms prior to puberty; and evidence of a temporal association between a preceding streptococcal infection and the onset or exacerbation of neuropsychiatric symptoms.	Children who had tics or OCD of such a severity that hospitalization was considered; who required treatment for severe, active comorbid major psychiatric disorders; who had with autism, pervasive developmental delay, or "mental retardation" or who had neurologic diagnoses other than tics and Tourette syndrome, serious concurrent or chronic medical disorders, and a personal history of penicillin allergy.	Penicillin versus placebo for 4 months; cross-over design meant that all participants received penicillin during the 8 months of the study	High
Perlmutter et al., 1999 ³² N = 29 1 month and 12 months	Children ages 5 to 14 years with a tic disorder and/or OCD; onset of neuropsychiatric signs and symptoms before puberty; a history of sudden onset of signs and symptoms, or an episodic course characterized by abrupt exacerbations and periods of partial or complete remission; evidence of, and association between, streptococcal infection and onset or exacerbation of signs and symptoms; and current exacerbation severe	Children with a history of Sydenham's chorea or rheumatic fever, autism, schizophrenia or other psychotic disorder, a neurological disorder other than a tic disorder, an autoimmune disorder, or other medical illness. Immunoglobulin concentrations were measured, and children were excluded from the study if they had IgA deficiency (a contraindication to IVIG administration).	Plasma exchange, IVIG, or placebo for 2 weeks	High

First Author, Year				
Number	Inclusion Criteria	Exclusion Criteria	Intervention	Risk of Bias
Follow-up(s)				
	enough to cause significant distress and interfere with the child's social functioning in at least 2 spheres (home, school, social relations).			
Comparative Cohort S	tudies			
Pavone et al., 2014 ²⁸ N = 120 Every 2 months for 2 years	Children with a tic disorder and/or OCD; who had infection-related symptom flare-ups, history of dramatic onset of either OCD or tics, new onset anxiety, sensory or motor abnormalities, behavioral regression, deterioration in school performance, emotional lability, or urinary symptoms (all these neuropsychiatric phenomena were in temporal association to streptococcal pharyngeal tonsillitis). The surgical group (n = 56) were referred to surgery based on a clinical history of recurrent inflammation in addition to the symptoms above.	No specific exclusion criteria listed	Surgery versus no surgery; surgery group had 25 tonsillectomies and 31 adenotonsillectomies	High
Murphy et al., 2013 ²⁹ N = 112 Not reported	Children with a tic disorder and/or OCD; and with infection-related symptom flare-ups, history of dramatic onset of either OCD or tics, new onset anxiety, sensory or motor	Children with a psychotic disorder, significant medical illness, or nontic neurologic disorder	Surgery versus no surgery; surgery group had 4 tonsillectomies, 10 adenoidectomies, and 22 had both procedures	High

First Author, Year Number Follow-up(s)	Inclusion Criteria	Exclusion Criteria	Intervention	Risk of Bias
	abnormalities, behavioral regression, deterioration in school performance, emotional lability, or urinary symptoms. Participants on stable doses of psychotropic medication for their condition were not excluded. The surgical group comprised children who had a tonsillectomy and/or adenoidectomy procedure, and were matched to nonsurgery participants on age and sex.			

Note. This language was taken directly from the study; the coverage guidance authors recognize this language is no longer acceptable.

Abbreviations. CANS: childhood acute neuropsychiatric syndromes; IgA: immunoglobulin A; IVIG: intravenous immunoglobulin; OCD: obsessive-compulsive disorder; PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections; PANS: pediatric acute-onset neuropsychiatric syndrome; PITAND: pediatric infection-triggered autoimmune neuropsychiatric disorders; RCT: randomized controlled trial; SSRI: selective serotonin reuptake inhibitor.

Antibiotics

We identified 3 RCTs that tested antibiotics as a primary intervention for children diagnosed with PANDAS or PANS. ^{25,30,31} As a reminder, children meeting the criteria for PANDAS also meet the criteria for PANS. Conclusions from both systematic reviews agreed with author conclusions of these 3 RCTs: there is some evidence that antibiotic prophylaxis might reduce exacerbations of neuropsychiatric symptoms for children diagnosed with PANDAS. ^{24,27}

Azithromycin

Murphy and colleagues conducted a double-blind RCT with 31 participants randomized to receive azithromycin prophylaxis (N = 17) for 4 weeks or to receive a placebo (N = 14) for 4 weeks; participants in the placebo group were then given the option to begin taking azithromycin, which launched the open-label observational portion of the study. Both groups also received twice daily probiotics. We rated the outcomes from the trial portion of this study as having a moderate risk of bias; no outcomes were reported for the open-label portion.

When comparing scores on the OCD Clinical Global Impressions Severity scale (which has a scale of 1 to 7), participants who received azithromycin reported statistically significantly greater reductions in symptom frequency 4 weeks after baseline (azithromycin group mean, 4.06; azithromycin group standard deviation [SD], 0.23; placebo group mean, 4.93; placebo group SD, 0.25; effect size, 0.11; P = .003). The effect size for the difference in symptoms between the azithromycin and placebo groups suggests that there was only a very small difference between the 2 groups, and that the difference was not likely to be clinically significant. No significant difference was found between the group on the Children's Yale-Brown Obsessive-Compulsive Scale, and no difference between groups for the severity of symptoms. The severity of symptoms.

Investigators also assessed whether participants responded to their assigned therapy, using a 30% or greater reduction in symptoms to judge whether a participant responded. In the azithromycin group, 52.9% (9 of 17) were categorized as responders, and 21.4% (3 of 14) were categorized as responders in the placebo group.²⁵

The authors reported that among participants with greater tic severity scores at baseline (measured as 1 standard deviation greater than average number of tics), participants in the azithromycin group were significantly more likely to have at least a 30% reduction in tic symptoms during the 4-week trial than control group participants (no statistics reported; P < .05). If there is a treatment benefit to azithromycin, this suggests that it might have greater benefit for children with more severe tics.

Penicillin

Garvey and colleagues conducted a double-blind, balanced crossover study with 37 participants randomized to receive either penicillin prophylaxis or a placebo for 4 months.³¹ After the first 4 months passed, the treatment assignment was reversed for 4 months; therefore, participants were followed for 8 months.³¹ There was no wash out period between the reversal of treatment assignment.³¹ We rated this study as having a high risk of bias. No statistically significant difference was reported between treatment groups for exacerbations of neuropsychiatric symptoms, with 38 exacerbations during the placebo phase and 35 exacerbations during the penicillin phase.³¹ There were no clinically meaningful differences in depression or anxiety symptoms between the treatment phases.³¹ Of the 27 parents who

provided global ratings of their child's behaviors, 22 reported an improvement of behavior during the penicillin phase; 18 of these parents correctly identified this as the active treatment phase when rating their child's behavior.³¹ There were no statistically significant differences in neuropsychiatric symptoms between the penicillin and placebo phases, as measured by the Children's Yale-Brown Obsessive Compulsive Scale (P = .16) or the Yale Global Tic Severity Scale (P = .28).³¹

Azithromycin Versus Penicillin

Snider and colleagues conducted a double-blind RCT with 23 participants randomized to receive either azithromycin or penicillin prophylaxis for 12 months. We rated this study as having a high risk of bias. The authors reported that both antibiotic therapies reduced the number of streptococcal infections during the study year compared to the year prior to the study (mean reduction of about 2 infections per year), with no significant difference between the 2 groups (mean for both groups, 0.1; SD for both groups, 0.3; P > .05). Parent and child reports at baseline and the end of the study were reviewed and rated by the study authors to determine the presence and frequency of exacerbations of neuropsychiatric symptoms. Both groups reported decreased neuropsychiatric exacerbations, but the participants who received penicillin reported significantly fewer exacerbations of neuropsychiatric symptoms (penicillin group mean, 0.5; penicillin group SD, 0.5; azithromycin group mean, 0.9; azithromycin group SD, 0.5; P < .01).

Tonsillectomies and Adenoidectomies

We identified 2 comparative cohort studies that examined the association of tonsillectomies and adenoidectomies with change in symptoms for children diagnosed with PANDAS, and both compared children with PANDAS who had either or both of these surgeries (N = 88) to children with PANDAS who had received neither surgery (N = 140).^{28,29} Both studies specifically named PANDAS as the diagnosis of focus.^{28,29} We rated both of these studies as having a high risk of bias. Both systematic reviews agreed with the conclusions of the authors from these 2 studies that tonsillectomy and adenoidectomy do not appear to reduce neuropsychiatric symptom severity or exacerbations.^{24,27} We did not identify any studies that tested the surgical interventions of tonsillectomies and adenoidectomies for the broader diagnosis of PANS.

In a prospective comparative cohort study including 120 participants, Pavone and colleagues reported that there was no significant difference in symptom remission rates between the surgery and nonsurgery groups (relative risk [RR], 1.39; 95% confidence interval [CI], 0.75 to 2.55; P = 0.29). The authors also reported no significant difference in days to first symptom relapse (surgery group mean, 45.1; surgery group SD, 17.8; nonsurgery group mean, 39.3; nonsurgery group SD, 14.2; P = .09).

Murphy and colleagues conducted a prospective comparative cohort study including 112 children who met the criteria for an OCD or tic diagnosis, and were divided into a group meeting the criteria for PANDAS and a group that did not meet criteria for PANDAS, according to a temporal relationship with a streptococcal infection. The authors reported no significant difference in OCD or tic severity between the surgery and nonsurgery groups among children with or without a PANDAS diagnosis, as measured by the Children's Yale-Brown Obsessive Compulsive Scale (surgery group mean, 17.9; surgery group SD, 9.9; nonsurgery group mean, 18.7; nonsurgery group SD, 10.5; P = .71) or the Yale Global Tic Severity Scale (surgery group mean, 33.4; surgery group SD, 23.5; nonsurgery group mean, 33.6; nonsurgery group SD, 21.6; P = .97). The authors also reported that there was no relationship between surgery status and

age of onset of OCD or tic symptoms (surgery group mean, 5.9 years; surgery group SD, 2.1 years; nonsurgery group mean, 6.5 years; nonsurgery group SD, 2.7 years; P = .32). There was no statistically significant relationship between surgery status and duration of symptoms (surgery group mean, 2.5 years; surgery group SD, 2.1 years; nonsurgery group mean, 3.3 years; nonsurgery group SD, 2.5 years; P = .09).

Both comparative cohort studies concluded that the surgical interventions had no effect on severity of symptoms or symptom progression. ^{28,29}

IVIG

We identified a single RCT that tested IVIG versus placebo, ^{11,26} and a single RCT that tested IVIG versus a placebo or plasma exchange. ³² Both RCTs enrolled children who met the diagnostic criteria for PANDAS and OCD. ^{26,32}

IVIG Versus Saline Placebo

Williams and colleagues randomized 35 children to receive IVIG or an intravenous saline placebo for 2 consecutive days at trial start.²⁶ All children were prescribed prophylactic antibiotics for the duration of the 6 months of this study, and penicillin was reported as the most commonly prescribed antibiotic (no number reported).¹¹ The investigators then offered the opportunity to children who had received the placebo to enter an open-label phase in which they received IVIG along with the children in the intervention group who were judged to be nonresponders to the treatment 6 weeks after the first infusion.²⁶ The investigators defined responding to treatment before the trial began as a 30% or greater decrease in symptoms as measured by the Children's Yale Brown Obsessive Compulsive Scale, or by a "much" or "very much" improved rating on the Clinical Global Impressions Improvement scale.²⁶ We rated the first phase of this trial as having a low risk of bias, and the 6- to 12-week open-label phase and the 24-week follow-up with any associated outcomes as having a high risk of bias.

At the conclusion of the 6-week blinded trial phase, there were no significant differences between the intervention and control groups for neuropsychiatric symptoms, as measured by changed in scores between baseline and 6-week follow-up on the Clinical Global Impressions Improvement scale and the Children's Yale Brown Obsessive Compulsive Scale.²⁶

- Seven of the participants in the intervention group (38.9%; intervention group N = 18) were classified as responders to the treatment, meaning that they either demonstrated a 30% or greater decrease in symptoms as measured by the Children's Yale Brown Obsessive Compulsive Scale, or by a "much" or "very much" improved rating on the Clinical Global Impressions Improvement scale. ²⁶ In the placebo group, 4 children were classified as having a significant decrease in symptoms (23.5%; placebo group N = 17). ²⁶
- There was not a significant difference in the number of children in each group who had a significant improvement in symptoms (P = .40). The authors also reported that was no significant difference in the average change in symptoms between the intervention group and placebo group, as measured by the Clinical Global Impressions Improvement scale (P = .12) or the Children's Yale-Brown Obsessive Compulsive Scale (P = .44).

During the nonblinded, open-label phase, 24 participants received IVIG.²⁶ This included 10 of 18 participants who were originally randomized to the intervention group and who were classified as

nonresponders at the end of the 6-week blinded phase; these participants therefore received doses of IVIG on 2 consecutive days twice: at baseline and 6 weeks after baseline.²⁶ Of the participants in the open-label phase, 17 (70.8%) were classified as responding to the treatment by 24 weeks.²⁶ However, there was no comparator group for this phase of the study and the authors did not report follow up at 24 weeks for the group of initial responders in the blinded phase of the RCT.

Leon and colleagues conducted additional follow-up interviews by telephone for all 35 original study participants for up to 5 years. ¹¹ The authors reported that after the trial, 6 participants had tonsillectomy, 11 participants were diagnosed with new psychiatric conditions (i.e., attention-deficit/hyperactivity disorder, depression, anxiety, phobia, or chronic tic disorder), and 24 (68.6%) had experienced an exacerbation of symptoms. ¹¹ Those exacerbations were treated with a variety of approaches, including additional IVIG, antibiotics, psychiatric medications, and cognitive behavioral therapy; treatments were often combined and used at the same time. ¹¹

IVIG Versus Plasma Exchange or Saline Placebo

Perlmutter and colleagues randomized 29 children who met the diagnostic criteria for PANDAS or OCD to receive IVIG, plasma exchange, or a saline placebo. 32 The authors compared symptoms at baseline to the same symptoms measured 1 month after treatment. 32 Participants in the plasma exchange group (N = 10) received 5 or 6 exchange transfusions, which required 85 to 121 minutes per transfusion. 32 Participants in the IVIG group (N = 9) received infusions during 2 days at the start of the trial; participants in the control group received a saline placebo (N = 10). 32 On average, participants in both the plasma exchange group and IVIG group reported significant reduction in symptoms from baseline to 1 month and between baseline and the 1-year follow-up, as measured by obsessive-compulsive symptoms, psychosocial functioning (i.e., anxiety, depression, and emotional lability), and global functioning. 32

The authors reported comparisons of the change in symptoms for the 2 intervention groups to the change in symptoms for the saline placebo group between baseline and 1-month follow-up.³² In comparison with the changes in scores in the saline placebo group (N = 10) 1 month after treatment, the IVIG group's (N = 9)³²:

- scores for obsessions and compulsions decreased (45% vs. 3%; P < .05);
- scores for tics did not decrease significantly (19% vs. 12%; P >.05);
- sum of obsessions, compulsions, and tics decreased (45% vs. 6%; P < .05);
- scores for global impairment improved (26% vs. 1%; P < .05);
- scores for psychosocial functioning did not significantly improve (20% vs. 0%; P > .05); and
- scores for global severity improved significantly (26% vs. 1%; *P* < .05).

One year after treatment, all 9 participants who received IVIG were successfully followed and readministered the measures described above; 7 of 9 were judged to be "much" or "very much" improved in a global assessment of symptoms by their parents.³² There were no comparisons made between the control group and the intravenous exchange group 1 year after baseline.³²

The authors noted that there was not a statistically significant difference between the IVIG and plasma exchange groups for improvement in any symptoms at 1 month or 1 year after treatment.³² They reported that scores for both active intervention groups remained significantly improved from baseline for obsessions and compulsions, psychosocial functioning, and global severity.³² However, the

participants who received IVIG did not show a statistically significant improvement in tics at 1 year after baseline when compared to their own scores. ³²

Plasma Exchange

We identified a single RCT that tested plasma exchange versus placebo or IVIG for children who met the criteria for PANDAS and OCD; this study conducted by Perlmutter and colleagues is also described in the section that describes studies of IVIG.³² We rated this study as having a high risk of bias. In comparison with the placebo group (N = 10) 1 month after treatment, the plasma exchange group's (N = 10)³²:

- scores for obsessions and compulsions decreased (58% vs. 3%; P < .05);
- scores for tics decreased (49% vs. 12%; P <.05);
- sum of obsessions, compulsions, and tics decreased (54% vs. 6%; P < .05);
- scores for global impairment improved (36% vs. 1%; P < .05);
- scores for psychosocial functioning did not significantly improve (30% vs. 3%; P > .05); and
- scores for global severity improved (26% vs. 1%; P < .05).

One year after baseline, 8 of 10 participants who received plasma exchange were successfully followed and readministered the measures described above; 7 of 8 were judged to be "much" or "very much" improved in a global assessment of symptoms by their parents.³² There were no comparisons made between the control group and the intravenous exchange group 1 year after treatment.³²

The authors noted that there was not a statistically significant difference between the IVIG and plasma exchange groups for improvement in any symptoms at 1 month or 1 year after treatment.³² They reported that scores for both active intervention groups remained significantly improved from baseline for obsessions and compulsions, psychosocial functioning, and global severity.³² In addition to those measures, the participants who received plasama exchange also remained significantly improved on the measure of tics when compared to their scores at baseline.³²

Harms

Sigra and colleagues' systematic review of any treatment for PANDAS, PANS, CANS or PITAND reported that adverse events reported in included studies were typically mild to moderate in nature, including nausea, vomiting, headache and stomachache.²⁴

Antibiotics

Murphy and colleagues reported that some participants who received prophylactic azithromycin had loose stools (no number reported), and 9 out of 12 children who received azithromycin had heart rate irregularities.²⁵

Other known adverse events associated with long-term antibiotic therapy include skin rashes, gastrointestinal disturbance, increased risk of childhood-onset chronic conditions (e.g., asthma, obesity, attention deficit hyperactivity disorder), and contribute to antibiotic resistance.^{35,36} Use of azithromycin may also result in changes in the electrical activity of the heart that can lead to fatal irregular heart rhythm.³⁷

Tonsillectomy and Adenoidectomy

Although the included studies did not report harms, adverse events associated with tonsillectomy and adenoidectomy may include hemorrhage, complications from anesthesia, and infection. 38,39

IVIG

Williams and colleagues reported that a single participant appeared to have an allergic reaction to the IVIG infusion, but that the reaction resolved without complication. The authors also reported that several participants noted minor discomforts during treatment, such as joint pain, headache, stomach pain, tiredness, and anxiety. Perlmutter and colleagues reported that 6 of 9 children receiving immunoglobulin infusions reported experiencing 1 or more of the following: nausea, vomiting, mild to moderately severe headache, and low grade fever. All of these symptoms were resolved with hydration therapy, paracetamol, or diphenhydramine. No long-term adverse events were reported, and none of the studies mentioned intending to collect information about long-term adverse events. L1,26,32

The FDA categorized IVIG as a biologic agent, and 8 of the 12 products listed are approved for use in children under 18 years of age (ASCENIV, Flebogamma, Gammagard Liquid, Gammagard S/D, Gammaplex, Gamunex-C, PANZYGA, and Privigen).⁴⁰ None of the approved indications include PANDAS or PANS for these products, and the age range for approved use vary by product.⁴⁰ The package inserts for IVIG products include black box warnings for thrombosis, renal dysfunction, and acute renal failure.⁴⁰

Plasma Exchange

Perlmutter and colleagues reported that 7 of 10 children who received plasma exchange reported pallor, dizziness, and nausea during the first exchange transfusions; 2 of these children also experienced vomiting.³² Three additional children reported feeling anxious during the exchange transfusions.³²

Known complications of plasma exchange include circuit clotting, low or high blood pressure, nausea, vomiting, itchy skin, hives, low calcium levels in the blood, venous access malfunction, infections, thrombosis, anaphylactic shock, and high fever.⁴¹⁻⁴⁴

Ongoing Studies

We identified 3 ongoing studies that might provide upcoming information about diagnosis and treatment of PANDAS or PANS. 45-47

A single double-blinded RCT plans to enroll 44 children diagnosed with PANDAS to test the effectiveness of taking naproxen sodium twice daily for 8 weeks on the severity of OCD symptoms, as measured by the second edition of the Children's Yale-Brown Obsessive-Compulsive Scale.⁴⁶ Enrolled participants will be between 6 and 15 years of age with first OCD symptom onset within 18 months prior to trial start date, and have symptoms that significantly interfere with daily life.⁴⁶ The estimated primary completion date is October 2022.⁴⁶

A single RCT plans to enroll 92 children from 6 to 17 years of age with a confirmed diagnosis of PANS or PANDAS, and will randomize participants to receive intravenous immunoglobulin therapy or a placebo; the participant, care provider, investigator, and outcomes assessor will all be blinded.⁴⁷ The estimated study start date is August 30, 2021, and the estimated primary completion date is March 2023.⁴⁷ The

primary outcome measure will be the Children's Yale-Brown Obsessive Compulsive Scale at 9 weeks after treatment, which will be measured as a secondary outcome at week 18 along with Clinical Global Impression assessment, the Parent Obsessive-Compulsive Impact Scale, the Child Obsessive-Compulsive Impact Scale, the Swanson, Nolan, And Pelham Scale - Version IV (SNAP-IV; measures symptoms and behaviors that could impact child's behaviors at school), and the Parent Tic Questionnaire.⁴⁷

This study will exclude children whose symptoms had first onset more than 6 months before the trial start date, children with current relapse of symptoms whose first onset was more than 12 months before the trial start date; who have a contraindication for intravenous immunoglobulin; who have severely restricted food intake, whose body mass index is 40 or greater; who have symptoms of autism or schizophrenia, bipolar disorder, or other psychotic disorder; who have serious or unstable mental illness; who have been treated with corticosteroids or began cognitive behavioral therapy within the 8 weeks prior to randomization; who have a history of rheumatic fever; who have had prior immunomodulatory treatment; who had taken antibiotics or antivirals for an acute infection within 1 week of randomization; who have severe liver disease; who have known hepatitis B, hepatitis C, or HIV infection; pregnant or lactating women or women unwilling to comply with contraception protocol; or who participated in another interventional trial within 3 months of randomization or during the course of this study.⁴⁷

A single observational matched cohort study plans to enroll 500 children diagnosed with PANS who have not yet received any treatment, whose symptoms began within 1 month of enrollment date, and who are 18 years of age or younger. The investigators plan to match these children with healthy children without a PANS diagnosis to examine immunologic, neurologic, genomic, and behavioral differences between the two cohorts. This study began in 2013 and has an estimated primary completion date of March 2028. Outcome measures include the following, measured ever every 2 to 4 weeks for up to 12 years: Global Impairment Score, Children's Yale-Brown Obsessive Compulsive Scale, Columbia Impairment Score, Caregiver Burden Inventory, and neurological findings (e.g., irregular movements).

Evidence Summary

The origins and progression of symptoms associated with PANDAS and PANS are still being studied and documented; there are few published studies that tested whether antibiotic therapy, surgical interventions, IVIG, or plasma exchange might improve symptoms in children diagnosed with these conditions. It is also difficult to know how long any improvements in symptoms last after children receive the treatments we reviewed in this coverage guidance, because they often receive multiple treatments (simultaneously or 1 after another). Additionally, it is hard to distinguish whether patterns of exacerbation and resolution of symptoms can be directly attributed to infections and treatments, or if there is an underlying pattern of increase of symptoms followed by a decrease of symptoms that would occur without these treatments. It is not clear how long any treatment benefit might be sustained before another exacerbation, or whether any treatment alone or in combination with other treatments can prevent or shorten the length of exacerbations.

 We have very low confidence that prophylactic antibiotic therapy reduces exacerbations of neuropsychiatric symptoms. Risks for long-term antibiotic use include skin rashes, gastrointestinal disturbance, increased risk of childhood-onset chronic conditions (e.g., asthma, obesity, attention deficit hyperactivity disorder), and contribute to antibiotic resistance.^{35,36}

- We did not identify any evidence testing antibiotics in response to current psychiatric exacerbation.
- We have low confidence that surgical interventions such as tonsillectomy and adenoidectomy do not reduce neuropsychiatric symptom exacerbations. Harms of tonsillectomy and adenoidectomy may include hemorrhage, complications from anesthesia, and infection.^{38,39}
- We have very low confidence that IVIG decreases neuropsychiatric symptoms. It is not clear how long any treatment benefit might be sustained. There is an ongoing trial of IVIG for children with PANS or PANDAS that might have published results in 2023 or 2024. The package inserts for IVIG products include serious warnings for thrombosis, renal dysfunction, and acute renal failure.⁴⁸
- We have very low confidence that plasma exchange decreases neuropsychiatric symptoms. It is not clear how long any treatment benefit might be sustained. Known complications of plasma exchange transfusions include high fever, blood clots, infection, minor or severe allergic reactions, and high or low blood pressure.⁴¹⁻⁴⁴

The very low and low confidence we have in the findings above means that findings from new comparative studies that test treatments for PANDAS or PANS could change the recommendations that we make for which treatments should be covered for children diagnosed with PANDAS or PANS.

Clinical Practice Guidelines

We identified 6 publications that included recent guidelines for the diagnosis and treatment of individuals with PANDAS or PANS.^{3,19-23} We rated all the guidelines as having poor methodological quality.

PANS/PANDAS Clinical Research Consortium

The most recent clinical guidelines written and published in the US for treating PANS was written by members of the PANS/PANDAS Research Consortium at workgroup meetings partially sponsored by the National Institutes of Health.³ The workgroups reviewed literature, reviewed more than 1,000 cases of children diagnosed with PANDAS/PANS, and then prepared summaries to be reviewed by review panels of clinical experts who either worked with children suspected of having PANDAS/PANS or were experts in child psychiatry, pediatrics, infectious diseases, microbiology, neurology, neuroimmunology, immunology, and rheumatology.³ Not all experts agreed on all treatments proposed in the guidelines, so the guideline committee opted to describe multiple treatment options beyond the treatments that had the highest consensus.³ The authors of the committee summary stated that they expect the guidelines to be altered over time in response to the initiation and completion of new controlled clinical trials testing the efficacy of treatments.³

As an overview, the guidelines recommend a 3-pronged approach to treating PANS^{3,19,20,22}:

- "treating the symptoms with psychoactive medications, psychotherapies (particularly cognitive behavioral therapy), and supportive interventions;
- removing the source of the inflammation with antimicrobial interventions; and
- treating disturbances of the immune system with immunomodulatory and/or anti-inflammatory therapies" (pp. 562; Swedo et al., 2017).

The guidelines presented the following 6 principles for the identification and treatment of PANS:

- 1. Establish that PANS is the correct "diagnosis of exclusion" by completing a comprehensive diagnostic evaluation.²³
- 2. Provide symptomatic relief with psychiatric medications and behavioral interventions, prioritizing treatment of symptoms causing the greatest distress and interference.²²
- 3. Treat underlying infections and consider use of therapeutic or prophylactic antibiotics. 20
- 4. Treat symptoms resulting from neuroinflammation or postinfectious autoimmunity with antiinflammatory or immunomodulatory therapies, chosen on the basis of symptom severity and disease trajectory.¹⁹
- 5. Evaluate effectiveness of the treatment regimen at frequent intervals, making modifications as warranted by improvement or worsening of symptoms.³
- 6. Treatment can be tapered downward or stopped when symptoms resolve. However, treatment may be necessary again at some point in the future, given the relapsing–remitting nature of PANS symptoms.³

Clinical Guidance About PANS from Nordic Countries

The Nordic Pediatric Immunopsychiatry group published guidance for diagnosis and management of suspected PANS in 2021, and included pediatric neurologists, child psychologists, and child psychiatrists from Denmark, Norway, Sweden and Great Britain.²¹ The authors intended this guidance to propose a standard set of diagnostic criteria for PANDAS and PANS, and to propose a standard process for diagnostic evaluation.²¹

The authors agreed to adopt the clinical criteria proposed by Chang and colleagues for PANS that was published in 2015^{21,23}:

- 1. Abrupt, dramatic onset (culmination within 72 hours) of OCD or severely restricted food intake.
- 2. Concurrent presence of additional neuropsychiatric symptoms, with similarly severe and acute onset, from at least two of the following seven categories (see reference for full description):
 - Anxiety,
 - Emotional liability and/or depression,
 - o Irritability, aggression and/or severely oppositional behaviors,
 - Behavioral (developmental) regression,
 - Deterioration in school performance,
 - Sensory or motor abnormalities and
 - Somatic signs and symptoms, including sleep disturbances, enuresis or increased urinary frequency.
- 3. Symptoms are not better explained by a known medical disorder, such as Sydenham's chorea, systemic lupus erythematosus, Tourette disorder or others.

The authors agreed to adopt Swedo and colleagues' diagnostic criteria for PANDAS that were published in 1998^{21,49}:

- Presence of Obsessive Compulsive Disorder and/or a tic disorder: The patient must meet lifetime diagnostic criteria (DSM- III- R or DSM- IV) for Obsessive Compulsive Disorder or a tic disorder.
- 2. Pediatric onset: Symptoms of the disorder first become evident between 3 years of age and the beginning of puberty.

- 3. Episodic course of symptom severity: Clinical course is characterized by the abrupt onset of symptoms or by dramatic symptom exacerbations. Symptoms usually decrease significantly between episodes and occasionally resolve completely between exacerbations.
- 4. Association with group A Beta- hemolytic streptococcus infection: Symptom exacerbations must be temporally related to group A Beta- hemolytic streptococcus infection, that is associated with positive throat culture and/or significantly elevated anti- group A Beta- hemolytic streptococcus antibody titers.
- 5. Association with neurological abnormalities: During symptom exacerbations, patients will have abnormal results on neurological examination. Motoric hyperactivity and adventitious movements (including choreiform movements or tics) are particularly common.

In addition to the criteria listed above, the authors further proposed a definition of severe symptoms and required that the child meet at least 1 major criteria and 1 minor criteria. The major criteria included: total Children's Yale- Brown Obsessive Compulsive Scale (CY- BOCS) score ≥24; reduced intake of food or fluid, leading to less urine production (less than three urinations daily) or weight loss (more than 10%); and severe tics (Yale Global Tic Severity Scale (YGTSS) total tic severity score ≥40 but <50). Minor criteria included being absent from school at least 50% of class days during 1 month, and inability to participate in leisure activities or loss of social contact. In the criteria included being absent from school at least 50% of class days during 1 month, and inability to participate in leisure activities or loss of social contact.

The authors then proposed a standard clinical work-up, which is described in Table 4.

Table 4. Nordic Pediatric Immunopsychiatry Group's Proposed Clinical Work-Up for PANS

Examination	Instrument or Analysis	Description		
Psychiatric	Psychiatric			
General	Achenbach System of Empirically Based Assessment (ASEBA),19 Mini international neuropsychiatric interview (M.I.N.I KID) or equivalent	General assessment of psychiatric conditions		
	Child and Adolescent Trauma Screen (CATS)	Trauma screening		
	Children's Global Assessment Scale (C- GAS)	Assessment of general functioning		
	Clinical Global Impression- Severity Scale (CGI- S)	Clinician- rated severity of the patient's illness at time of assessment		
	Pediatric Quality of Life Inventory (PedsQL)	Assessment of quality of life		
	Optional: Work and Social Adjustment Scale (WSAS) 2	Measure of impaired functioning		
	Optional: KIDSCREEN	Assessment of subjective health and well- being		
Symptom-specific	Children's Yale- Brown Obsessive Compulsive Scale (CY- BOCS)	OCD inventory		
	The Screen for Child Anxiety Related Disorders (SCARED)	Screening for child anxiety related disorders		
	Yale Global Tic Severity Scale (YGTSS)	Tics inventory		

Examination	Instrument or Analysis	Description
	Kiddie Schedule for Affective Disorders and Schizophrenia (Kiddie- SADS)	Interview screening for psychiatric diagnoses
	ADHD rating scale (ADHD- RS)	Questionnaire related to inattention, hyperactivity and impulsivity
	Behavior Rating Inventory of Executive Function (BRIEF)	Behavior Rating Inventory of Executive Function
Infectious		
General	Throat: bacterial culture	No description
	Blood: complete blood cell count with differential count, antistreptolysin- O and antideoxyribonuclease B antibodies	No description
Symptom-specific	Throat: Mycoplasma Polymerase Chain Reaction (PCR)	No description
	Nasopharynx: Aspirate PCR panel	Common viral airway infections such as influenza virus and enterovirus
	Urine analysis and culture	No description
Extended workup	Cerebrospinal fluid cell count, protein, glucose, lactate; Epstein- Barr-virus/cytomegalovirus/varicella zoster virus/herpes simplex virus/Mycoplasma/enterovirus/influenza virus immunoglobulin G and immunoglobulin M +Polymerase Chain Reaction (PCR); Borrelia burgdorferi immunoglobulin G and immunoglobulin M (paired with serum)	No description
Immunological		,
General	Blood: erythrocyte sedimentation rate (ESR), antiphospholipid antibodies (anticardiolipin and beta2 glycoprotein 1 antibodies), antinuclear antibodies (antidsDNA, ANA IIF, anti- ENA screen: Anti- SSA, anti- SSA, anti- SSB, anti- Sm, anti- Scl-70, anti- Jo1, anti- Centromer B (- CENP- B) and anti- U1- RNP), immunoglobulins subclasses, tissue- transglutaminase IgA and deamined gliadinpeptide IgG (Celiac disease), neuronal antibodies, Myelin oligodendrocyte glycoprotein (MOG) antibodies, antithyroperoxidase (TPO), thyroid stimulating hormone (TSH) receptor antibodies, TSH, T3 and free T4, complement C3 and C4, angiotensin- converting enzyme (ACE), Vitamin- D, Vitamin B12, ferritin, cupper, ceruloplasmin, cytokines	No description

Examination	Instrument or Analysis	Description		
Extended work-up	Cerebrospinal fluid Lumbar opening pressure, neuronal antibodies (standard panel), immunoglobulin G, index and electrophoresis for oligoclonal bands (paired with serum), and cytokines	No description		
Toxicological				
Symptom-specific	Drug screening	No description		
Metabolic				
Symptom-specific	Urine metabolic screening	No description		
Radiological				
Extended work-up	Cerebral MRI including contrast: structural, diffusion and FLAIR sequences	No description		
Neuropsychological	Neuropsychological			
Extended work-up	Standard or sleep electroencephalogram	No description		

Note. This table is reproduced from Tables 3 and 4 on pages 4 and 5 of the Nordic Pediatric Immunopsychiatry group's published guidance for diagnosis and management of suspected PANS. Abbreviations. FLAIR: fluid attenuated inversion recovery; MRI: magnetic resonance imaging; OCD: obsessive-compulsive disorder; PANS: pediatric acute-onset neuropsychiatric syndrome.

The authors recommended that verified or strongly suspected bacterial infections should be treated at the discretion of the provider for a maximum of 14 days; however, they do not recommend prophylactic antibiotic therapy. They further recommended that any other treatment occur within ongoing clinical research or under the guidance of centers that specialize in the care of children with suspected PANS. Such treatments for children with severe symptoms might begin with oral non-steroidal anti-inflammatory drugs, proceed to steroids if ineffective, and finally proceed to intravenous immunoglobulin. The authors state that plasma exchange, and cytostatic and immunomodulatory drugs are only clinically indicated when a child has been diagnosed with autoimmune encephalitis.

Policy Landscape

Payer Coverage Policies

We did not identify coverage policies for Washington State's Medicaid program or national or local coverage determinations for Medicare related to PANDAS or PANS.

We identified coverage policies related to PANDAS and PANS from 2 private payers (Aetna and Cigna), but we did not identify coverage policies related to PANDAS or PANS for BlueCross BlueShield or for Moda.

Private Payers

Aetna considers parenteral immunoglobulins, rituximab, and plasmapheresis to be investigational or experimental for PANDAS and autoimmune encephalitis.⁵⁰⁻⁵²

Cigna considers plasmapheresis, immune globulin, and rituximab to be investigational or experimental for PANDAS and PANS in policies last updated in 2021.⁵³⁻⁵⁵ These coverage policies consider plasmapheresis to be medically necessary as a primary therapy for autoimmune encephalitis characterized by the presence of the n-methyl D-aspartate receptor antibody.⁵⁵

Recommendations from Others

We did not identify policy statements or recommendations for PANDAS or PANS from the American Neurology Association, the American Academy of Pediatrics, the American Association of Immunologists, the Infectious Diseases Society of America, or the American Psychiatric Association.

PANDAS Physician Network

The PANDAS Physician Network maintains a <u>website</u> with tools such as flowcharts for diagnosing and treating PANS and PANDAS, and for classifying symptoms into mild, moderate, or severe cases.⁵⁶ The authors recommend that children with moderate or severe symptoms be treated by an experienced team of multidisciplinary providers or a PANS/PANDAS specialist.⁵⁶ To summarize the proposed elements of the treatment guidelines (please note that this list is simplified)⁵⁶:

- 1. Start with 14 days of antibiotic therapy, and consider the appropriateness of prophylactic antibiotic therapy; lengthen therapy if infection is not resolved or symptoms persist.
- 2. Consider 5 to 7 days of non-steroidal anti-inflammatory drugs.
- 3. Ensure family access to cognitive behavioral therapy, and parenting management techniques.
- 4. Consider steroid course if no improvement from first 3 steps.
- 5. Escalate to intravenous immunoglobulin therapy if first 4 steps have not resolved symptoms.
- 6. If symptoms do not resolve, consider a second course of intravenous immunoglobulin or evaluate the need for plasma exchange, and prescribe prophylactic antibiotic therapy.

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Coverage guidance is prepared by the Health Evidence Review Commission (HERC), HERC staff, and subcommittee members. The evidence summary is prepared by the Center for Evidence-based Policy at Oregon Health & Science University (the Center). This document is intended to guide public and private purchasers in Oregon in making informed decisions about health care services.

The Center is not engaged in rendering any clinical, legal, business or other professional advice. The statements in this document do not represent official policy positions of the Center. Researchers involved in preparing this document have no affiliations or financial involvement that conflict with material presented in this document.

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Appendix A. GRADE Table Element Descriptions

Element	Description
Balance of benefits and harms	The larger the difference between the desirable and undesirable effects, the higher the likelihood that a strong recommendation is warranted. An estimate that is not statistically significant or has a confidence interval crossing a predetermined clinical decision threshold will be downgraded.
Quality of evidence	The higher the quality of evidence, the higher the likelihood that a strong recommendation is warranted
Resource allocation	The higher the costs of an intervention—that is, the greater the resources consumed in the absence of likely cost offsets—the lower the likelihood that a strong recommendation is warranted
Values and preferences	The more values and preferences vary, or the greater the uncertainty in values and preferences, the higher the likelihood that a weak recommendation is warranted
Other considerations	Other considerations include issues about the implementation and operationalization of the technology or intervention in health systems and practices within Oregon.

Strong recommendation

In Favor: The subcommittee concludes that the desirable effects of adherence to a recommendation outweigh the undesirable effects, considering the balance of benefits and harms, resource allocation, values and preferences and other factors.

Against: The subcommittee concludes that the undesirable effects of adherence to a recommendation outweigh the desirable effects, considering the balance of benefits and harms, resource allocation, values and preferences and other factors.

Weak recommendation

In Favor: The subcommittee concludes that the desirable effects of adherence to a recommendation probably outweigh the undesirable effects, considering the balance of benefits and harms, resource allocation, values and preferences and other factors., but further research or additional information could lead to a different conclusion.

Against: The subcommittee concludes that the undesirable effects of adherence to a recommendation probably outweigh the desirable effects, considering the balance of benefits and harms, cost and resource allocation, and values and preferences, but further research or additional information could lead to a different conclusion.

Confidence in estimate rating across studies for the intervention/outcome

Assessment of confidence in estimate includes factors such as risk of bias, precision, directness, consistency and publication bias.

High: The subcommittee is very confident that the true effect lies close to that of the estimate of the effect. Typical sets of studies are RCTs with few or no limitations and the estimate of effect is likely stable.

Moderate: The subcommittee is moderately confident in the estimate of effect: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different. Typical sets of studies are RCTs with some limitations or well-performed nonrandomized studies with additional strengths that guard against potential bias and have large estimates of effects.

Low: The subcommittee's confidence in the estimate of effect is limited: The true effect may be substantially different from the estimate of the effect. Typical sets of studies are RCTs with serious limitations or nonrandomized studies without special strengths.

Very low: The subcommittee has very little confidence in the estimate of effect: The true effect is likely to be substantially different from the estimate of effect. Typical sets of studies are nonrandomized studies with serious limitations or inconsistent results across studies.



Appendix B. GRADE Evidence Profile

Quality Assessment (Confidence in Estimate of Effect) for Antibiotics										
No. of Studies	Study Design(s)	Risk of Bias	Inconsistency	Indirectness	Imprecision	Other Factors	Level of Confidence			
Change in Psychiatric Symptoms										
3	RCTs	Moderate to High	Not serious	Not serious	Serious	Small sample sizes, short follow-	Very Low ●○○			
Hospital	izations					ир				
Harms										
1	RCT	High	Unable to rate	Not serious	Serious	Small sample sizes, short follow- up	Very Low ●○○			
Function	Function or Quality of Life for Patient									
0 Function	Function or Quality of Life for Parent									
0										

Abbreviation. RCT: randomized controlled trial.

Quality Assessment (Confidence in Estimate of Effect) for Tonsillectomy or Adenoidectomy										
No. of Studies	Study Design(s)	Risk of Bias	Inconsistency	Indirectness	Imprecision	Other Factors	Level of Confidence			
Change i	Change in Psychiatric Symptoms									
2	Comparative	High	Not serious	Serious	Not serious	None	Low			
	cohort						••0			
Hospitali	Hospitalizations									
0										
Harms	Harms									
0										
Function or Quality of Life for Patient										
0										
Function or Quality of Life for Parent										
0										

	Quality Assessment (Confidence in Estimate of Effect) for IVIG									
No. of Studies	Study Design(s)	Risk of Bias	Inconsistency	Indirectness	Imprecision	Other Factors	Level of Confidence			
Change i	Change in Psychiatric Symptoms									
2	RCTs	High	Not serious	Not serious	Not Serious	Very small samples, incomplete reporting	Very low ●○○			
Hospitali	izations									
Harms										
2	RCTs	High	Not serious	Not Serious	Not Serious	Very small samples, incomplete reporting	Very low ●○○			
Function	Function or Quality of Life for Patient									
Function	or Quality	of Life for	Parent							
Tanction	Quality	or Life 101	Tarent							

Abbreviation. RCT: randomized controlled trial.

Quality Assessment (Confidence in Estimate of Effect) for Plasma Exchange										
No. of Studies	Study Design(s)	Risk of Bias	Inconsistency	Indirectness	Imprecision	Other Factors	Level of Confidence			
Change i	Change in Psychiatric Symptoms									
1	RCT	High	Not serious	Not serious	Not Serious	Very small sample, incomplete reporting	Very low ●○○			
Hospitali	izations									
Harms										
1	RCT	High	Not serious	Not Serious	Not Serious	Very small sample, incomplete reporting	Very low ●○○			
Function	Function or Quality of Life for Patient									
Function	or Quality	of Life for	Daront							
Function or Quality of Life for Parent										

Abbreviation. RCT: randomized controlled trial.

Appendix C. Methods

Scope Statement

Populations

Children diagnosed with:

- Pediatric Autoimmune Neuropsychiatric Disorders Associated with Streptococcal Infections (PANDAS),
- Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS)

Population scoping notes: Patients without either of the above conditions are excluded

Interventions

Therapeutic plasma exchange; intravenous immunoglobulin (IVIG); antibiotics; tonsillectomy and/or adenoidectomy

Intervention exclusions: Behavioral interventions, selective serotonin reuptake inhibitors, nonsteroidal inflammatory drugs

Comparators

Usual care or other interventions

Outcomes

Critical: Change in psychiatric symptom scores (e.g., Children's Yale-Brown Obsessive Compulsive Scale, Clinical Global Impressions-Improvement, Yale Global Tic Severity scale); Hospitalizations, including institutionalization or emergency visits

Important: Harms; standardized measures of function or quality of life for patients and caregivers

Considered but not selected for the GRADE table: None

Key Questions

KQ1: What is the effectiveness of treatments for PANDAS/PANS as compared to the named comparators?

KQ2: Does the comparative effectiveness of treatments for PANDAS/PANS differ by:

- a. Patient characteristics
- b. Condition characteristics
- c. Intervention
- d. Provider characteristics (e.g., Center of Excellence)

KQ3: What are the harms of interventions for PANDAS/PANS in children?

Contextual Questions

CQ1: What are the evidence-based criteria available for the diagnosis of PANDAS/PANS, and what are the diagnostic accuracy of available criteria or tests?

Search Strategy

A full search of the core sources was conducted to identify systematic reviews, meta-analyses, and technology assessments that meet the criteria for the scope described above. Searches of core sources were limited to citations published after 2015.

The following core sources were searched:

Agency for Healthcare Research and Quality (AHRQ)

Canadian Agency for Drugs and Technologies in Health (CADTH)

Cochrane Library (Wiley Online Library)

Institute for Clinical and Economic Review (ICER)

Medicaid Evidence-based Decisions Project (MED)

National Institute for Health and Care Excellence (NICE)

Tufts Cost-effectiveness Analysis Registry

Veterans Administration Evidence-based Synthesis Program (ESP)

Washington State Health Technology Assessment Program

An Ovid MEDLINE® search was also conducted to identify systematic reviews, meta-analyses, and technology assessments, using the search terms paediatric autoimmune neuropsychiatric disorder associated with streptococcal infection, pediatric autoimmune neuropsychiatric disorder associated with streptococcal infection, pediatric acute-onset neuropsychiatric syndrome, pediatric infection triggered autoimmune neuropsychiatric disorder, childhood acute onset neuropsychiatric syndrome, paediatric acute-onset neuropsychiatric syndrome, autoimmune encephalitis. The search was limited to publications in English published since 2012. In addition, a MEDLINE® search was conducted for randomized controlled trials and comparative cohort studies.

Searches for clinical practice guidelines were limited to those published since 2015. A search for relevant clinical practice guidelines was also conducted using MEDLINE® and the following sources:

Australian Government National Health and Medical Research Council (NHMRC)

Canadian Agency for Drugs and Technologies in Health (CADTH)

Centers for Disease Control and Prevention (CDC), Community Preventive Services

National Institute for Health and Care Excellence (NICE)

Scottish Intercollegiate Guidelines Network (SIGN)

United States Preventive Services Task Force (USPSTF)

Veterans Administration/Department of Defense (VA/DoD) Clinical Practice Guidelines

Inclusion/Exclusion Criteria

Studies were excluded if they were not published in English, did not address the scope statement, or were study designs other than systematic reviews, meta-analyses, technology assessments, randomized controlled trials, comparative cohort studies, or clinical practice guidelines.

Appendix D. Applicable Codes

Coding note: PANS does not have ICD-10-CM index entries; PANDAS is indexed to D89.89.

CODES	DESCRIPTION	
ICD-10-CM Codes		
D89.89	Other specified disorders involving the immune mechanism, not elsewhere classified	
D89.9	Disorder involving the immune mechanism, unspecified	
G04.81	Other encephalitis and encephalomyelitis	
CPT Cod	es	
	Behavioral therapy	
90832	Psychotherapy, 30 minutes with patient	
90833	Psychotherapy, 30 minutes with patient when performed with an evaluation and management	
90033	service (List separately in addition to the code for primary procedure)	
90834	Psychotherapy, 45 minutes with patient	
90836	Psychotherapy, 45 minutes with patient when performed with an evaluation and management	
	service (List separately in addition to the code for primary procedure)	
90837	Psychotherapy, 60 minutes with patient	
90838	Psychotherapy, 60 minutes with patient when performed with an evaluation and management	
	service (List separately in addition to the code for primary procedure)	
90839	Psychotherapy for crisis; first 60 minutes	
	Intravenous immunoglobulin therapy	
90283	Immune globulin (IVIG), human, for intravenous use	
96365	Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); initial, up to 1 hour	
96366	Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); each	
	additional hour (List separately in addition to code for primary procedure)	
99601	Home infusion/specialty drug administration, per visit (up to 2 hours)	
26544	Plasma exchange	
36514	Therapeutic apheresis; for plasma pheresis	
42020	Tonsillectomy and adenoidectomy	
42820	Tonsillectomy and adenoidectomy; younger than age 12	
42821 42825	Tonsillectomy and adenoidectomy; age 12 or over	
42825	Tonsillectomy, primary or secondary, younger than age 12 Tonsillectomy, primary or secondary, age 12 or over	
42830	Adenoidectomy, primary or secondary, age 12 or over	
42831	Adenoidectomy, primary, younger than age 12 Adenoidectomy, primary; age 12 or over	
42835	Adenoidectomy, secondary; younger than age 12	
42836	Adenoidectomy, secondary; younger than age 12 Adenoidectomy, secondary; age 12 or over	
HCPCS Level II Codes Intravenous immunoglobulin therapy		
J1459	Injection, immune globulin (privigen), intravenous, non-lyophilized (e.g., liquid), 500 mg	
J1555	Injection, immune globulin (cuvitru), 100 mg	
J1556	Injection, immune globulin (bivigam), 500 mg	
J1557	Injection, immune globulin, (gammaplex), intravenous, non-lyophilized (e.g., liquid), 500 mg	
J1558	Injection, immune globulin (xembify), 100 mg	
J1559	Injection, immune globulin (hizentra), 100 mg	
J1561	Injection, immune globulin, (gamunex-c/gammaked), non-lyophilized (e.g., liquid), 500 mg	
J1562	Injection, immune globulin (vivaglobin), 100 mg	
J1566	Injection, immune globulin, intravenous, lyophilized (e.g., powder), not otherwise specified, 500 mg	

J1568	Injection, immune globulin, (octagam), intravenous, non-lyophilized (e.g., liquid), 500 mg		
J1569	Injection, immune globulin, (gammagard liquid), non-lyophilized, (e.g., liquid), 500 mg		
J1572	Injection, immune globulin, (flebogamma/flebogamma dif), intravenous, non-lyophilized (e.g., liquid), 500 mg		
J1599	Injection, immune globulin, intravenous, non-lyophilized (e.g., liquid), not otherwise specified, 500 mg		
S9338	Home infusion therapy, immunotherapy, administrative services, professional pharmacy services, care coordination, and all necessary supplies and equipment (drugs and nursing visits coded separately), per diem		
	SSRIs, NSAIDs, and corticosteroids		
J1094	Injection, dexamethasone acetate, 1 mg		
J1100	Injection, dexamethasone sodium phosphate, 1 mg		
J0702	Injection, betamethasone acetate 3 mg and betamethasone sodium phosphate 3 mg		
J1700	Injection, hydrocortisone acetate, up to 25 mg		
J1710	Injection, hydrocortisone sodium phosphate, up to 50 mg		
J1720	Injection, hydrocortisone sodium succinate, up to 100 mg		
J2650	Injection, prednisolone acetate, up to 1 ml		
J7510	Prednisolone oral, per 5 mg		
J7512	Prednisone, immediate release or delayed release, oral, 1 mg		
J8540	Dexamethasone, oral, 0.25 mg		
J7624	Betamethasone, inhalation solution, compounded product, administered through DME, unit dose form, per mg		
J1130	Injection, diclofenac sodium, 0.5 mg		

Note. Inclusion on this list does not guarantee coverage.

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Public Comments		Δ:
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Discussion Table

IDs/#s	Summary of Issue	Draft Subcommittee Response
A1, C1, D1, G3, I1, J1, K1, L1, M1, N1, O1, Q1, R1, S1, U1	The proposed requirement to have two pediatric subspecialists evaluate a patient and recommend IVIG and/or plasmapheresis would be difficult to implement due to access issues and may prevent timely access to care. A patient's primary care provider should be considered as one of the two clinicians who determines eligibility for IVIG and/or plasmapheresis. A primary care provider in Oregon can include naturopaths.	For EbGS discussion: Due to the complex nature of these conditions and the need to rule out serious alternative diagnoses, at least 1 pediatric subspecialist should be involved in the patient's care. Such subspecialist involvement will ensure the best possible diagnostic and treatment plan for these severely affected children. Adolescents may also be adequately evaluated by an adult subspecialist who feels comfortable caring for their age group. However, to address concerns raised regarding access, EbGS can consider modifying the 2-subspecialist consultation requirement to 1 subspecialist in addition to the recommendation of the primary care provider, who could be a physician, naturopath, nurse practitioner, etc. If the requirement is reduced to 1 subspecialist, then the consultation should be in-person or via teleconsultation (not provider-to-provider or e-consult). The relevant clause would be modified as follows: Option 1: A consultation with and recommendation by 2 pediatric subspecialists (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist). One of these





IDs/#s	Summary of Issue	Draft Subcommittee Response
		consultations may consist of a provider-to-provider consultation or e-consultation. Option 2: b) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult. Similarly, the reevaluation clause could be modified to include a reevaluation
		 by the primary care provider and 1 subspecialist: Option 1: A reevaluation at 3 months by both pediatric experts is required for continued therapy of IVIG or plasma exchange. Option 2: A reevaluation at 3 months by both the primary care provider and subspecialist is required for continued therapy of IVIG or plasma exchange.
A3	The proposed criteria do not account for a level of severity of illness which would allow for immediate access to IVIG or plasmapheresis.	In instances in which a patient faces incapacitating or life-threatening illness, the patient is best treated in the hospital environment. Hospital-level care is beyond the scope of this coverage guidance.
A4	More clarification is needed regarding the requirement that a patient try and fail two therapies prior to being considered for IVIG and/or plasmapheresis, including 1) the duration of a trial of therapy and 2) how to determine if the less intensive therapy is not effective.	The duration of a trial of therapy will differ by type of therapy. For example, an appropriate course of antibiotics may be a 14-day course while an appropriate trial of an SSRI might be 6 weeks. Additionally, these therapies may be tried concurrently. However, as these therapies have few serious side effects and may be effective, the subcommittee feels that a trial of such less intensive therapy prior to therapy escalation is important to the care of children with PANDAS/PANS.





IDs/#s	Summary of Issue	Draft Subcommittee Response
		The trial of less intensive therapy would be considered to have not been effective when no significant clinical improvement was found on whatever objective clinical instrument is used for the most concerning clinical presenting symptom(s), for example a scale for rating OCD symptoms or for rating depression symptoms. Staff have also noted in discussion with experts that a clinical improvement that is nonsustained, such as with a course of steroids, is also an indication for IVIG or plasmapheresis.
		For EbGS discussion: Consider modifying the current requirement wording to clarify what is
		considered a "trial" of an alternative therapy. Option 1: At least 2 less-intensive therapies (for example, appropriate limited course of nonsteroidal antiinflammatory drugs [NSAIDs], corticosteroids, selective serotonin reuptake inhibitors [SSRIs], behavioral therapy, short-course antibiotic therapy) have been tried and were not effective.
		Option 2: A clinically appropriate trial of at least 2 less-intensive treatments (for example, appropriate limited course of nonsteroidal antiinflammatory drugs [NSAIDs], corticosteroids, selective serotonin reuptake inhibitors [SSRIs], behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of significant improvement on a validated objective instrument directed at the patient's primary symptom complex). These trials may be performed concurrently.





IDs/#s	Summary of Issue	Draft Subcommittee Response
		For EbGS discussion:
		Similarly, the reevaluation at 3 months should be done with a validated objective instrument.
		Option 1: This evaluation must include objective clinical testing, which must be performed pretreatment and posttreatment to demonstrate significant clinical improvement.
		Option 2: This evaluation must include objective clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate significant clinical improvement.
A1, A5, C1,	There is concern about workforce training, education, and	Training and education of providers is beyond the scope of the HERC. It is also
H1, I1, L1, M1, N1, O1,	willingness to see patients with PANDAS/PANS symptoms, as	beyond the scope of the HERC to require providers to see certain patients or
Q1, R1, S1	well as lengthy wait times to access appointments with	prescribe certain treatments. Likewise, workforce issues regarding the
	subspecialists.	number of specialists and access to specialists is beyond the scope of the
		HERC, though it is important context for HERC decisions.
		Previously suggested modifications (see above) would allow for use of
		telemedicine to accomplish visits and consultations, and would improve
		access to specialists for patients in rural or underserved areas. Other
		modifications would allow adolescents to be evaluated by adult subspecialists
		which should also help to address the access to care issue.
D1, E1, F1,	Description of personal stories of affected children and	We thank you for taking the time to share your and your loved ones'
G1, H1, I1,	caregiver experience.	experiences and stories. Such real-life stories add needed context to the
J1, K1, M1, N1, O1, Q1,		subcommittee's deliberations.
S1		While individual stories provide context for the subcommittee's decisions, the
		subcommittee makes coverage decisions on a population-level basis and





IDs/#s	Summary of Issue	Draft Subcommittee Response
		must base these decisions on evidence and other factors with respect to the
		population in general.
G2	There is concern about the credentials of the appointed	For this topic, OHA selected 2 Oregon experts knowledgeable of local practice
	experts for this topic.	patterns and standards of care regarding PANDAS/PANS. One out-of-state
		expert was nominated by the PANS Consortium as a subject matter expert.
		The subcommittee places a preference on local expertise to shape care
		guidelines that will affect Oregon patients, payers and providers.
A2, B7	The coverage exclusion of other interventions (for example,	Our review found published evidence from 2 comparative observational
	tonsillectomy) does not align with PANS Consortium (advocacy	studies and 2 systematic reviews indicating that tonsillectomies are not
	organization) treatment guidelines, and is supported by	effective for reducing neuropsychiatric symptom severity or exacerbations in
	unpublished studies.	children with a PANDAS diagnosis. Unpublished literature is not eligible for
		inclusion in the coverage guidance evidence review.

Commenters

Identification	Stakeholder Stakeholder	
Α	Sarah Lemley, Executive Director, Northwest PANDAS/PANS Network [Submitted December 7, 2021]	
В	Dritan Agalliu, PhD; Associate Professor; Departments of Neurology, Pathology and Cell Biology; Columbia University Medical Center [Submitted December 8, 2021]	
С	Donna Kirchoff, MD, Integrative Developmental & Behavioral Pediatrics [Submitted December 9, 2021]	
D	Deborah Miller, parent/caregiver of a child with PANDAS [Submitted December 9, 2021]	
E	Heather Winkeljohn, parent/caregiver of a child with PANDAS/PANS [Submitted December 9, 2021]	
F	Jennifer Rowan, parent/caregiver of a child with PANDAS/PANS [Submitted December 10, 2021]	
G	Cathy Daraee, RN, grandparent of a child with PANDAS/PANS [Submitted December 13, 2021]	
Н	Meggan Bennett, parent/caregiver of a child with PANDAS/PANS [Submitted December 15, 2021]	
1	Carrie Ann McGowan, parent/caregiver of a child with PANDAS/PANS [Submitted December 17, 2021]	
J	Andrea Jones, RN, and parent/caregiver of a child with PANDAS/PANS [Submitted December 17, 2021]	





K	Kamiar Daraee, grandparent of a child with PANDAS/PANS [Submitted December 20, 2021]	
L	Lara Winn, MSW student clinician, Collective Care Clinic [Submitted December 23, 2021]	
M	Christina and Ivan Vejar, parents/caregivers of a child with PANDAS/PANS [Submitted January 3, 2022]	
N	Jeremy Johnson, relative of a child with PANDAS/PANS [Submitted January 5, 2022]	
0	Jessica Johnson, relative of a child with PANDAS/PANS [Submitted January 5, 2022]	
Р	Ivan Vejar, Board Member, Northwest PANDAS/PANS Network [Submitted January 5, 2022]	
Q	Carly Absher, parent/caregiver of a child with PANDAS/PANS [Submitted January 6, 2022]	
R	Kym McCornack, NWPPN Oregon Outreach Coordinator [Submitted January 6, 2022]	
S	Sara E. Zeman, JD, LLM, and parent/caregiver of a child with PANDAS/PANS [Submitted January 6, 2022]	
Т	Paul Terdal, management consultant, Terdal Consulting LLC [Submitted January 6, 2022]	
U	Paul Ryan, President, PACE Foundation [Submitted January 6, 2022]	

Public Comments

ID/#	Comment	Disposition
A1	Option #1 Guidance Concerns: The revised HERC guidance unnecessarily perpetuates barriers to care by requiring 2 physician specialists attest to the need for IVIG and/or Plasma Exchange when the list of specialists specified by the Evidence Based SubCommittee are NOT those who are well versed in treating this population and in fact, 2 of the 4	Thank you for your comments. We have written specific responses to individual sections of your public comment in the rows that follow. Due to the complex nature of these conditions and the
	specialties identified REFUSE TO SEE THESE PATIENTS. These requirements would further perpetuate health inequity in our state, especially those who live in rural areas or who don't have the resources needed to access 2 specialists (location, transportation, trauma, time off work, etc).	need to rule out serious alternative diagnoses, at least 1 pediatric subspecialist should be involved in the patient's care. Such subspecialist involvement will ensure the best possible diagnostic and treatment plan
	Oregon already faces a critical shortage of PANDAS/PANS-competent providers, especially within the restricted list of specialties identified in the revised HERC report. These requirements would inappropriately exclude many disciplines and providers with PANDAS/PANS expertise who are currently caring for this population and are well versed in the diagnostic and treatment guidelines for care.	for these severely affected children. Additionally, adolescents may be adequately evaluated by an adult subspecialist who feels comfortable caring for their age group.





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A2	Imposing cumbersome requirements in order to access medically necessary care only compounds the health disparities that already exist for these disorders. We would ask the HERC EbGS Committee to remove the 2-physician consultation and recommendation requirement for accessing IVIG or Plasma Exchange. The removal of this requirement will ensure inclusivity of medical providers with PANDAS/PANS expertise who are currently being excluded by the drafted HERC requirement language and allow for more timely access to care and treatment interventions. Alternatively, drop the requirement to one physician specialist which will alleviate some of the barrier and still ensure you have some oversight. Please note, accessing consultation with Pediatric Neurologists and Pediatric Psychiatrists, especially those who take Oregon Health Plan, can be 6 months to a year out for a new patient. In addition, the exclusion of tonsillectomy coverage overlooks PANDAS Physician	Our review found published evidence from 2
AZ	In addition, the exclusion of tonsillectomy coverage overlooks PANDAS Physician Network's guidance on this issue: "Many PANS/PANDAS patients have damaged or cryptic tonsils, but the potential benefit of tonsillectomy is not limited to patients with those tonsil characteristics. In an unpublished research study done at Georgetown Medical Center, PANDAS patients had their tonsils removed, analyzed, and the children subsequently tracked for over six months. The tonsils relative to non-PANDAS patients had many pathogens, most prominent being staphylococcus (staph). Streptococcus pyogenes was not found in PANDAS patients but was present in non-PANDAS controls. Other notable pathogens included MRSA, E. coli, Pseudomonas and Serratia marcens. The absence of Streptococcus in the PANDAS cohort suggests that once the patient has been "sensitized" other pathogens can induce neurologic symptoms in susceptible patients. In addition, the tonsils belonging to PANDAS patients contained elevated levels of TH17, indicating a consistent immune response to the pathogens lodged within the tonsils. TH17 has been found in animal PANDAS research to be a potential agent for opening the blood brain barrier, allowing inflammation in targeted regions of the brain.	Cur review found published evidence from 2 comparative observational studies and 2 systematic reviews indicating that tonsillectomies are not effective for reducing neuropsychiatric symptom severity or exacerbations in children with a PANDAS diagnosis. Unpublished literature, such as the referenced Georgetown study, is not eligible for inclusion in the coverage guidance evidence review.





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	The Georgetown study and physician experience indicates that removal of the tonsils can provide remission of PANS & PANDAS symptoms for some patients. There is no marker to determine which patient a tonsillectomy will result in remission of PANS/PANDAS symptoms. A clear benefit of tonsillectomy that was found in the Georgetown study and further observed by practitioners who see many PANS/PANDAS patients, is that those PANS/PANDAS cases that have undergone tonsillectomy, have a significantly lower chance of recurrence post-immunotherapy such as IVIG. Since immunotherapy suppresses the potential cause of basal ganglia encephalitis and in some cases like IVIG "reboots" the immune response, then removing a consistent infectious trigger housed within the tonsil or removing a repository for new pathogen agitators would most likely be beneficial." We would also ask for the committee to take more time to understand the importance of tonsillectomy in these disorders by utilizing a subject expert who has been on faculty at Georgetown for more than 25 years (Dr. Earl Harley) and has put forth written comments and will be available during the hearing. He can also be reached at [email redacted].	
A3	Lastly, these requirements dismiss the PANS Consortium treatment guidelines which indicate the necessity to progress to these lines of treatment (IVIG and Plasma Exchange) for those patients whose severity is "incapacitating, life threatening, or occupy 71%-100% of waking hours". Delaying needed medical care for a child suffering from a severe presentation of PANDAS/PANS because of these imposed requirements would be inappropriate in a life-threatening situation. We would ask the HERC EbGS Committee to remove the requirement of two failed therapies before proceeding to IVIG or PE as treatment guidelines published by the PANS Consortium and PANDAS Physician Network recommend treatment based on severity. To force a child with life-threatening presentation of these disorders to	In instances in which a patient faces incapacitating or life-threatening illness, the patient is best treated in the hospital environment. Hospital-level care is beyond the scope of this coverage guidance. We reviewed both the PANS/PANDAS Clinical Research Consortium treatment guidelines and the Nordic Pediatric Immunopsychiatry guidelines and included a discussion in the coverage guidance (pg. 32-36). Both guidelines were rated as having a high risk of methodological bias. Authors of the PANS/PANDAS Consortium guidelines recognized that there was





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	fail less-intensive therapies first can result in irreversible health outcomes because of delayed medical care.	disagreement regarding proposed treatments among contributing clinicians and that the recommended treatment pathway is subject to amendment pending further evidence.
A4	We would also request clarification on: The duration required of the less-intensive therapies (NSAIDS, corticosteroids, SSRIS, therapy and/or antibiotics)? How would effectiveness or failed effectiveness of these therapies be measured and by whom? The reevaluation post 3 months of IVIG or PE, what specific 'objective clinical testing' would be utilized and how would providers be trained to ensure they are using it consistently?	The duration of a trial of therapy will differ by type of therapy. For example, an appropriate trial of antibiotics may be a 14-day course, while an appropriate trial of an SSRI might be 6 weeks. Additionally, these therapies may be tried concurrently. As these therapies have few serious side effects and may be effective, the subcommittee feels that a trial of such less intensive therapy prior to therapy escalation is important to the care of children with PANDAS/PANS.
		The trial of less intensive therapy would be considered to have not been effective when no significant clinical improvement was found on whichever validated instrument is used for the most concerning clinical presenting symptom(s), for example, a scale for rating OCD symptoms or for rating depression symptoms.





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A5	Why the Committee would not trust current clinicians to prescribe medically	Training and education of providers is beyond the
	necessary care and instead, require referral to specialists that lack expertise,	scope of the HERC. It is also beyond the scope of the
	experience and acceptance of these disorders with access issues that would only	HERC to require providers to see certain patients or
	delay needed medical care?	prescribe certain treatments. Likewise, workforce
	How the Committee would provide training and education to the identified	issues regarding the number of specialists and access
	specialists and ensure their willingness to treat this population given the historical	to specialists is beyond the scope of the HERC.
	reports of almost all Oregon families having to seek medical care outside of these	
	specialties and even outside of our state as there is a lack of expertise locally?	
B1	My laboratory has a long-standing interest in understanding the role of the	Thank you for your comments. We have written
	adaptive immune system (Th17/Th1 lymphocytes) in autoimmune central nervous	specific responses to individual sections of your letter in
	system (CNS) sequelae following recurring S. pyogenes (Group A Streptococcus;	the rows that follow.
	GAS) infections, specifically how CD4+ T cell subtypes induce neurovascular	
	damage, neuroinflammation and neuronal circuit dysfunction in animal models for	
	PANDAS/PANS. We have shown that S. pyogenes-specific Th17 cells from the nose	
	enter the olfactory bulb and brain via olfactory sensory axons, where they damage	
	the structural and functional integrity of the blood-brain barrier (BBB) and allow	
	autoantibodies to enter the brain. Recently, we have demonstrated that while the	
	loss of excitatory synapses in the olfactory bulb is transient after multiple	
	infections, there is a persistent functional deficit in odor processing and neuronal	
	function. Moreover, using mice that lack Th17 lymphocytes, we have shown that	
	they are critical for selective CNS entry of autoantibodies across the blood-brain	
	barrier, microglial activation and neural circuit impairment during post-infectious	
	basal ganglia encephalitis. We are currently working to further understand the	
	origin of inflammatory chemokines and cytokines present in the blood of children	
	with PANDAS/PANS and elucidate how they affect the pathology in the brain.	
	Furthermore, my laboratory has significant expertise regarding the	





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	neuroimmunological mechanisms that promote blood-brain barrier breakdown and	
	immune cell infiltration into the CNS in Experimental Autoimmune	
	Encephalomyelitis, a mouse model for multiple sclerosis. As a basic scientist, I have	
	the required expertise to evaluate the literature in the field of PANDAS/PANS and	
	to provide an independent assessment whether the "literature-based evidence"	
	presented to the HERC Evidence-Based Guideline Subcommittee via the HERC	
	Review of the Literature and Coverage Guidance for PANDAS/PANS/Pediatric AE on	
	September 9, 2021 (Pages 70 - 117) is accurate.	
	After reviewing the HERC document in detail, I regret to inform the committee	
	that: a) the literature presented in the document is INCOMPLETE and b) the papers	
	presented as evidence against treatment do not accurately reflect the conclusions	
	of the study. I have attached a rebuttal outlining the missing literature and I have	
	attached the PDFs for some of the recent studies that the committee needs to take	
	into consideration.	
	Pediatric Acute-onset Neuropsychiatric Syndrome (PANS) is characterized by the	
	abrupt and dramatic onset of obsessive-compulsive symptoms, restricted intake of	
	food or fluids (sometimes to the point of starvation or dehydration), anxiety,	
	depression and suicidality, emotional lability, personality changes, sensory	
	hypersensitivity, cognitive deficits and physical symptoms, such as arthralgias,	
	urinary dysfunction, and severe insomnia. As its name implies, PANS affect	
	children, primarily those aged 4 - 9 years. When Group A streptococcal infections	
	(such as strep throat) triggers symptoms, the disorder is known as Pediatric	
	Autoimmune Neuropsychiatric Disorders Associated with Streptococcal Infections	
	(PANDAS). Recently, a number of studies have demonstrated that PANS/PANDAS is	
	a form of autoimmune encephalopathy—or inflammation of the brain. Below, I	
	outline several critical old and recent basic and clinical studies that demonstrate a	





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very strong association of GAS with PANDAS and treatment strategies for PANDAS	
and PANS.	
and PANS. Basic studies in animal models of PANDAS/PANS have demonstrated that both cellular (Th17 lymphocytes) and humoral (antibodies) adaptive immunity, generated in response to multiple GAS infections, target the brain and trigger neuroinflammation, blood-brain barrier damage, neuroinflammation and neuronal dysfunction (Brimberg et al., 2012; Dileepan et al., 2016; Hoffman et al., 2004; Platt et al., 2020; Yaddanapudi et al., 2010). These animal models have been focused on demonstrating the ability of GAS to prime development of an autoimmune reaction by stimulating adaptive cellular and humoral immune responses. In the mouse, intranasal (i.n.) infections with live bacteria polarize T cells located in the nasal-associated lymphoid tissue (NALT, the mouse structural analog of human tonsils and adenoids) toward a Th17 phenotype, a T cell subtype that is both essential for mucosal immune protection against bacteria but also strongly implicated in many autoimmune diseases. Multiple i.n. S. pyogenes infections strengthen this Th17 immune response, largely due to induction of IL-6 and TGF-β1, which are two proinflammatory cytokines essential for Th17 differentiation. This model has been used to demonstrate that repeated i.n. infections with S. pyogenes induce migration of GAS-specific Th17 cells and other T cell subtypes from the nasal epithelium to the olfactory bulb (OB), where sensory axons make connections with projection interneurons to form the neural circuitry essential for odor discrimination, as well as to other CNS regions (Dileepan et al., 2016). The presence of Streptococcus-specific Th17 cells in the CNS after repeated i.n. infections increases the permeability of capillaries in several CNS regions thereby enabling	We reviewed the references provided in this public comment. Animal model studies are not eligible for inclusion in HERC guidance documents as they provide indirect evidence and are outside of the scope of the coverage guidance evidence review. Your work to address the evidence gaps is helpful and may motivate others to perform more rigorous human subjects research on these conditions. However, the subcommittee uses only peer-reviewed studies regarding important clinical outcomes performed on human subjects and generally requires between-group comparison for evidence of treatment effectiveness.
	very strong association of GAS with PANDAS and treatment strategies for PANDAS and PANS. Basic studies in animal models of PANDAS/PANS have demonstrated that both cellular (Th17 lymphocytes) and humoral (antibodies) adaptive immunity, generated in response to multiple GAS infections, target the brain and trigger neuroinflammation, blood-brain barrier damage, neuroinflammation and neuronal dysfunction (Brimberg et al., 2012; Dileepan et al., 2016; Hoffman et al., 2004; Platt et al., 2020; Yaddanapudi et al., 2010). These animal models have been focused on demonstrating the ability of GAS to prime development of an autoimmune reaction by stimulating adaptive cellular and humoral immune responses. In the mouse, intranasal (i.n.) infections with live bacteria polarize T cells located in the nasal-associated lymphoid tissue (NALT, the mouse structural analog of human tonsils and adenoids) toward a Th17 phenotype, a T cell subtype that is both essential for mucosal immune protection against bacteria but also strongly implicated in many autoimmune diseases. Multiple i.n. S. pyogenes infections strengthen this Th17 immune response, largely due to induction of IL-6 and TGF-β1, which are two proinflammatory cytokines essential for Th17 differentiation. This model has been used to demonstrate that repeated i.n. infections with S. pyogenes induce migration of GAS-specific Th17 cells and other T cell subtypes from the nasal epithelium to the olfactory bulb (OB), where sensory axons make connections with projection interneurons to form the neural circuitry essential for odor discrimination, as well as to other CNS regions (Dileepan et al., 2016). The presence of Streptococcus-specific Th17 cells in the CNS after repeated i.n. infections





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	control an essential aspect of blood-brain barrier function. The intranasal model	
	produces profound changes in olfactory neural circuitry by reducing excitatory	
	input at the presynaptic terminals of olfactory sensory axons and perturbing the	
	excitatory/inhibitory balance within the primary olfactory circuit. This model of	
	post-S. pyogenes autoimmunity demonstrates a central role for the cellular	
	adaptive immune response (for example, bacterial-specific Th17 cells in the CNS) in	
	disrupting blood-brain barrier function, thus promoting entry of antibodies into the	
	CNS and inducing changes in synaptic signaling. Although such a cellular adaptive	
	immune response has not been identified to date in the nervous systems of	
	children suffering from BGE, S. pyogenes-specific Th17 cells can be found in the	
	tonsils of human patients (Dileepan et al., 2016), making Th17 lymphocytes a	
	potential causative agent in either initiation or persistence of disease pathogenesis.	
	Moreover, mice that lack Th17 lymphocytes, have reduced blood-brain barrier	
	damage and antibody entry into the CNS, reduced microglial activation and	
	preservation of neural circuit function in the mouse model for PANDAS (Platt et al.,	
	2020). Since Th17 lymphocytes are critical for pathogenesis in multiple	
	autoimmune diseases such as Multiple Sclerosis, Lupus, and Psoriasis and they are	
	also necessary for disease pathogenesis in rodent models for PANDAS (Platt et al.,	
	2020), these findings suggest a critical requirement for the adaptive cellular	
	immune response in PANDAS pathogenesis in addition to the role of the humoral	
	immune response.	
	A second group of rodent models for BGE employs subcutaneous immunization	
	with an antigenic target (bacterial homogenate) plus complete Freund's adjuvant	
	to activate the immune system, in conjunction with agents (i.e., B. pertussis toxin)	
	that open the BBB to provide access to brain targets. In this model, mice and rats	
	develop a strong humoral immune response toward S. pyogenes and show	





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	behavioral abnormalities. Specifically, GAS-immunized rodents display increased	
	rearing and decreased locomotion, as well as increased repetitive and	
	perseverative behaviors, impaired pre-pulse inhibition, and reduced concentrations	
	of serotonin in the prefrontal cortex as compared to controls (Brimberg et al.,	
	2012; Hoffman et al., 2004; Yaddanapudi et al., 2010). Moreover, adoptive transfer	
	of serum IgGs from S. pyogenes-immunized mice to naive recipient mice, or direct	
	infusion of sera into rat brains, recapitulates some of the behavioral deficits in	
	recipient rodents, whereas no effects were observed after adoptive transfer of IgG-	
	depleted serum. Histological examination of brain tissue revealed antibody	
	deposition in the deep cerebellar nuclei and hippocampus in mice and the	
	striatum, cortex, and thalamus in rats (Brimberg et al., 2012; Hoffman et al., 2004;	
	Yaddanapudi et al., 2010). Serum IgG isolated from immunized rodents recognizes	
	both cerebellar targets and human D1/ D2 dopamine receptors by either western	
	blotting or ELISA. Thus, the subcutaneous animal models for BGE have provided	
	useful information regarding the humoral immune response after bacterial	
	infection (i.e., the presence of antibodies directed against GAS and CNS) and	
	demonstrate a clear link between S. pyogenes exposure and behavioral	
	abnormalities.	
	A comprehensive review of animal models for PANDAS/PANS is discussed in a	
	recent review that we published in Frontiers in Immunology (Platt et al., 2017).	
	Overall, the summary outlined above from studies in animal models of	
	PANDAS/PANS provides a very strong evidence that there is an immune-based	
	component (cellular and humoral) underlying the pathogenesis of disease that	
	should be taken into serious consideration for recommendation to treat children	
	suffering from PANDAS/PANS. I want to remind the committee that the mouse	
	model for multiple sclerosis, Experimental Autoimmune Encephalomyelitis, which	





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	has some similarities to the human MS, has been proven very valuable to	
	understand a) the mechanisms of human MS and b) develop treatments for the	
	disease. Therefore, the committee should take into consideration the animal	
	studies for PANDAS/PANS for their decision.	
В3	Studies in sera of patients suffering from Sydenham's chorea, PANDAS and PANS	We reviewed the references you provided and
	have identified anti-neuronal autoantibodies targeting the basal ganglia, including	determined these studies are ineligible for inclusion in
	the D1 and D2 dopamine receptors and recently cholinergic interneurons (Cox et	the coverage guidance report, given that these are
	al., 2013; Dale et al., 2012; Kirvan et al., 2003; Kirvan et al., 2006; Sinmaz et al.,	basic science research papers, animal model studies,
	2015; Xu et al., 2020). These antibodies induce neuronal dysfunction in vitro	and/or do not report clinical outcomes that are
	(Kirvan et al., 2003; Xu et al., 2020) and elicit behavioral abnormalities in rodents	relevant to the evidence review. We recognize the
	after adoptive transfer [reviewed in (Platt et al., 2017)], suggesting a critical role for	important role of basic science research, but coverage
	the humoral immune response in the pathogenesis of these diseases. Moreover,	decisions regarding interventions require evidence of
	the titer of these pathological antibodies is reduced in the sera of Sydenham's	clinically important benefits in humans.
	chorea, PANDAS or PANS patients during the convalescence period that	
	corresponds with improved symptomatology (neurological and psychiatric	
	manifestations) (Chain et al., 2020; Xu et al., 2020). Finally, a recent clinical study	
	has shown that in 41 pediatric subjects, followed for over a 24-month period, 65%	
	of new GAS [Group A streptococcal] infections caused no symptoms, yet these	
	subjects developed antibodies against GAS suggesting that the majority of GAS	
	infections are not detected in clinic (Hysmith et al., 2017). This could result in	
	missed opportunities for primary prevention of rheumatic fever and rheumatic	
	heart disease, Sydenham's chorea or PANDAS with appropriate antimicrobial	
	therapy.	
	In addition to the presence of D2R antibodies, a recent study (Pilli et al, 2020)	
	characterized the presence of proinflammatory D2R-specific T cells in movement	
	and psychiatric disorders in 24 cases and 16 controls. D2R-specific T cells were	





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	identified by flow cytometric quantification of CD4+CD25+CD134+ T cells. Cytokine secretion was analyzed using a cytometric bead array and ELISA. HLA genotypes were examined in D2R-specific T-cell-positive patients. D2R antibody seropositivity was determined using a flow cytometry live cell-based assay. The study identified three immunodominant regions of D2R that specifically activated CD4+ T cells in 8/24 patients. Peptides corresponding to these regions were predicted to bind with high affinity to the HLA of the eight positive patients and had also elicited the secretion of pro-inflammatory cytokines IL-2, IFN-g, TNF, IL-6, IL-17A and IL-17F. Therefore, autoreactive D2R-specific T cells and a proinflammatory Th1 and Th17 cytokine profile characterize a subset of pediatric patients with movement and psychiatric disorders, further underpinning the theory of immune dysregulation in these disorders.	
B4	Recently, two large epidemiological cohort studies of children in Europe (N=1,068,000) (Orlovska et al., 2017) and Asia (N=28,600) (Wang et al., 2016) reported that children hospitalized with GAS infections had a 96% higher risk of neuropsychiatric disorders (Taiwan) (Wang et al., 2016), 51% higher risk for obsessive-compulsive disorder (OCD) and a 35% higher risk for tic disorders (Denmark) (Orlovska et al., 2017). These recent epidemiological studies together with previous findings that more than 25% of pediatric cases presenting with obsessive-compulsive disorders (OCD) and tic disorders (for example, Tourette syndrome) originate as PANDAS (Swedo et al., 1998) strongly argue for a critical role of recurrent GAS infections in the etiology of PANDAS or PANS and that these diseases are rare similar in incidence to Lupus.	Thank you for providing this background. We reviewed these references and determined that, while they are out of scope for inclusion in the evidence review since they do not evaluate the effectiveness of potential treatments for PANDAS/PANS, they do provide helpful epidemiologic context. Therefore, the Orlovska et al., 2017 and Wang et al., 2016 references have been added to the coverage guidance background section after the submission of this comment.
B5	PANDAS and PANS cases are increasingly being classified as a form of Autoimmune Encephalitis. The Mayo Clinic conducted a study in 2018 warning that more than 90,000 Autoimmune Encephalitis cases are being missed on an annual basis	After reviewing the literature on autoimmune encephalitis, the subcommittee determined that autoimmune encephalitis is a life-threatening condition





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	worldwide (Dubey et al., 2018). We contend that many PANDAS and PANS cases fall within that category as recently discussed in detail in studies published in the American Academy of Neurology (Cellucci et al., 2020) and Lancet Psychiatry (Pollak et al., 2020). Furthermore, PANDAS and PANS are now considered as a form of basal ganglia encephalitis demanding attention and urgent care, as argued in recent editorial by esteemed physicians in Immunology, Neurology & Psychiatry of PANDAS/PANS. (Dale et al., 2017).	that could be associated with an acute infection or presence of a tumor and is, therefore, acutely treated in a hospital setting. The specific diagnostic criteria currently in broad clinical use for autoimmune encephalitis is unique and has biological markers. Because of the differences in diagnostic criteria and disease processes between autoimmune encephalitis and PANDAS/PANS, the subcommittee voted to exclude autoimmune encephalitis from the scope of this report in its September 9, 2021 meeting.
B6	A recent Italian study (Murgia et al., 2021) used a metabolomics approach to identify a specific metabolic pattern in patients affected by PANS compared to healthy subjects. Thirty-four outpatients referred for PANS and 25 neurotypical subjects matched for age and gender, were subjected to metabolite analysis. The study found a unique plasma metabolic profile in PANS patients, significantly differing from that of healthy children, that suggests the involvement of specific patterns of neurotransmission (tryptophan, glycine, histamine/histidine) as well as a state of neuroinflammation and oxidative stress in the disorder. This metabolomics study offers new insights into biological mechanisms underpinning the disorder and supports research to identify potential biomarkers implicated in PANS.	We reviewed this reference and determined that it is out of scope for inclusion in the coverage guidance as it does not evaluate any of the prespecified interventions or clinical outcomes for the evidence review. See also response to B3.
В7	The NIMH PANS consortium formed by a large number of experts from the disciplines of pediatrics, infectious disease, neurology, immunology and psychiatry have published the guidelines for treatment of PANDAS/PANS which rely on antibiotic therapy, steroids, IVIG, and psychiatric treatments (Thienemann et al., 2017; Frankovich et al., 2017: Cooperstock et al., 2017). The PANS Research	Thank you for your comment. The coverage guidance currently contains a summary of the PANDAS/PANS Clinical Research Consortium guidelines based on the same sources cited in this section.





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	Consortium has based its diagnosis and treatment guidelines on their experience of managing more than 1,000 patients in the U.S. The majority of the children are under age 13 and those who are left untreated can suffer dire consequences into young adulthood, including suicide. Treatment of PANS/PANDAS involves a three-pronged approach that utilizes psychiatric medications to provide symptomatic relief, antibiotics to eliminate the source of neuroinflammation and immune-modulating therapies to treat disturbances of the immune system. When these therapies are instituted promptly, many children recover completely and return to full functioning. Delays in obtaining treatment not only prolong the child's suffering needlessly but also increase the risk that the PANS/PANDAS symptoms will become entrenched, leading to long-term psychiatric, neurologic, and cognitive dysfunction.	
B8	A Stage 3 Clinical Trial of IVIG will be conducted in January 2021, "A Superiority Study to Compare Panzyga Versus Placebo in Patients with PANS," Clinical Trials.gov, NCT04508530 in both Europe and USA in approximately 200 children to examine the effectiveness of IVIG in PANDAS and PANS children in a larger cohort.	Thank you for your comment. We are currently monitoring the status of this trial for future consideration.
C1	I would like to provide written testimony regarding Medicaid coverage of specific treatments for PANDAS/PANS for children who have Medicad [sic] insurance. My understanding is that one coverage option under consideration is that in order to qualify for PE or IVIG, a Medicaid patient would require both documentation of failed treatment (which I think is very reasonable) AND would REQUIRE 2 MD Subspecialists (Pediatric Neurology, Rheumatology, Psychiatry or Neurodevelopmental) to consult and recommend treatment (which I do not think is reasonable). Unfortunately, the requirement to have 2 specific subspecialists recommend treatment creates an unnecessary and cumbersome barrier to appropriate care, as	Thank you for your comment. Please see the response to A5. Additionally, while it is outside of the scope of this coverage guidance review process to specify or include a pool of eligible providers who treat a specific condition in its report, we have listed examples of the types of providers, such as naturopaths, that can provide consultations for PANDAS/PANS in our draft Coverage Guidance recommendation.





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	our state not only lacks access to physicians in these specialties (a 12+ month wait	
	to see both clinicians would be very common), but there are also only a very small	
	number of physicians within those specialties that are willing to evaluate or treat a	
	child for PANS/PANDAS. The language also excludes currently treating clinicians	
	(Pediatricians, Naturopaths and Nurse Practitioners) who may have the required	
	expertise to diagnose and treat PANS/PANDAS. Unlike AZ and CA, OR does not have	
	CPAE/PANS clinic to serve this need.	
	PANS/PANDAS is an extremely challenging disorder to diagnose and treat, and	
	when it is severe enough that IVIG and/or PE is indicated, is devastating for not	
	only the children who are impacted but for their entire families, due to the	
	extreme impact it has on mood and behavior. Please consider changing the	
	coverage language to allow a wider range of clinicians to verify this diagnosis and	
	recommend IVIG/PE treatment, as the current language will essentially make	
	IVIG/PE impossible for Medicaid patients to get in a timely manner.	
D1	I am the mother of a 12 year old son with PANDAS. I am opposed to requiring 2	Thank you for taking the time to share your and your
	physicians to approve IVIG for children with PANDAS/PANS.	loved ones' experiences and stories. Such real-life
	My son, [name redacted], is not able to get the treatment that he needs despite	stories add needed context to the subcommittee's
	the fact that there is a National Standard of Care for children suffering from	deliberations.
	PANDAS and PANS.	
	Despite the fact that there are multiple multi disciplinary institutions across the	
	country with clinics dedicated to treating children with PANDAS/PANS using this	
	national standard of care with success. Making it a requirement for two physicians	
	to approve IVIG will only make it harder for my son and other children suffering	
	from PANDAS/PANS to get the treatment that they require to heal.	
	It is unethical to withhold treatment for these children and leave them to suffer	
	when there is, once again, a National Standard of Care.	





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	My son is a victim of a broken health care system in the state of Oregon that has	
	left him a shell of the child he should be because he does not have access to the	
	lifesaving treatment that he requires.	
	I say lifesaving because my son has spoken multiple times about wanting to kill	
	himself. He would rather be dead then deal with the effects of PANDAS. Tics,	
	uncontrollable rage, hallucinations, black out episodes, OCD, extreme anxiety, baby	
	talking and loss of friendships to name a few.	
	[Name redacted]'s story does not have to end this way. His story can end happily if	
	he is given access to the treatment that he requires to save him from irreversible	
	brain damage. [Name redacted] needs IVIG ever 4 weeks for a recommended 5-6	
	months at a cost of over \$5500 a round. It is sickening that the state of Oregon add	
	another road block with a 2 physician requirement for IVIG. It is sickening that the	
	state of Oregon would leave my son to suffer rather than giving access to	
	treatment.	
	My 12 year old son has been suffering with PANDAS for 10 long years. Enough is	
	enough. He deserves to know what a healthy childhood looks like. We should be	
	removing roadblock to treatment not adding them. I plead with you to make	
	treatment for children suffering from PANDAS/PANS accessible so that other	
	children like [name redacted] won't be left to suffer and can live a happy healthy	
	life.	
E1	Your decision and failure to recognize PANDAS/PANS as a legitimate condition is	Our review found published evidence from 2
	not only damaging but it is dire and detrimental to the health and wellness of	comparative observational studies and 2 systematic
	children. Six years ago, here in New Mexico, I heard those words that any	reviews indicating that tonsillectomies are not effective
	PANDAS/PANS parents hates to hear from a provider, "PANDAS/PANS is	for reducing neuropsychiatric symptom severity or
	controversial." Luckily, I did not accept that and pursued proper treatment from a	exacerbations in children with a PANDAS diagnosis.
	well versed Internal Medicine Physician and also from the PANDAS clinic in Tucson,	Unpublished literature is not eligible for inclusion in the





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	Arizona. Thankfully, my child has improved but not without lasting effects. Her pediatrician was knowledgeable about the disease, but the Neurologist was not. We were luckily in a position to pay for our child's treatments, one of which involved tonsillectomy. Many families are not so lucky. EARLY treatment and recognition is key. You must stop this draconian, shameful and archaic belief. Look at the research out of some of the leading Universities in America. Put your egos aside and look at the SCIENCE!	coverage guidance evidence review. We hope that future research provides effective treatments available to children experiencing these symptoms. Thank you for taking the time to share your and your loved ones' experiences and stories. Such real-life stories add needed context to the subcommittee's deliberations.
F1	I am writing to express my deep distress that a Center for Neurodevelopmental Pediatrics in Oregon is refusing to treat children with PANS/PANDAS. I implore you to please work to remove barriers to care for our children. My son developed PANS at the age of 4. My well-adjusted, happy boy became a completely different child overnight, losing language and social awareness, developing extreme OCD and anorexia, and attempting suicide twice, among other terrifying symptoms. Getting appropriate treatment was a frightening, exhausting ordeal. A psychologist and colleague first identified his symptoms as consistent with PANS, and encouraged me to seek medical treatment. His initial diagnosis by our local doctor was 'autism with psychosis', which makes zero sense for a child who was neurotypical a week ago. After a night at OHSU which included a spinal tap, MRI and various other tests, no diagnosis was offered, except that I was an anxious mom who was causing his symptoms. At the third hospital we visited, Randall's, we finally met with reason. The doctor we saw said, "In the absence of another diagnosis I am willing to treat this as PANS." We went through all of that simply to get my son antibiotics, and we are so lucky that antibiotics and other over the counter treatments were all he needed to heal. Two days after his first dose his symptoms improved dramatically, and now at age eight years he is happy, healthy	Thank you for taking the time to share your and your loved ones' experiences and stories. Such real-life stories add needed context to the subcommittee's deliberations.





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	and doing well in all respects. He has had setbacks however, and I live with the fear that I will lose him again and be unable to afford or access the care he will need to regain his personality and lead a normal life. It is a cloud that haunts us always. For those families whose children need treatment now, my heart is utterly broken. An estimated 1:200 children are affected by PANS/PANDAS. Will your child, or grandchild, niece or nephew be one of them? I wish that upon no one, but I also wish you could walk in our shoes for just a day so you would know how critical it is for our children to be assured access to care. Mitigate the personal and societal costs of lifelong psychiatric treatment, special education, and unemployment. Make the sensible choice and do everything possible to remove barriers to care. Insist that our hospitals recognize PANS/PANDAS as the very real disorder it is, and demand that they provide the treatment our children need. Thank you.	
G1	I am an Oregonian, having lived here for over 40 years. I am also a retired Registered Nurse having been in practice for 40 + years. I understand the importance of evidence-based medicine. We must acknowledge science evolves. Learning should never stop and we should always seek to understand without doing harm. Just because the medical community in Oregon is behind the curve on the scientific evidence behind PANDAS/PANS and the efficacy of treatment, does not signify its non existence.	Thank you for your comments. We agree that science evolves, and the subcommittee supports making coverage decisions based on the best currently-available evidence, and updating policy when new evidence becomes available. We have written specific responses to individual sections of your letter in the rows that follow.
G2	The lack of true expertise in our state is evidenced by the 2 out of the 3 subjects experts chosen to advice this subcommittee on PANDAS/PANS. Medical ethics should require that an "expert" is well vetted when contributing to the decision making process specific to access of medically needed care. This is especially pertinent when those 2 subject experts have something to gain by being labeled an "expert" in a region that lacks treating providers.	Thank you for your comment. For this topic, OHA selected two Oregon experts that were knowledgeable of local practice patterns and standards of care regarding PANDAS/PANS. One out-of-state expert was nominated by the PANS Consortium as a subject matter expert. The subcommittee places a preference on local expertise to





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	This is not only misleading, but appears unethical to accept someone as a expert who lacks the true experience needed to contribute to such important conversations. The NW, specifically Oregon, is negligently behind the science when it comes to providing care to children and youth with post - infectious storms of encephalitis. One only has to look at the fact that 8 other states have mandated coverage for their children to understand the dire need parents face in lobbying for access to critically necessary care. I am the Grandmother to an Oregon child with PANS. The HERC's role is the review of clinical evidence in order to guide the OHA in making benefit-related decisions for its health plans. You have the evidence. Your local non-profit has brought in far superior PANDAS/PANS expertise then any "self-proclaimed expert" on your panel (other then Dr. Daines). Dr. Agaillu wrote wrote a very detailed rebuttal of the evidence presented during the first hearing while identifying the lack of evidence included and considered pertinent to this decision.	shape care guidelines that will affect Oregon patients, payers and providers. We have responded to Dr. Agalliu's comments during this public comment period; see responses to B1-B8.
G3	The HERC's role is NOT to impose additional barriers to care by requiring 2 subspecialists in order to access IVIG. You do RECOGNIZE THAT A LACK OF SUPPORTIVE AND ACCESSIBLE subspecialists exist in the disciplines you identified. There are NO Rheumatologists willing to treat this population. The Neuro developmental center at OHSU REFUSES TO TREAT THIS POPULATION. While access issues are not in your scope, do NOT IMPOSE IMPOSSIBLE HURDLES TO CARE for children who are suffering from brain inflammation and families who are suffering levels of trauma equal to that of caring for an Alzheimer's patient.	Thank you for your comment. See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms. See response to A1 regarding the 2-subspecialist requirement. See response to C1 regarding provider eligibility for treatment consultation.





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	Please listen to your community members affected by this disorder. You have the evidence that IVIG can result in symptom reduction or alleviation for moderate to severe children with PANDAS/PANS.	
	PLEASE CHANGE THE GUIDANCE TO ONLY REQUIRE 1 PHYSICIAN SUBSPECIALIST and for this to include pediatric or adult subspecialists since access in many parts of Oregon struggle with the presence of any of the sub specialties.	
	Also, in reference to those in rural areas and a lack of subspecialists, please allow for a physician to physician consult or file review to ensure a timely turnaround on a critical medical treatment Lastly, please consider the inclusion of a Naturopathic Physician in the list of subspecialists.	
	The LEADING, TREATING MEDICAL PROVIDER IN OUR STATE, who head of Children Psychiatry at OHSU and the Medical Director at Unity Psych Hospital refer patients to a NATUROPATHIC PHYSICIAN in our state who has more expertise in treating these cases the 2 of the 3 "experts" on your committee.	
H1	I am writing you in hopes that you will limit the road blocks for children with PANDAS/PANS to receive IVIG. My son has seen plenty of specialty providers to include a neurologist and psychiatrist. Neither of these providers acknowledged my son's illness as being legitimate. My son was put on psychiatric medication which almost killed him as it caused serotonin syndrome. The only providers that have offered us any help is our Pediatrician who ordered IVIG when I had private insurance (it was denied coverage) and our ND. I currently pay out of pocket for Ozone infusions, supplements, and hyperberic [sic] chamber treatments.	Thank you for taking the time to share your and your loved ones' experiences and stories. Such real-life stories add needed context to the subcommittee's deliberations. Ozone infusions, hyperbaric oxygen therapy and supplements are outside the scope of this coverage guidance.
	Requiring "specialists" or potentially harmful medications for a child causes a delay in life saving treatment and is an unnecessary roadblock. In our State there are no experts when it comes to PANDAS/PANS. I implore you to listen to us as parents	See response to A1 regarding the 2-subspecialist requirement.





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	when we say we our children deserve the right to treatment to save them, to solve the underlying problem, what they do not need is a band-aid to mask the symptoms or potentially make them worse. A pediatrician or ND is perfectly capable of making the decision on the health care needs of our children. I appreciate your time.	See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms. See response to C1 regarding provider eligibility for treatment consultation.
II	I am writing with concerns about your guidelines for the treatment of PANS/PANDAS (referred to as PANS most of the time from this point). First, as a parent of a child with PANDAS, I have had a very hard time finding treatment options in Oregon. It is my understanding that you have made it a requirement for children who have not responded adequately to the first level treatments of PANS to have the acquiescence of two specialists from delineated specialties to access the more advanced immunotherapies. Two of these specialties have no specialists who are willing to even consider seeing a PANS patient (developmental pediatricians and rheumatologists). The other two have few specialists that are willing or competent enough to see PANS patients and as a result, they are hard to get into. It has taken us 2.5 years to assemble a team to treat our daughter—in the meantime, she has lost IQ points, dropped from above average to the 25th percentile in working memory, dropped from average to the 2nd percentile for processing speed, developed dyspraxia, a tremor, and various other movement abnormalities, and spent years stewing in anger, rage, OCD, and misery. My 7-year-old daughter is losing her childhood while we try to get her the treatment she needs. Steroids are not a permanent solution. Your guidelines are simply a barrier to treatment and will lead to suffering and permanent damage to the children and families dealing with this condition. In addition to affecting the children on OHP, it will trickle out to affect how private insurance treats PANS patients. We currently	Thank you for taking the time to share your and your loved ones' experiences. Such real-life stories add needed context to the subcommittee's deliberations. See response to A1 regarding the 2-subspecialist requirement. See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms. See response to C1 regarding provider eligibility for treatment consultation. See response to G2 regarding the selection of appointed experts for this coverage guidance report.





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	have the full endorsement for IVIG from 3 different specialists (neurologist,	
	rheumatologist-we had to leave the state to find one, psychiatrist, and then also	
	our PANS specialist) on your list and we still cannot gain pre-approval. There are	
	implications beyond just what happens with OHP. Insurance companies will look at	
	this as tacit approval to deny treatment to desperately sick children.	
	I believe that you likely have made this a requirement because you have been	
	misled by people who have labeled themselves "experts" when they have no	
	qualifications to do so. These people flout the research done by those actually	
	researching and treating PANS, to offer their own unfounded opinions about what	
	PANS is, the way it is treated, and the state of PANS treatment in Oregon. You	
	should be looking for actual renowned experts, not people who call themselves	
	experts but do not treat PANS patients regularly and are out of step with the	
	research and conduct none themselves. Talk to Dr. Latimer in DC or Dr. Kovacevic	
	in Chicago. Talk to Dr. Swedo. I cannot stress this enough, your experts arenot	
	experts. I speak from personal experience.	
	I will be frankour family is struggling. I mean, really, really struggling, due to the	
	lack of understanding of these sorts of conditions. I cannot even begin to count	
	how many times I have felt hopeless, despondent, and emotionally destroyed by	
	this disorder and the lack of affordable treatment options. My daughter is the	
	center of my universe and I'm losing her. I'm losing her due to the failures of our	
	medical system to understand, study, and treat this condition. The future she has	
	now is bleak without further treatment. Please do not leave children and their	
	families in this position. Give them options. Include PANS practitioners on your list	
	of specialists. Lower your requirements for treatment1 specialist should be	
	enough. This is such an awful illness made even worse by the barriers to care and	
	the seeming indifference of those in power to the suffering of children. I would be	





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J1	happy to send you audio recordings of my daughter during some of her more mild flares. The more intense episodes require too much to have the time or thought to record her. I don't think you realize what this condition is like. If you did, I firmly believe that you would be moved to help these children. Please reconsider. I am a former pediatric nurse of over 15 years and have worked at Randall	Thank you for taking the time to share your and your
	Children's Hospital for that duration with some of the best doctors in the United States. I am well educated, and resourceful, however nothing could prepare me for my daughters diagnosis with PANDAS 3 years ago. I am adept at advocating for patients and know the lingo with physicians, and yet it was nearly impossible to find a physician who would look into possible treatment for my daughter. We have amazing insurance, but because of the rulings previously made about pans/pandas, only 5% of her treatments are actually covered by insurance. We have had to refinance our home 3 times just to cover the tens of thousands of dollars in medical bills we accrue each year for her illness. Why am I sharing this with you? Because this committee is ruling to have 2 subspecialists to approve for the treatment of IVIG. IVIG is LIFE CHANGING for pandas and is currently the gold standard nationwide. My question for you is, are two subspecialists required to sign off for IVIG in the case of Kawasaki's? (Another rare childhood illness that causes the body to attack itself). If not, than this added measure will create yet another impossible barrier for parents to jump over. I implore you- follow the research. Finding qualified providers who understand PANDAS and are willing to touch it is hard enough- don't make it even harder for parents who are already exhausted, poor and in desperate need.	loved ones' experiences. Such real-life stories add needed context to the subcommittee's deliberations. See response to A1 regarding the 2-subspecialist requirement.





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K1	I understand your committee is imposing requirements for children on Medicaid that would make accessing care near impossible. These cumbersome requirements are unnecessary. Oregon lacks true experience within the medical communities when it comes to diagnosing and treating PANDAS/PANS. This is why it was necessary for my Granddaughter to travel to Southern California to be evaluated and treated by a PANDAS/PANS expert. To impose restrictions that require TWO subspecialists to recommend treatment is impossible in this state. If the committee desires oversight, ADJUST THE SUBSPECIALIST RECOMMENDATION TO ONE REQUIRED INSTEAD OF TWO. These families are traumatized. They are barely getting by day by day. Many on Medicaid may lack the resources that other families navigate support and care for their severely psychiatric child. Requiring that a traumatized parent, who has probably struggled to find a treating provider has to obtain TWO additional recommendations for care, and timely treatment is VITAL when addressing a child with brain inflammation is not realistic. This type of requirement, imposed on those who are less resourced, is inequitable and the committee should take this into consideration. TO IMPOSE REQUIREMENTS WITHOUT RECOGNIZING THEIR ILL EFFECT IS NEGLIGENT. I ask that the committee change the requirement to ONE subspecialist and include naturopathic physician to those who can be consulted with. The subspecialist you list are unsupportive and not willing to treat this population.	Thank you for taking the time to share your and your loved ones' experiences. Such real-life stories add needed context to the subcommittee's deliberations. See response to A1 regarding the 2-subspecialist requirement. See response to C1 regarding provider eligibility for treatment consultation.
L1	As someone who works with patients with PANS/PANDAS, I have significant concerns that the requirements imposed by the committee will increase barriers to care.	Thank you for your comment. See response to A1 regarding the 2-subspecialist requirement.





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	Requiring that 2 MD subspecialists recommend IVIG and/or plasma exchange means that virtually no families will qualify because Oregon has a critical shortage of PANS/PANDAS competent providers.	See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms.
	These requirements would also exclude providers from other disciplines who do have PANS/PANDAS experience. I urge you to reconsider these requirements because they increase the substantial barriers to care that families suffering with PANS/PANDAS already face.	
M1	We have suffered for 6 years (diagnosis at age 6), to help my child live a live free from neuropsychiatric symptoms that have prevented her from going to school, sleeping, eating, and performing the activities of daily life - while trying to get help from specialists in the state of Oregon. There are NO medical sub-specialists in the state of Oregon who are experts in diagnosing and treating PANDAS/PANS - no rheumatologists, neurologists, psychiatrists, who work in tandem to support the "guardrails" that these HERC guidelines are imposing on us. We live in HELL with our child on a daily basis, trying whatever we can to help and support her while trying to find insurance coverage for IVIG. We have waiting LONG ENOUGH! We are unable to own a home; we live on a single income because my child has been unable to consistently attend school due to her illness (adding on the anxiety of living for 2+ years in a pandemic, being completely isolated from the world, due to worries that contracting COVID would exacerbate her autoimmune disorder); and trying to manage our own anxiety and PTSD of living with a child who could effectively be treated with IVIG, but being prevented from doing so by the impossible standards of insurance companies. Children with pediatric cancer or juvenile rheumatoid arthritis are NOT denied treatment (including IVIG), while our children are ignored.	Thank you for taking the time to share your and your loved ones' experiences. Such real-life stories add needed context to the subcommittee's deliberations. See response to A1 regarding the 2-subspecialist requirement. See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms.





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	Please consider changing these guidelines. The system is NOT being abused. There	
	are not lines of children waiting outside clinics for IVIG. It should be for children like	
	mine, who suffer daily FOR YEARS, and run the risk of permanent brain and bodily	
	damage, due to the constant flaring of her untreated autoimmune disorder.	
N1	I understand your committee is imposing requirements for PANDAS/PANS children	Thank you for taking the time to share your and your
	on Medicaid that would make accessing medical care near impossible. These	loved ones' experiences. Such real-life stories add
	cumbersome requirements are unnecessary.	needed context to the subcommittee's deliberations.
	Oregon already lacks true expertise within the medical communities when it comes	See response to A1 regarding the 2-subspecialist
	to diagnosing and treating PANDAS/PANS. This is why many of these families who	requirement.
	have the financial resources to do so, travel outside of the state to be evaluated by	See response to A5 regarding provider willingness to
	a competent and supportive PANDAS/PANS expert.	treat patients with PANDAS/PANS symptoms.
	Your guidance report, as it stands now with the requirement to somehow find 2 specialists supportive enough to see PANDAS/PANS patients, is impossible in this state. Many of these families, depending on where they live in Oregon, cannot find one medical provider in their area knowledgeable enough or willing to see these cases. If the committee desires oversight, adjust the subspecialist recommendation to one required instead of two.	See response to C1 regarding provider eligibility for treatment consultation.
	Our family was traumatized by this disorder. Previously normal lives were turned upside down by the devastating onset of psychiatric symptoms in a previously healthy child. The grief and trauma sustained by these disorders should not be further burdened by state imposed restrictions that do nothing but create additional barriers to medical care.	
	Many on Medicaid may lack the resources that other families have to navigate support and care for their severely psychiatric child. Requiring that a traumatized parent, who has already probably struggled to find a treating provider, have to	





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	obtain TWO additional recommendations for care when timely treatment is vital, is	
	not realistic. This type of requirement, imposed on those who are less resourced, is	
	inequitable and the committee should take that into consideration. To impose	
	requirements without recognizing their ill effect is negligent.	
	I ask that the committee change the requirement to ONE subspecialist requirement	
	and include naturopathic physicians, nurse practitioners and physician assistants in	
	those who can be consulted with. The subspecialists you list are unsupportive and	
	not willing to treat this population.	
01	I understand your committee is imposing requirements for PANDAS/PANS children	Thank you for taking the time to share your and your
	on Medicaid that would make accessing medical care near impossible. These	loved ones' experiences. Such real-life stories add
	cumbersome requirements are unnecessary.	needed context to the subcommittee's deliberations.
	Oregon already lacks true expertise within the medical communities when it comes	See response to A1 regarding the 2-subspecialist
	to diagnosing and treating PANDAS/PANS. This is why many of these families who	requirement.
	have the financial resources to do so, travel outside of the state to be evaluated by a competent and supportive PANDAS/PANS expert.	See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms.
	Your guidance report, as it stands now with the requirement to somehow find 2 specialists supportive enough to see PANDAS/PANS patients, is impossible in this state. Many of these families, depending on where they live in Oregon, cannot find	See response to C1 regarding provider eligibility for treatment consultation.
	one medical provider in their area knowledgeable enough or willing to see these	
	cases. If the committee desires oversight, adjust the subspecialist recommendation	
	to one required instead of two.	
	Our family was traumatized by this disorder. Previously normal lives were turned	
	upside down by the devastating onset of psychiatric symptoms in a previously	
	healthy child. The grief and trauma sustained by these disorders should not be	





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	further burdened by state imposed restrictions that do nothing but create additional barriers to medical care.	
	Many on Medicaid may lack the resources that other families have to navigate support and care for their severely psychiatric child. Requiring that a traumatized parent, who has already probably struggled to find a treating provider, have to obtain TWO additional recommendations for care when timely treatment is vital, is not realistic. This type of requirement, imposed on those who are less resourced, is inequitable and the committee should take that into consideration. To impose requirements without recognizing their ill effect is negligent.	
	I ask that the committee change the requirement to ONE subspecialist requirement and include naturopathic physicians, nurse practitioners and physician assistants in those who can be consulted with. The subspecialists you list are unsupportive and not willing to treat this population.	
P1	In regards to the current draft guidance on PANDAS/PANS, Oregon's only PANDAS/PANS non-profit, Northwest PANDAS/PANS Network, along with the PACE Foundation and a number of leading PANS/CPAE clinics and centers from around the country, including University of Arizona, Stanford, Harvard/Massachusetts General, Dartmouth and University of Arkansas strongly recommend the below revised verbiage for item 2b of the current HERC guidance. These clinics and centers are leaders in this management of these disorders and are the most well versed clinicians when it comes to best practices. Their recommendation to change the HERC's verbiage should be weighed heavily and given the strongest consideration for effective implementation and oversight of the treatments being considered.	Thank you for your comment. The proposed amendment submitted to modify item 2b is neither feasible nor enforceable, as OHA does not have the authority to compel providers of any scope or specialty to respond to consultation requests.





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	Proposed Draft 2b) A consultation with and recommendation by 1 pediatric	
	subspecialist (for example, neurologist, psychiatrist, neurodevelopmental	
	specialist, immunologist, infectious disease, rheumatologist). Consultation can be	
	provider-to-provider in-person, or via an e-consultation (including a file review). If	
	no response is received within 96 hours from consulting subspecialist, once	
	information received, then item 2b of this HERC requirement is deemed to be	
	waived.	
	We hope that Oregon will benefit from the guidance that is being provided by	
	these leading PANS/CPAE clinics and centers, especially given that our region lacks	
	any such expertise.	
	Their efforts, experience and dedication to treating this population should	
	absolutely be recognized in the absence of any equivalent.	
	Thank you for your consideration on this extremely important matter and	
	recognizing the utter importance of modifying the language in line 2b.	
Q1	I am writing as the mother of two daughters diagnosed with post-infectious	Thank you for taking the time to share your and your
	encephalopathies (NMDAR and PANS/PANDAS) in Washington. These disorders	loved ones' experiences. Such real-life stories add
	have the same type of acute onset, extremely similar symptomsjust different	needed context to the subcommittee's deliberations.
	antibodies responsible for the suffering they inflict. IVIG and Super high dose	See response to A1 regarding the 2-subspecialist
	steroids are bottom tier treatment options for NMDAR encaphalitisand it is	requirement.
	covered by insurance.	·
	Because of a complete lack of awareness, or perhaps willful and criminal ignorance	See response to A5 regarding provider willingness to
	of the latest research concerning PANS/PANDAS by NW Higher Education	treat patients with PANDAS/PANS symptoms.
	institutions, a total lack of providers who treat these horrifying disorders, and	
	because suffering families already experience extreme duress & medical	
	discrimination while trying to access healing and relief for their children the HERC	
	Committee's requirement of 2-sub-specialists to verify a child's need for IVIG or	





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	plasmapheresis is simply adding to the caseloads of the VERY few doctors who do	
	successfully treat it and this requirement will continue to send families traveling	
	outside the Pacific NW to access treatment for their children. And worst of all,	
	families like mine will have to pay exhorbitant fees to heal their children.	
	All providers who are genuinely knowledgable of PANS/PANDAS go through a	
	tiered treatment process of NSAIDS, steroids, antibiotics and anti-virals, and many	
	other immune suppressing medications before ever considering IVIG for their	
	patients. Once all else has failed, only then does a knowledgable PANS/PANDAS	
	physicians AND naturopaths prescribe the use of IVIG. Only about 10-20% of	
	PANDAS patients require this treatment. 1 in 200 kids have PANDAS/PANS. There	
	are roughly over 4,000 PANS/PANDAS patients (mostly undiagnosed) in	
	Oregonthat means this committee could change the lives of approximately 400	
	Oregon children and their families.	
	My NDMAR daughter was misdiagnosed for 4 years. Because doctors did not	
	understand the signs and symptoms of her disease. Same is true for my	
	PANS/PANDAS daughter who suffered misdiagnosis for 6 years: she was prescribed	
	unnecessary and ineffective psychiatric medications, endured 4 inpatient	
	psychiatric stays, 1 long term residential treatment stay out of state, unsuccessful	
	wrap-around servicesand a host of other horrifying problems that I cannot begin	
	to describe in an email, before we traveled to Chicago to visit a PANDAS specialist	
	who saw her dire need for proper medical intervention. He immediately ordered	
	steroids and high dose IVIG for my daughter. After losing over 6 years of her	
	childhood, she finally began healing with relief of symptoms. Today, she does not	
	take a single psychiatric medication.	
	We paid for her IVIG completely out of pocket.	





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R1	Her current state of health is saving my insurance company huge sums of money every year. She will graduate from high school this year and attend Portland State University. This is no less than a miracle for our family - we never imagined a future was possible for her and it wasn't possibleuntil IVIG. Do not make the mistake of causing more harm to families who desperately need help to heal their children by adding more obstacles to their children's paths to healing. What PANDAS/PANS families endure is unimaginable to clinicians and to the rest of the world. Please do the right thing: help our children who suffer the most with no relief, who have no hope of a future. How much is a life worth? I'm writing about PANDAS/PANS treatment. I very much appreciate the work you have done to date and understand you want to provide medically necessary care to this population. Allowing patient to access IVIG as part of treatment for PANDAS/PANS is a huge step to ensure the best outcome for children who need that level of care. It also aligns more closely with the National Standard of Care for the treatment of PANDAS/PANS. Working at Northwest PANDAS/PANS Network the last 3+ years has allowed me to become very familiar with the treating landscape in Oregon. There are very few providers treating this population across all disciplines. Because of this, the requirement of 2 subspecialist to access IVIG will act as a roadblock. I urge you to revise the guidance to allow access with 1 subspecialists. Also, removing pediatric as to include adult subspecialists would be helpful for rural areas that may not have a pediatric subspecialist.	Thank you for your comments in support of the recommendation for coverage. See response to A1 regarding the 2-subspecialist requirement. Additionally, adolescents may be adequately evaluated by an adult subspecialist who feels comfortable caring for their age group. See response to A5 regarding provider willingness to treat patients with PANDAS/PANS symptoms.
S1	I am a retired health care attorney who's spent the last year singularly focused on helping my PANS/PANDAS-diagnosed child. I am writing to urge you NOT to require two specialists' opinions for Medicaid coverage of IVIG treatment. I further urge you not to impose any specialist requirement but allow coverage where generalists	Thank you for taking the time to share your and your loved ones' experiences. Such real-life stories add needed context to the subcommittee's deliberations.





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	prescribe this treatment. I will describe what my family has endured to	See response to A1 regarding the 2-subspecialist
	demonstrate what barriers to care already exist and why your proposal creates	requirement.
	additional, virtually impassible barriers.	See response to A5 regarding provider willingness to
	The amount of trauma this illness has inflicted upon my child and family cannot be	treat patients with PANDAS/PANS symptoms.
	overstated. Our daughter went from being a bright, empathic, highly functional, socially-engaged child—a gifted/talented student, award-winning artist, former symphony member, "the life of the party," etc., to a child who's so emotionally labile and psychologically distraught that she runs out of the house barefoot at night, intent on throwing herself into traffic. She's so adversely mentally affected at times that she has not recognized her own parents. She's been unable to even attend school for the past year. She barely ever leaves her dark bedroom and cannot tolerate lights, sounds, smells. She spends her days trapped in unending cycles of elaborate OCD rituals that are a hallmark of PANS. As parents feeling helpless to break our child free from this suffering—we've experienced nothing worse. Indeed, my whole family suffers immense, ongoing trauma living with the psychiatric chaos this disease has wrought. There has been no stability, no predictability, no security, living with this illness. Further, my child is aware of her suffering and of its impact on our family; she begs us almost daily to 'let her die.'	treat patients with PANDAS/PANS symptoms. See response to C1 regarding provider eligibility for treatment consultation.
	We've been living through hell, and we've had no hope that our beloved child	
	would ever return, or that any of us could live a normal life again. Not, that is, until	
	about four months ago, when we found the Doctor of Naturopath who is now actually treating our daughter. With this doctor, we have finally felt hope, because,	
	at last, an experienced, well-regarded clinician understands what's happening to	
	our child, is confident in her ability to treat her, and is committed to seeing her	
	through to recovery. Indeed, my daughter's severe psychiatric illness has been	
	refractory to all standard treatments, save the protocols for PANS/PANDAS that	





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	this doctor has been leading us through. We anticipate and discuss that my child	
	likely will undergo IVIG therapy soon. Because Medicaid coverage decisions tend to	
	influence private insurers' and medical practices' policies, your proposed	
	restriction on IVIG therapy has the potential to directly impede my child's	
	treatment and recovery and to seriously harm my family's finances, health, and	
	welfare.	
	1. Pediatric mental health care is in extremely short supply in Oregon, which delays	
	crucial PANS/PANDAS treatment.	
	Advocates of the proposed rule assert it is easy to access specialists. However, that	
	was not my experience. Like most families with a child with PANS/PANDAS, we	
	began our search for care with our pediatrician and standard psychiatric care and	
	therapy. This was prior to the pandemic, but, even then, psychiatric appointments	
	were near impossible to obtain, with practices closed or having months-long wait	
	lists. As it became clear that our child's needs exceeded the scope of typical	
	outpatient services, we quickly learned of the inadequacy of mental health services	
	for Oregon children. As our child's crisis grew more urgent, her doctors failed to	
	respond appropriately; her treating psychologist even terminated her services,	
	saying our child needed a higher level of care (of course, none was readily	
	available). Despite our best efforts, we found practice after practice closed to new	
	patients; program wait lists months-long or even closed. OHSU said it would take a	
	year to get a Neurocognitive evaluation. We learned adolescent psychiatric beds	
	are virtually nonexistent—only 20 beds for all of Oregon's adolescents. Our	
	daughter spent one week essentially warehoused in an ER, waiting for inpatient	
	psychiatric care that never materialized. She was kept in virtual solitary	
	confinement and provided with no psychiatric care despite being in crisis. Referrals	
	to specialty mental health care dissolved as program directors and intake	





ID/#	Comment	Disposition
	coordinators advised our child was not within their purview. None of the mental	
	health providers we saw recognized our child's severe PANS/PANDAS crisis; this	
	went on for months.	
	2. Medical specialists in Oregon are not experienced with PANS/PANDAS	
	treatment.	
	Unfortunately, my husband (a university professor) and I have found that medical	
	and psychological specialists in Oregon likewise are not experienced in recognizing,	
	diagnosing, or treating PANS/PANDAS. Despite discussing our child's refractory-to-	
	treatment OCD in clinic visits with pediatric specialists in Psychiatry, Neurology,	
	Neurosurgery, Psychology, Neurocognitive Psychology, Pediatric Emergency	
	Medicine, Otolaryngology, and Gastroenterology, none recognized her	
	PANS/PANDAS crisis. None offered any diagnostic work-up, treatment, or referral	
	for PANS/PANDAS. One specialist even canceled scheduled treatment due to our	
	child's deteriorating mental status, despite that being a standard sign of	
	PANS/PANDAS. Our exhaustive search for help belies that Oregon lacks a supply of	
	specialists ready, willing, and able to treat PANS/PANDAS. Our Naturopath is the	
	only provider we've found competent and available to treat our child's	
	PANS/PANDAS.	
	3. Physicians in Oregon will decline to offer treatments that invite scrutiny.	
	Eventually, relevant Oregon clinicians will adopt national PANS/PANDAS diagnostic	
	and treatment protocols. Imposing extraordinary practice oversight, of the kind	
	routine for medical education or licensing discipline, will disincentivize clinicians	
	willing to fill this crucial role. Doctors already contend with insurers' medical	
	necessity and pre-authorization burdens. There's no reason these customary	
	safeguards cannot serve this committee's well-intentioned goals. The proposed	
	rule of two specialists for IVIG treatment coverage adds unnecessary impediments	





ID/#	Comment	Disposition
T1	to delivery of care. In violation of the physician's oath, this will directly harm critically ill children by preventing them from timely accessing healing therapies. Your committee reviewed that it is not in the business of creating a center for care. Neither should it be in the business of creating roadblocks to care. I am writing to express concern about the proposed treatment limitations on coverage of PANDAS and PANS that: (1) Limit coverage to "Up to 3 monthlycourses of(IVIG) therapy" (2) Require "fail first" protocols ("Two or more less-intensive therapieshave been tried and were not effective") (3) Require "A consultation with and recommendation by 2 pediatric subspecialists" PANDAS and PANS are neuropsychiatric disorders that fall within the scope of the Paul Wellstone and Pete Domenici Mental Health Parity and Addiction Equity Act of 2008 (MHPAEA). The Oregon Department of Justice has affirmed, in a publicly released legal opinion¹, that MHPAEA applies to Coordinated Care Organizations. The Center for Medicare and Medicaid Services (CMS) has released a "Parity Compliance Toolkit" with detailed technical information and guidance to help states assess compliance with the final Medicaid/CHIP parity rule. As specified in the final parity rule, and the CMS toolkit, Medicaid programs must not impose either quantitative or non-quantitative treatment limitations (NQTLs) on mental health benefits that are not comparable to, or applied more stringently that, those used with respect to the medical / surgical benefits in the same classification.	Thank you for your comment. The Commission recognizes the importance of complying with these and other laws, and develops coverage criteria that can be implemented in a compliant fashion. Our understanding is that PANDAS and PANS are not mental health conditions as defined by state and federal law because they do not appear under any of the diagnostic categories listed in the mental disorders section of the current edition of the International Classification of Disease, and do not appear in the Diagnostic and Statistical Manual of Mental Disorders.
	The limit on coverage of "Up to 3 monthly courses of (IVIG) therapy" is a quantitative limitation for the purposes of MHPAEA • CMS guidance on MHPAEA specifically identifies "fail first" protocols, such as requiring "Two or more less-intensive therapies," as NQTLs	





ID/#	Comment	Disposition
	CMS guidance also identifies preauthorization requirements, such as the	
	requirement for "A consultation with and recommendation by 2 pediatric	
	subspecialists," as NQTLs	
	The proposed PANDAS / PANS benefits are classified as "outpatient" benefits under	
	the final parity rule. This means that the Oregon Health Plan cannot impose these	
	kinds of quantitative and non-quantitative treatment limitations on treatment for	
	PANDAS / PANS unless it imposes comparable limits on substantially all medical /	
	surgical outpatient benefits.	
	A review of the prioritized list indicates that these proposed limitations for PANDAS	
	/ PANS are unique:	
	The predominant medical / surgical outpatient benefits on the list are NOT	
	limited to "3 monthly courses of therapy"	
	• The predominant medical / surgical outpatient benefits on the list do NOT have	
	stringent fail-first requirements	
	• The predominant medical / surgical outpatient benefits on the list do NOT require	
	consultation and referrals from multiple specialists	
	Therefore, the proposed limitations on PANDAS / PANS are unlawful. A CCO that	
	attempted to implement those limitations could be subject to legal action under	
	MHPAEA.	
	I urge you instead to provide coverage of the proposed treatment for PANDAS /	
	PANS without these onerous and unlawful limitations.	
	Note also that the Early and Periodic Screening, Diagnostic and Treatment (EPSDT)	
	requires states "to provide any additional health care services" (for children under	
	age 21) "that are coverable under the Federal Medicaid program and found to be	
	medically necessary to treat, correct or reduce illnesses and conditions discovered	
	regardless of whether the service is covered in a state's Medicaid plan."	





ID/#	Comment	Disposition
	Health services for PANDAS / PANS must be covered for children under age 21 as	
	an EPSDT benefit.	
U1	The PACE Foundation, established in 2016, is dedicated to improving the diagnosis, treatment and quality of life for persons with Pediatric Autoimmune Neurological Disorders through advocacy, treatment and research. In the past few years, with the assistance from the NIMH, PACE has established or partnered with a number of leading medical institutions, to create a national standard of care for pediatric Postinfectious Autoimmune Encephalopathy disorders like PANS and PANDAS (see attached clinical map). PACE representatives have attended and presented both written and verbal testimony at each of the prior HERC hearings on PANS/PANDAS Following the in-depth discussion at the 9/9 HERC session, regarding item 2b of the current HERC PANS/PANDAS guidance, PACE would recommend that the committee change item 2b as follows: 1. Require a consultation and recommendation by only 1 pediatric subspecialist 2. Supply a specific list of sub-specialists (for example, neurologist, neurodevelopmental specialist, immunologist, infectious disease, rheumatologist, etc.) 3. Provide a specified timeframe for sub-specialists to respond to a request for consultation before item 2b is waived PACE is aware that the Oregon organization named NWPPN, has drafted a substitute amendment for the current item 2b. PACE fully supports the NWPPN proposed amendment. Our organization strongly believes that if the HERC Committee does not adopt the	Thank you for your comments. See response to P1 regarding the proposed amendment to item 2b. See response to A1 regarding the 2-subspecialist requirement. Finally, it is outside of the scope of this coverage guidance review process to specify or include a pool of eligible providers who treat a specific condition in its report. We have listed examples of the types of providers that can provide consultations for PANDAS/PANS in our draft Coverage Guidance recommendation.
	changes to item 2b noted above, Oregon may become the most restrictive	





ID/#	Comment	Disposition
	PANS/PANDAS treatment state in the country. Conversely, if the HERC Committee	
	does adopt the suggested changes to item 2b, it will mirror the requirement for the	
	majority of states in the US.	





References Provided by Commenters

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C–U	None provided	





Pediatric Autoimmune Neuropsychiatric Disorders Associated with Streptococcal Infections (PANDAS) and Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS)

Draft Coverage Guidance for HERC Consideration May 19, 2022





Disclosures

- None of the authors have any conflicts of interest to disclose
- This slide set is designed for a live presentation with commentary, to accompany the full report. Each slide will contain the accompanying page number(s) from the draft coverage guidance (CG)
- The full draft CG can be found elsewhere in these meeting materials.
- Citation of studies corresponds with the reference number assigned in the CG (e.g., Sigra et al., 2018 = CG ref #24)





Appointed Ad Hoc Experts

Alison Christy, MD, PhD

- Specialty: pediatric neuroimmunology, neuroimmunological disorders, pediatric neurology, and movement disorders
- Conflicts to disclose: None

Michael Daines, MD

- Specialty: pediatric allergy and immunology, rheumatology
- Conflicts to disclose: Lead investigator of Phase 3 clinical trial of IVIG for PANS; industry funding for design of clinical trial (Octapharma); travel reimbursements from PACE Foundation, a PANDAS/PANS advocacy group

Paria Zarrinnegar, MD

- Specialty: psychiatry, biopsychosocial assessment of children and adolescents
- Conflicts to disclose: None





Scope Adjustments

- HERC approved a scope statement for the following interventions:
 - IVIG, plasma exchange, tonsillectomy, adenoidectomy, corticosteroids, SSRIs, short-course antibiotics, prophylactic antibiotic therapy, behavioral therapies, NSAIDs
- However, EbGS excluded interventions not subject to utilization controls, such as corticosteroids, NSAIDs, SSRIs, behavioral therapies, prophylactic antibiotics
- Pediatric autoimmune encephalitis was removed as a Condition from the scope due to different diagnostic criteria and higher acuity of the condition





Background



- Sudden onset of changes or regressions in behaviors in multiple domains, typically including symptoms of OCD and verbal or motor tics
- Symptom severity can range from mild to severe and can significantly interfere with daily activities (e.g., school)
- PANDAS and PANS are diagnoses of exclusion and include ruling out:
 - Primary psychiatric diagnoses, such as OCD
 - Sydenham chorea, pediatric autoimmune encephalitis, neuropsychiatric lupus, central nervous system vasculitis
 - Other conditions that better account for the symptoms





Background



- Prevalence of PANDAS and PANS is unknown
- Natural histories of PANDAS and PANS still being studied
 - Emerging research suggests that 60% to 80% of pediatric patients have significant reduction in symptoms over time
 - American Academy of Child and Adolescent Psychiatry practice parameter suggests small subset of children diagnosed with OCD or Tourette disorder may experience exacerbations related to streptococcal infection





Scope Statement



Populations

Children diagnosed with PANDAS or PANS

Interventions

 Therapeutic plasma exchange; IVIG; antibiotics; tonsillectomy and/or adenoidectomy

Comparators

Usual care or other interventions





Scope Statement



Critical Outcomes

- Change in psychiatric symptom scores
- Hospitalizations, including institutionalization or emergency visits

Important Outcomes

- Harms
- Standardized measures of function or quality of life for patients and caregivers





Scope Statement



Key Questions

- 1. Effectiveness of treatments
- 2. Comparative effectiveness for populations based on:
 - a. Patient characteristics
 - b. Condition characteristics
 - c. Intervention
 - d. Provider characteristics (e.g., Center of Excellence)
- 3. Harms of treatments





Evidence Review





Evidence Sources



- Most recent systematic review: Sigra et al., 2018 (CG ref #24)
- 5 RCTs (CG refs #25, 26, 30-32)
- 2 comparative cohort studies (CG ref #28-29)

Populations

- PANDAS
- PANS
- CANS
- PITAND
- Abrupt-onset OCD

Interventions

- Antibiotics
- Tonsillectomy
- Therapeutic plasma exchange
- IVIG
- Tonsillectomy
- Adenoidectomy
- Adenotonsillectomies





Evidence Section Overview

- Overall, small sample sizes and short follow-ups in comparative studies
- Reported outcomes
 - No evidence identified for hospitalizations and function or quality of life for patient or caregivers
 - Very little data on comparative effectiveness by subpopulations
 - Change in psychiatric symptoms and harms summarized by intervention type on following slides





GRADE Table: Prophylactic Antibiotics

Outcomes	Estimate of Effect for Outcome Confidence in Estimate
Change in psychiatric symptom scores (Critical outcome) Hospitalizations (Critical outcome)	Mixed results for antibiotics (i.e., penicillin, azithromycin) administered for 4 weeks to 1 year, compared to placebo or other antibiotic. ● ○ (very low confidence, based on 3 RCTs, n = 91) No evidence identified.
outcome) Harms (Important outcome)	The few harms that were reported included heart rate irregularity (9/12) for children who received azithromycin, and loose stool (no statistics reported). ● ○ (very low confidence, based on 1 RCTs, n = 23)
Function or quality of life for patient (Important outcome)	No evidence identified.
Function or quality of life for patient (Important outcome)	No evidence identified.



GRADE Table: Tonsillectomy and Adenoidectomy

Outcomes	Estimate of Effect for Outcome Confidence in Estimate
Change in psychiatric symptom scores (Critical outcome)	No difference in neuropsychiatric symptoms between surgery and non-surgery groups among children diagnosed with PANDAS.
	•• (low confidence, based on 2 comparative cohort studies, $n = 232$)
Hospitalizations (Critical outcome)	No evidence identified.
Harms (Important outcome)	No evidence identified.
Function or quality of life for patient (Important outcome)	No evidence identified.
Function or quality of life for patient (Important outcome)	No evidence identified.





GRADE Table: IVIG

Outcomes	Estimate of Effect for Outcome/ Confidence in Estimate
Change in psychiatric symptom scores (Critical outcome) Hospitalizations (Critical outcome)	Compared to saline placebo, some children had decreased symptoms. Compared to plasma exchange, no significant difference. ● ○ (very low confidence, based on 2 RCTs, n = 54) No evidence identified.
Harms (Important outcome)	1/33 had an allergic reaction to the IVIG infusion that resolved without complication. 31/33 children reported mild or moderate adverse events (e.g., nausea, vomiting, headache, fever, joint pain). ● ○ (very low confidence, based on 2 RCTs, n = 64)
Function or quality of life for patient (Important outcome)	No evidence identified.
Function or quality of life for patient (Important outcome)	No evidence identified.

GRADE Table: Plasma Exchange

Outcomes	Estimate of Effect for Outcome Confidence in Estimate
Change in psychiatric	Compared to saline placebo, improvements on most measures.
symptom scores (Critical	Compared to IVIG, no significant difference.
outcome)	● ○ (very low confidence, based on 1 RCT, n = 29)
Hospitalizations (Critical outcome)	No evidence found.
Harms (Important	All children who received plasma exchange (10/10) experienced mild
outcome)	side effects such as nausea, vomiting, anxiety, or fever.
	● ○ (very low confidence, based on 1 RCT, n = 29)
Function or quality of life	No evidence found.
for patient (Important	
outcome)	
Function or quality of life	No evidence found.
for patient (Important	
outcome)	





Ongoing Studies



- Double-blind RCT (N = 44) to test naproxen sodium for 8 weeks in children with PANDAS
 - Estimated completion: October 2022
- Quadruple-blind RCT (N = 92) to test IVIG in children with PANDAS or PANS
 - Estimated completion: March 2023
- Observational matched cohort study (N = 500) to examine immunologic, neurologic, genomic, and behavioral differences between children with PANS and healthy children for up to 15 years
 - Estimated completion: March 2028





Evidence Summary



- The natural histories of PANDAS and PANS are still being studied
- The very low and low confidence we have in the findings means that findings from new comparative studies that test treatments for PANDAS or PANS could easily change the evidence underpinning decisions about which treatments should be covered for children with PANDAS or PANS





Clinical Practice Guidelines





PANS/PANDAS Clinical Research Consortium



- Proposed a 3-pronged approach to treatment (CG ref #3, 19, 20, 22):
 - Treat psychiatric symptoms with psychoactive medications and psychotherapy;
 - Remove source of inflammation with antimicrobial interventions; and
 - Treat immune system disturbances with immunomodulatory and/or anti-inflammatory therapies
- Not all involved experts agreed on treatments proposed in publication (CG ref #3)





Nordic Countries Guideline



- Treatment should be overseen at specialized centers
- Verified or strongly suspected infections should be treated with antibiotics for no longer than 14 days
 - Do not recommend prophylactic antibiotic therapy
- Severe symptoms may be treated first with NSAIDs, then possibly escalate to IVIG
- Authors state plasma exchange and immunomodulatory drugs are only clinically indicated for children diagnosed with autoimmune encephalitis (CG ref #21)





Policy Landscape





Payer Policies



No policies identified for Washington State Medicaid,
 Medicare, Moda, or BlueCross BlueShield

Aetna

- The following are considered investigational or experimental for PANDAS:
 - Parenteral immunoglobulins
 - Plasma exchange
 - Rituximab

Cigna

- The following are considered investigational or experimental for PANDAS and PANS:
 - IVIG
 - Plasma exchange
 - Rituximab





Other Recommendations



- PANDAS Physician Network (CG ref #56)
 - Recommends that children with moderate to severe symptoms be treated by experienced multidisciplinary team or PANDAS/PANS specialist
 - Summary of proposed treatment sequence (see <u>website</u> for details):
 - Start with 14 days of antibiotic therapy, lengthen therapy if infection is not resolved or symptoms persist. Consider prophylactic antibiotic therapy
 - 2. Consider 5 to 7 day course of NSAIDs
 - 3. Ensure access to CBT and parenting management techniques
 - 4. Consider steroid course
 - Escalate to IVIG
 - Administer second course of IVIG or plasma exchange plus prophylactic antibiotics





Discussion





Discussion: General

Resource Allocation

Some of the proposed treatments (antibiotics, NSAIDs, SSRIs) are inexpensive and widely available. Other proposed treatments are more expensive and may have limited access (IVIG, therapeutic plasmapheresis, behavioral therapy).

Untreated PANDAS/PANS has high costs with frequent medical visits, hospitalizations, school disruption, parental inability to work, and other factors.

Values and Preferences

Some parents would value any treatment that might help their child's symptoms.

Other parents would have concerns about the risks and side effects of the therapies being considered, and/or the unproven efficacy of the intervention.





Discussion: Prophylactic Antibiotic Therapy

Other Considerations

Long-term or frequent antibiotic use is associated with a range of negative consequences, including but not limited to C. difficile infection, diarrhea, and increased antibiotic resistance leading to reduced ability to treat other infections with antibiotics. Most health plans cover short-term antibiotics without prior authorization criteria but may scrutinize or not cover long-term prescriptions.

Balance of Benefits and Harms

We have very low confidence that prophylactic antibiotic use is helpful in PANDAS/PANS, given the small sample sizes in the included studies. There are concerning known harms of frequent or long-term antibiotic use.

Rationale

Prophylactic antibiotic therapy is not recommended for coverage for PANDAS/PANS because of insufficient comparative evidence that prophylactic antibiotic use leads to any measurable benefit for these conditions. The known risks of prophylactic or long-term antibiotic use outweigh potential benefits in these conditions.





Discussion: Tonsillectomy and Adenoidectomy

Other Considerations

Tonsillectomy and/or adenoidectomy have known harms such as bleeding, anesthesia reactions, and death. Tonsillectomy and/or adenoidectomy frequently have coverage limitations, such as multiple streptococcal infections in one year. Historically, this procedure has been overused.

Balance of Benefits and Harms

We have low confidence that that there is no benefit from tonsillectomy and/or adenoidectomy for PANDAS, and this procedure has known harms. This treatment has not been proposed for PANS.

Rationale

Tonsillectomy and/or adenoidectomy are not recommended for coverage for treatment of PANDAS/PANS because these procedures have known harms and because evidence shows that these procedures do not improve the outcomes in this condition.





Discussion: IVIG

Other Considerations

IVIG is a blood product with the inherent risks that accompany accepting any form of blood product. IVIG has a significant rate of mild side effects including fever, body aches, nausea, rash, and fatigue. Severe side effects include thrombosis, renal dysfunction, and acute renal failure, and life-threatening allergic reaction. IVIG can interfere with vaccine effectiveness for vaccines given within several months of IVIG. Several products on the market are FDA-approved for people under the age of 19.

Balance of Benefits and Harms

There were mixed results from 2 very small trials regarding the clinical effectiveness of IVIG. Outside of PANDAS, no evidence met inclusion criteria for PANS. IVIG has a significant rate of known harms.

Rationale

Expert opinion and lower-quality observational data indicate there may be benefit for some patients with these conditions, but recommended treatment protocols and criteria for treatment vary widely. Due to the severe impact of symptoms associated with PANDAS/PANS on child health, growth and development, and the lack of known effective treatments, coverage of IVIG is recommended when recommended by the patient's PCP and a pediatric subspecialist and after less-intensive therapies were not effective, were not tolerated or did not result in sustained improvement in symptoms.





Discussion: Plasma Exchange

Other Considerations

High rates of patients undergoing plasma exchange report side effects, including fever, chills, and muscle cramps. Known complications of plasma exchange include circuit clotting, low or high blood pressure, nausea, vomiting, itchy skin, hives, low calcium levels in the blood, venous access malfunction, infections, thrombosis, and anaphylactic shock.

Balance of Benefits and Harms

The comparative evidence that plasma exchange is effective at treating PANDAS is very limited and shows no clear benefit. The harms of this treatment are generally mild but serious complications can occur.

<u>Rationale</u>

Expert opinion and lower-quality observational data indicate there may be benefit for some patients with these conditions, but recommended treatment protocols and criteria for treatment vary widely. Due to the severe impact of symptoms associated with PANDAS/PANS on child health, growth and development, and the lack of known effective treatments, coverage of plasma exchange is recommended when recommended by the patient's PCP and a pediatric subspecialist and after less-intensive therapies were not effective, were not tolerated or did not result in sustained improvement in symptoms.





Recommendation

Tonsillectomy, adenoidectomy, adenotonsillectomy, and prophylactic antibiotic therapy are not recommended for coverage to treat pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) and pediatric acute-onset neuropsychiatric syndrome (PANS) (weak recommendation).

Up to 3 monthly immunomodulatory courses of intravenous immunoglobulin (IVIG) therapy or therapeutic plasma exchange are recommended for coverage to treat PANDAS and PANS (weak recommendation) when both of the following are met:

- A) A clinically appropriate trial of two or more less-intensive treatments (for example, appropriate limited course of nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, selective serotonin reuptake inhibitors (SSRIs), behavioral therapy, short-course antibiotic therapy) was either not effective, not tolerated, or did not result in sustained improvement in symptoms (as measured by a lack of clinically meaningful improvement on a validated instrument directed at the patient's primary symptom complex). These trials may be done concurrently, AND
- B) A consultation with and recommendation from a pediatric subspecialist (for example, pediatric neurologist, pediatric psychiatrist, neurodevelopmental pediatrician, pediatric rheumatologist, pediatric allergist/immunologist, pediatric infectious disease specialist) as well as the recommendation of the patient's primary care provider (for example, family physician, pediatrician, pediatric nurse practitioner, naturopath). The subspecialist consultation may be a teleconsultation. For adolescents, an adult subspecialist consult may replace a pediatric subspecialist consult.

A reevaluation at 3 months by both the primary care provider and pediatric expert is required for continued therapy of IVIG or plasma exchange. This evaluation must include clinical testing with a validated instrument, which must be performed pretreatment and posttreatment to demonstrate clinically meaningful improvement.





Supplemental Materials





Abbreviations

Conditions

- CANS: childhood acute neuropsychiatric syndromes
- OCD: obsessive-compulsive disorder
- PANDAS: pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections
- PANS: pediatric acute-onset neuropsychiatric syndrome
- PITAND: pediatric infection-triggered autoimmune neuropsychiatric disorders

Treatment

IVIG: intravenous immunoglobulin

Other abbreviations

RCT: randomized controlled trial



