

## EXPERIMENTAL/INVESTIGATIONAL/UNPROVEN CARE/ BENEFIT EXCEPTIONS

Effective Date: July 1, 2024 Review Dates: 1/93, 4/95, 10/97, 4/99, 2/01, 12/01,

11/02, 11/03, 11/04, 10/05, 10/06, 10/07, 10/08, 4/09, 4/10, 4/11, 4/12, 4/13, 12/13, 11/14, 11/15, 11/16, 11/17, 11/18, 11/19, 11/20, 11/21, 5/22, 5/23, 5/24,

5/25

Date of Origin: June 30, 1988 Status: Current

#### Related medical policies:

• 91636 - Category III Current Procedural Terminology (CPT®) Codes

• 91448 - Clinical Trials for Self-Funded Groups Opting Out of PPACA

• 91606 - Clinical Trials

#### I. POLICY/CRITERIA

- A. Any drug, device, treatment, or procedure that is experimental, investigational or unproven is not a covered benefit. A drug, device, treatment, or procedure is experimental, investigational or unproven if *any* of the following apply:
  - 1. The drug or device final marketing approval or clearance has not been granted by the Food and Drug Administration (FDA); or
  - 2. The drug, device, treatment, or procedure is provided pursuant to oversight by an institutional review board (IRB) or other body that approves or reviews research concerning safety, toxicity or efficacy; or
  - 3. The patient informed consent documents describe the drug, device, treatment, or procedure as experimental or investigational or in other terms that indicate the service is being evaluated for its safety, toxicity, or efficacy; or
  - 4. Reliable evidence shows that the drug, device, treatment, or procedure is the subject of on-going Phase I or Phase II clinical trials or is the research, experimental, study or investigational arm of on-going Phase III clinical trials; or is otherwise under study to determine its toxicity, safety, or efficacy as compared with a standard means of treatment or diagnosis; or
  - 5. Reliable evidence shows that the prevailing opinion among experts regarding the drug, device, treatment, or procedure is that further studies or clinical trials are necessary to determine its toxicity, safety, or efficacy as compared with a standard means of treatment or diagnosis.
  - 6. The drug, device, treatment, or procedure is not widely used or generally accepted as standard medical care for the condition, disease, illness or injury being treated as reported in nationally recognized peer-reviewed medical literature published in the English language.



- B. Category III codes: Unless there is a Priority Health Medical Policy that specifically addresses coverage or medical necessity for a particular code the item, service or procedure represented by any Category III code is considered experimental, investigational, or unproven. See medical policy Category III Current Procedural Terminology (CPT®) Codes #91636.
- C. Individual case review may allow coverage for care or treatment that is investigational yet promising for the conditions described. Requests for individual consideration require prior Plan approval. All determinations of coverage for experimental, investigational, or unproven treatment will be made by a Priority Health medical director or clinical pharmacist. The exclusion of coverage for experimental, investigational, or unproven treatment may be **reviewed for exception** if the condition is:
  - a. A terminal illness, or
  - b. A chronic, life threatening, severely disabling disease that is causing serious clinical deterioration.
  - 1. All accepted standard treatments and technologies must be considered or used prior to review for exception under this policy.
  - 2. Any treatment or evaluation (including additional opinions) authorized under this policy must be received at a participating facility or a facility within the Plan's network.
  - 3. Any treatment authorized must be under the auspices of a nationally recognized sponsor such as the National Institutes of Health (NIH) and adhere to the US regulation standards of being approved and monitored by an Institutional Review Board (IRB) to make sure the risks are as low as possible and are worth any potential benefits.
  - 4. When care is available both within a clinical trial and outside a clinical trial, coverage preference will be given to the clinical trial. When care is available within multiple trials, coverage will be given to the more definitive trial (e.g., Phase III over Phase II).
  - 5. Informed consent must be documented.
  - 6. An independent expert physician review panel may be consulted to determine the appropriateness of the recommended treatment. The panel members will each provide their opinion on whether the treatment is promising and likely to be effective for that individual patient.
  - 7. Costs associated with experimental care: Funding for experimental care, which covers the cost of protocol development and data collection traditionally comes from a variety of sources including pharmaceutical companies, research institutions and government agencies (referred to as "sponsors"). The following is intended to clarify what the plan will cover and what the sponsoring facility is expected to cover.
    - a. The administrative costs are borne by the facility or sponsor, including:
      - 1. Data gathering
      - 2. Statistical study
      - 3. Regulatory requirements



- 4. Contractual agreements
- 5. Meetings and travel
- b. The routine patient care costs (conventional care) are covered by Priority Health.
  - 1. Routine patient care costs are items or services that are typically covered benefits when provided outside a clinical trial or experimental care.
  - 2. Routine services include services that would be approved for coverage under this policy, even when delivered within the context of a clinical trial or experimental care.
- c. Coverage for devices classified under the FDA Investigational Device Exemption (IDE) or Humanitarian Use Device (HUD)/Humanitarian Device Exemption (HDE). See definitions in Description Section & Appendix B for product specific coverage
  - 1. IDEs
    - a. Category A IDEs and associated care and services are not covered benefits
    - b. Category B IDEs when used in a clinical trial and prior authorized by Priority Health:
      - 1. Routine patient care costs in a clinical trial are covered as defined above.
      - 2. The device is not a covered benefit
  - 2. HUD/HDEs. Devices that have FDA approval for humanitarian use or as HDEs are considered experimental and investigational and excluded from coverage unless they are listed as covered in Appendix C.
- d. The costs associated in the delivery of the investigational agent are covered by Priority Health.
  - 1. Services required solely for the provision of the investigational item shall be provided in accordance with the benefits of the patient's health plan. Coverage would include procedures, drugs or devices approved for coverage for any medical indication.
  - 2. The clinically appropriate monitoring of the effects of the item or service should be considered routine patient care costs.
  - 3. The prevention of complications of the item or service should be considered routine patient care costs.
  - 4. This coverage shall include payment for reasonable and medically necessary services to administer the drug or use the device under evaluation in the clinical trial.
- e. Costs incurred for patient care generated specifically by the clinical trial or experimental care shall be borne by the facility or sponsor.
  - 1. The cost of the investigational drug, device, or service itself.



#### Experimental/Investigational/ Unproven Care/Benefit Exceptions

- 2. Costs incurred for patient care generated specifically by the clinical trial. Examples of these are costs for additional medication, laboratory studies, or diagnostic imaging.
- 3. The health plan's coverage of "routine costs" would *not* include non-FDA approved drugs or devices or unapproved medical procedures.
- 4. Coverage would *not* include diagnostic tests that are performed for investigational purposes but not necessary for the member's medical management.
- 5. It would also *not* include services beyond the scope of the member's contract.
- f. Costs of treating adverse side effects experienced during treatment are covered by Priority Health. Priority Health will cover medical care needed to treat any complications arising from the experimental and investigational service when the medical services provided are otherwise covered under the member's contract.
- g. Care outside the United States is not covered.

Coverage for care and services received in a clinical trial is defined in the Clinical Trials Medical Policy #91606. Refer to the "Clinical Trials" policy for benefits and limitations.

Member must have an advance care planning assessment (see Appendix A at the end of this medical policy) completed by a qualified provider. The assessment should accompany the request for a benefit exception.

#### III. MEDICAL NECESSITY REVIEW

□ Required	☐ Not Required	☐ Not Applicable
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#### IV. APPLICATION TO PRODUCTS

Coverage is subject to member's specific benefits. Group specific policy will supersede this policy when applicable.

- **❖** HMO/EPO: *This policy applies to insured HMO/EPO plans.*
- **POS:** This policy applies to insured POS plans.
- \* PPO: This policy applies to insured PPO plans. Consult individual plan documents as state mandated benefits may apply. If there is a conflict between this policy and a plan document, the provisions of the plan document will govern.
- ASO: For self-funded plans, consult individual plan documents. If there is a conflict between this policy and a self-funded plan document, the provisions of the plan document will govern.
- \* INDIVIDUAL: For individual policies, consult the individual insurance policy. If there is a conflict between this medical policy and the individual insurance policy document, the provisions of the individual insurance policy will govern.



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- MEDICARE: Coverage is determined by the Centers for Medicare and Medicaid Services (CMS) and/or the Evidence of Coverage (EOC); if a coverage determination has not been adopted by CMS, this policy applies.
- \* MEDICAID/HEALTHY MICHIGAN PLAN: For Medicaid/Healthy Michigan Plan members, this policy will apply. Coverage is based on medical necessity criteria being met and the appropriate code(s) from the coding section of this policy being included on the Michigan Medicaid Fee Schedule located at: <a href="http://www.michigan.gov/mdch/0,1607,7-132-2945">http://www.michigan.gov/mdch/0,1607,7-132-2945</a> 42542 42543 42546 42551-159815--,00.html. If there is a discrepancy between this policy and the Michigan Medicaid Provider Manual located at: <a href="http://www.michigan.gov/mdch/0,1607,7-132-2945">http://www.michigan.gov/mdch/0,1607,7-132-2945</a> 5100-87572--,00.html, the Michigan Medicaid Provider Manual will govern. If there is a discrepancy or lack of guidance in the Michigan Medicaid Provider Manual, the Priority Health contract with Michigan Medicaid will govern. For Medical Supplies/DME/Prosthetics and Orthotics, please refer to the Michigan Medicaid Fee Schedule to verify coverage.

#### V. DESCRIPTION

Experimental and investigational (with respect to medical research), refers to a procedure, device or pharmaceutical agent that is still undergoing pre-clinical or clinical evaluation, and/or has not yet received regulatory approval or is not recognized as standard medical care for the condition, disease, illness or injury being treated.

Criteria used in determining whether the technologies, equipment, supplies, treatments, procedures, therapies, biologics, drugs, or devices is considered experimental or investigational include, but are not limited to:

- 1. Whether it is commonly performed or used for the disease or condition;
- 2. Whether it is generally accepted as standard treatment or diagnosis for the disease or condition by the medical professionals or medical professional societies in the United States:
- 3. Whether it is medically indicated;
- 4. Whether there is sufficient or conclusive data to assess the therapeutic value or positive effects on short and long-term health outcomes (e.g., safety and effectiveness, failure rate, and side effects)

Medical research is conducted to aid the body of knowledge in the field of medicine. This can be divided into two general categories: New treatments that are tested in clinical trials, and all other research contributing to the development of new treatments. A new treatment refers to any form of previously untested treatment for a particular pathology. This can take the form of a new surgical procedure, a new drug, or a new treatment regimen. These are extensively tested in clinical trials prior to wide-spread use. Formal clinical trials have, among other aspects, extensive written research protocols that adhere to established research principles and study design.

At the early stages, study protocols usually focus on the safety of the new drug, device, or procedure using a single group of research subjects. Such "single arm" trials generally are followed by more extensive studies that measure the experimental

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intervention against alternative therapies and/or involve a rudimentary comparison between experimental and control subject groups. When basic safety and efficacy have been demonstrated by the experimental scientific process the investigational phase begins. As the research further matures, the new intervention will be tested in double-blind randomized studies, the so-called "gold-standard" of research. Depending on study results, the intervention may become a generally recognized standard of care.

The FDA defines Humanitarian Use Device (HUD) as a medical device intended to benefit patients in the treatment or diagnosis of a disease or condition that affects or is manifested in not more than 8,000 individuals in the United States per year. Humanitarian Device Exemption (HDE) is a marketing application for an HUD (Section 520(m) of the Federal Food, Drug, and Cosmetic Act (FD&C Act)). An HDE is exempt from the effectiveness requirements of Sections 514 and 515 of the FD&C Act and is subject to certain profit and use restrictions. HDE approval authorizes marketing of an HUD device for its specified indication for use. HDE approval is based upon, among other criteria, a determination by the FDA that the HUD will not expose patients to an unreasonable or significant risk of illness or injury and the probable benefit to health from use of the device outweighs the risk of injury or illness from its use (while taking into account the probable risks and benefits of currently available devices or alternative forms of treatment). The law exempts HDE devices from demonstrating a reasonable assurance of effectiveness, and instead requires demonstration of probable benefit. This difference in determination of effectiveness is a key difference between applications for premarket approval (PMA) and HDE devices. The table below compares some key aspects of HDEs and PMAs.

#### **Definitions:**

<u>Clinical Trials</u> (from the National Cancer Institute)

Clinical trials in cancer therapy are conducted to decrease morbidity and mortality from cancer. New drug development is one part of this effort, but other parts include the integration of multiple treatment modalities, the testing of new combinations of existing drugs, the testing of new dose schedules and routes of administration, the application of new diagnostic tests in choosing treatment regimens, and the evaluation of supportive care methods.

*Phase I* — The initial clinical test of a new treatment modality. Most Phase I patients have cancer for which no other effective therapeutic options are known, and patients with any type of cancer are admitted to most Phase I trials.

*Phase II* — The initial efficacy trial of a new cancer agent. The trial is done on groups of patients with one type of cancer.

Phase III — Designed to compare one or more treatments. A new drug or drug combination ("research arm") may be tested against a drug combination of proven efficacy. The patients are randomly allocated to the treatment options.



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Clinical Trials for Investigational New Drugs (from the Food & Drug Administration)

*Phase I* — Testing concerned primarily with the safety of the drug and normally done on a small number (20-100) of healthy volunteers.

*Phase II* — This phase of drug testing involves a few hundred patients and is designed to show whether the drug is effective in treating the disease or condition for which it is intended. Most Phase II studies are randomized controlled trials.

*Phase III* — The population size is expanded to several hundred to several thousand to clarify the drug's benefit-risk relationship and discover side effects and adverse reactions.

These three phases are necessary for FDA marketing approval of a new drug. Post marketing surveillance (*Phase IV*) is done to detect adverse reactions that might not have been detected in earlier trials.

Current Procedural Terminology (CPT®) Category III codes: Developed by the American Medical Association (AMA) and defined as a set of temporary ("T") codes that allow data collection for emerging technologies, services, procedures, and service paradigms. These codes are intended to be used for data collection to substantiate widespread usage or to provide documentation for the Food and Drug Administration (FDA) approval process. Unlike Category I CPT® codes, the procedures and services described by Category III CPT® codes do not necessitate FDA approval and therefore have been placed in a separate section of the CPT book. Per the AMA, "the inclusion of a service or procedure in this section does not constitute a finding of support, or lack thereof, with regard to clinical efficacy, safety, applicability to clinical practice, or payer coverage." The Category III CPT® Code description does not establish a service or procedure as safe, effective or applicable to the clinical practice of medicine.

#### Investigational Device Studies (IDEs)

Category A (Experimental) device refers to a device for which "absolute risk" of the device type has not been established (that is, initial questions of safety and effectiveness have not been resolved) and the FDA is unsure whether the device type can be safe and effective.

Category B (Non-experimental/investigational) device refers to a device for which the incremental risk is the primary risk in question (that is, initial questions of safety and effectiveness of that device type have been resolved), or it is known that the device type can be safe and effective because, for example, other manufacturers have obtained FDA premarket approval or clearance for that device type.

<u>Peer-reviewed literature</u> Articles or reports that have gone through an evaluation process in which journal editors and other expert scholars critically assess the quality and scientific merit of the article and its research. Articles that pass this process are published in the peer-

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reviewed literature. Peer-reviewed journals may include the research of scholars who have collected their own data using an experimental study design, survey, or various other study methodologies. They also present the work of researchers who have performed novel analyses of existing data sources, such as the ones described in this section.

Peer-reviewed literature is accessible via academic databases that enable users to execute searches across multiple journals.

<u>Promising</u> — Preliminary scientific data supports reasonable likelihood of success of the treatment for the diagnosis.

Reliable evidence means published reports and articles in the authoritative medical and scientific literature; the written protocol or protocols used by the treating facility or the protocol(s) of another facility studying substantially the same drug, device, treatment or procedure.

<u>Terminal illness</u> — A disease that can be expected to result in death within 1 year in the absence of effective treatment.

#### VI. REFERENCES

- 1. Centers for Medicare and Medicaid Services. <u>National Coverage Determination (NCD)</u> 310.1 Routine Costs in Clinical Trials. (Accessed March 12, 2025).
- Centers for Medicare and Medicaid Services. <u>Medicare Coverage of Items and Services in Category A and B Investigational Device Exemption (IDE) Studies</u>, CMS MLN Matters, MM8921, effective January 1, 2015.
- 3. Centers for Medicare and Medicaid Services, Internet Only Manual (IOM), Publication 100-08, Medicare Program Integrity Manual, Chapter 13.5.4, Local Coverage.

  Reasonable and Necessary Provision in an LCD.
- 4. National Library of Medicine. Health Data Sources. Peer-reviewed Literature. at
- 5. U.S. Food and Drug Administration. <u>FDA Categorization of Investigational Device</u> Exemption (IDE) Devices to Assist the Centers for Medicare and Medicaid Services (CMS) with Coverage Decisions.

#### VII. CODING

**See Policies:** 

91448 Clinical Trials for Self Funded Groups Opting Out of PPACA

91606 Clinical Trials

91636 Category III Current Procedural Terminology (CPT®) Codes



## Experimental/Investigational/ Unproven Care/Benefit Exceptions

**GENERAL NOT COVERED** services based on Experimental, Investigational, Unproven Care and plan document language. *This List is not inclusive. These codes are not included in any specific medical policy*.

#### **CPT/HCPCS codes:**

СРТ/НС	T/HCPCS codes:			
20560	Needle insertion(s) without injection(s); 1 or 2 muscle(s)			
20561	Needle insertion(s) without injection(s); 3 or more muscles			
	3 (7)			
34839	Physician planning of a patient-specific fenestrated visceral aortic endograft			
3 1037	requiring a minimum of 90 minutes of physician time			
43206	Esophagoscopy, rigid or flexible; with optical endomicroscopy			
43252	Upper gastrointestinal endoscopy including esophagus, stomach, and either the			
50004	duodenum and/or jejunum as appropriate; with optical endomicroscopy			
52284	Cystourethroscopy, with mechanical urethral dilation and urethral therapeutic drug			
	delivery by drug-coated balloon catheter for urethral stricture or stenosis, male,			
	including fluoroscopy, when performed			
53451	Periurethral transperineal adjustable balloon continence device; bilateral insertion,			
	including cystourethroscopy and imaging guidance			
53452	Periurethral transperineal adjustable balloon continence device; unilateral insertion,			
	including cystourethroscopy and imaging guidance			
53453	Periurethral transperineal adjustable balloon continence device; removal, each			
	balloon			
53454	Periurethral transperineal adjustable balloon continence device; percutaneous			
	adjustment of balloon(s) fluid volume			
	adjustificity of our con(s) rule volume			
55400	Vasovasostomy, vasovasorrhaphy			
58750	Tubotubal anastomosis			
69090	Ear piercing			
09090	Lai picienig			
82075	Alcohol (ethanol), breath			
83006	Growth stimulation expressed gene 2 (ST2, Interleukin 1 receptor like-1)			
	Myeloperoxidase (MPO)			
83876				
83951	Oncoprotein; des-gamma-carboxy-prothrombin (DCP)			
84145	Procalcitonin (PCT)			
84393	Tau, phosphorylated (eg, pTau 181, pTau 217), each (Covered for Medicare &			
0.400.4	Medicaid)			
84394	Tau, total (tTau) (Covered for Medicare & Medicaid)			
84431	Thromboxane metabolite(s), including thromboxane if performed, urine			
86305	Human epididymis protein 4 (HE4)			
86352	Cellular function assay involving stimulation (e.g., mitogen or antigen) and detection			
	of biomarker (e.g., ATP)			
87513	Infectious agent detection by nucleic acid (DNA or RNA); Helicobacter pylori (H.			
	pylori), clarithromycin resistance, amplified probe technique (Covered for Medicare			
	& Medicaid)			
88130	Sex chromatin identification; Barr bodies			
90865	Narcosynthesis for psychiatric diagnostic and therapeutic purposes (eg, sodium			
	amobarbital (Amytal) interview)			
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91117 92145	Colon motility (manometric) study, minimum 6 hours continuous recording (including provocation tests, e.g., meal, intracolonic balloon distension, pharmacologic agents, if performed), with interpretation and report Corneal hysteresis determination, by air impulse stimulation, unilateral or bilateral,
	with interpretation and report
93740 93895	Temperature gradient studies Quantitative carotid intima media thickness and carotid atheroma evaluation, bilateral
96020	Neurofunctional testing selection and administration during noninvasive imaging functional brain mapping, with test administered entirely by a physician or other qualified health care professional (i.e., psychologist), with review of test results and report
96931	Reflectance confocal microscopy (RCM) for cellular and sub-cellular imaging of skin; image acquisition and interpretation and report, first lesion
96932	Reflectance confocal microscopy (RCM) for cellular and sub-cellular imaging of skin; image acquisition only, first lesion
96933	Reflectance confocal microscopy (RCM) for cellular and sub-cellular imaging of skin; interpretation and report only, first lesion
96934	Reflectance confocal microscopy (RCM) for cellular and sub-cellular imaging of skin; image acquisition and interpretation and report, each additional lesion (List separately in addition to code for primary procedure)
96935	Reflectance confocal microscopy (RCM) for cellular and sub-cellular imaging of skin; image acquisition only, each additional lesion (List separately in addition to code for primary procedure)
96936	Reflectance confocal microscopy (RCM) for cellular and sub-cellular imaging of skin; interpretation and report only, each additional lesion (List separately in addition to code for primary procedure)
97610	Low frequency, non-contact, non-thermal ultrasound, including topical application(s), when performed, wound assessment, and instruction(s) for ongoing care, per day
99026	Hospital mandated on call service; in-hospital, each hour
99027	Hospital mandated on call service; out-of-hospital, each hour
99070	Supplies and materials (except spectacles), provided by the physician or other qualified health care professional over and above those usually included with the office visit or other services rendered (list drugs, trays, supplies, or materials provided)
99071	Educational supplies, such as books, tapes, and pamphlets, for the patient's education at cost to physician or other qualified health care professional
99072	Additional supplies, materials, and clinical staff time over and above those usually included in an office visit or other nonfacility service(s), when performed during a Public Health Emergency, as defined by law, due to respiratory-transmitted infectious disease
99075	Medical testimony
99080	Special reports such as insurance forms, more than the information conveyed in the usual medical communications or standard reporting form
99082	Unusual travel (e.g., transportation and escort of patient



- A4341 Indwelling intraurethral drainage device with valve, patient inserted, replacement only, each
   A4342 Accessories for patient inserted indwelling intraurethral drainage device with valve, replacement only, each
- A4563 Rectal control system for vaginal insertion, for long term use, includes pump and all supplies and accessories, any type each
- A6590 External urinary catheters; disposable, with wicking material, for use with suction pump, per month
- A6591 External urinary catheter; non-disposable, for use with suction pump, per month
- A9291 Prescription digital behavioral therapy, FDA-cleared, per course of treatment
- A9292 Prescription digital visual therapy, software-only, fda cleared, per course of treatment
- C1600 Catheter, transluminal intravascular lesion preparation device, bladed, sheathed (insertable) (Covered for Medicare)
- C1603 Retrieval device, insertable, laser (used to retrieve intravascular inferior vena cava filter) (Covered for Medicare)
- C1604 Graft, transmural transvenous arterial bypass (implantable), with all delivery system components (Covered for Medicare)
- C1824 Generator, cardiac contractility modulation (implantable)
- C1839 Iris prosthesis
- C1982 Catheter, pressure-generating, one-way valve, intermittently occlusive
- C8003 Implantation of medial knee extraarticular implantable shock absorber spanning the knee joint from distal femur to proximal tibia, open, includes measurements, positioning and adjustments, with imaging guidance (eg, fluoroscopy) (Covered for Medicare & Medicaid)
- C8004 Simulation angiogram with use of a pressure-generating catheter (e.g., one-way valve, intermittently occluding), inclusive of all radiological supervision and interpretation, intraprocedural roadmapping, and imaging guidance necessary to complete the angiogram, for subsequent therapeutic radioembolization of tumors (Covered for Medicare)
- C9759 Transcatheter intraoperative blood vessel microinfusion(s) (e.g., intraluminal, vascular wall and/or perivascular) therapy, any vessel, including radiological supervision and interpretation, when performed
- Non-randomized, non-blinded procedure for nyha class ii, iii, iv heart failure; transcatheter implantation of interatrial shunt or placebo control, including right and left heart catheterization, transeptal puncture, trans-esophageal echocardiography (tee)/intracardiac echocardiography (ice), and all imaging with or without guidance (e.g., ultrasound, fluoroscopy), performed in an approved investigational device exemption (ide) study (Covered with Prior Authorization for Medicare)
- C9782 Blinded procedure for New York Heart Association (NYHA) Class II or III heart failure, or Canadian Cardiovascular Society (CCS) Class III or IV chronic refractory angina; transcatheter intramyocardial transplantation of autologous bone marrow cells (e.g., mononuclear) or placebo control, autologous bone marrow harvesting and preparation for transplantation, left heart catheterization including ventriculography, all laboratory services, and all imaging with or without guidance (e.g., transthoracic echocardiography, ultrasound, fluoroscopy), performed in an approved investigational device exemption (IDE) study (Covered for Medicare)
- C9783 Blinded procedure for transcatheter implantation of coronary sinus reduction device or placebo control, including vascular access and closure, right heart catherization,



- venous and coronary sinus angiography, imaging guidance and supervision and interpretation when performed in an approved investigational device exemption (IDE) study (Covered for Medicare)
- C9796 Repair of enterocutaneous fistula small intestine or colon (excluding anorectal fistula) with plug (e.g., porcine small intestine submucosa [sis]) (Covered for Medicare)
- C9797 Vascular embolization or occlusion procedure with use of a pressure-generating catheter (e.g., one-way valve, intermittently occluding), inclusive of all radiological supervision and interpretation, intraprocedural roadmapping, and imaging guidance necessary to complete the intervention; for tumors, organ ischemia, or infarction (Covered for Medicare)
- C9808 Nerve cryoablation probe (e.g., cryoice, cryosphere, cryosphere max, cryoice cryosphere, cryoice cryo2), including probe and all disposable system components, non-opioid medical device (must be a qualifying medicare non-opioid medical device for post-surgical pain relief in accordance with section 4135 of the caa, 2023) (Covered for Medicare & Medicaid)
- E0490 Power source and control electronics unit for oral device/appliance for neuromuscular electrical stimulation of the tongue muscle, controlled by hardware remote
- E0491 Oral device/appliance for neuromuscular electrical stimulation of the tongue muscle, used in conjunction with the power source and control electronics unit, controlled by hardware remote, 90-day supply
- E0677 Nonpneumatic sequential compression garment, trunk
- E1905 Virtual reality cognitive behavioral therapy device (cbt), including preprogrammed therapy software
- G0183 Quantitative software measurements of cardiac volume, cardiac chambers volumes and left ventricular wall mass derived from ct scan(s) data of the chest/heart (with or without contrast) (Covered for Medicare)
- G0219 PET imaging whole body; melanoma for noncovered indications
- G0235 PET imaging, any site, not otherwise specified
- G0252 PET imaging, full and partial-ring PET scanners only, for initial diagnosis of breast cancer and/or surgical planning for breast cancer (e.g., initial staging of axillary lymph nodes)
- G0276 Blinded procedure for lumbar stenosis, percutaneous image-guided lumbar decompression (PILD) or placebo-control, performed in an approved coverage with evidence development (CED) clinical trial (Exception: Covered ONLY for Medicare, and ONLY when performed in a Coverage with Evidence Development (CED) clinical trial)
- G0429 Dermal filler injection(s) for the treatment of facial lipodystrophy syndrome (LDS) (e.g., as a result of highly active antiretroviral therapy) (Covered for Medicare per LCD L39051)
- G0513 Prolonged preventive service(s) (beyond the typical service time of the primary procedure), in the office or other outpatient setting requiring direct patient contact beyond the usual service; first 30 minutes (list separately in addition to code for preventive service) (payable for Medicare only)
- G0514 Prolonged preventive service(s) (beyond the typical service time of the primary procedure), in the office or other outpatient setting requiring direct patient contact beyond the usual service; each additional 30 minutes (list separately in addition to



- code G0513 for additional 30 minutes of preventive service) (payable for Medicare only)
- G0516 Insertion of non-biodegradable drug delivery implants, 4 or more (services for subdermal rod implant)
- G0517 Removal of non-biodegradable drug delivery implants, 4 or more (services for subdermal implants)
- G0518 Removal with reinsertion, non-biodegradable drug delivery implants, 4 or more (services for subdermal implants)
- G0561 Tympanostomy with local or topical anesthesia and insertion of a ventilating tube when performed with tympanostomy tube delivery device, unilateral
- J7318 J7329 Hyaluronic acid derivatives (Covered for Priority Medicare only)
- J8670 Rolapitant, oral, 1 mg
- J3490 Unclassified Drugs (Explanatory notes must accompany claims billed with unlisted codes) Not covered when submitted for Ketamine or other not covered drugs.

  All associated services are also excluded.
- K1007 Bilateral hip, knee, ankle, foot device, powered, includes pelvic component, single or double upright(s), knee joints any type, with or without ankle joints any type, includes all components and accessories, motors, microprocessors, sensors
- K1030 External recharging system for battery (internal) for use with implanted cardiac contractility modulation generator, replacement only (Covered for Medicaid)
- K1035 Molecular diagnostic test reader, nonprescription self-administered and self-collected use, fda approved, authorized or cleared
- Q2026 Injection, Radiesse, 0.1 ml (Covered for Medicare per LCD L39051)
- Q2028 Injection, sculptra, 0.5 mg (Covered for Medicare per LCD L39051)
- Oncology (colorectal), quantitative assessment of three urine metabolites (ascorbic acid, succinic acid and carnitine) by liquid chromatography with tandem mass spectrometry (LC-MS/MS) using multiple reaction monitoring acquisition, algorithm reported as likelihood of adenomatous polyps
- 0007U Drug test(s), presumptive, with definitive confirmation of positive results, any number of drug classes, urine, includes specimen verification including DNA authentication in comparison to buccal DNA, per date of service
- 0008U Heliobacter pylori detection and antibiotic resistance, DNA, 16S and 23S rRNA, gyrA, pbp1, rdxA and rpoB, next generation sequencing, formalin-fixed paraffin embedded or or fresh tissue, predictive, reported as positive or negative for resistance to clarithryomycin, fluoroquinolones, metronidazole, amoxicillin, tetracycline and rifabutin
- 0009U Oncology (breast cancer), ERBB2 (HER2) copy number by FISH, tumor cells from formalin fixed paraffin embedded tissue isolated using image-based dielectrophoresis (DEP) sorting, reported as ERBB2 gene amplified or non-amplified
- 0010U Infectious disease (bacterial) strain typing by whole genome sequencing, phylogenetic-based report of strain relatedness, per submitted isolate
- O011U Prescription drug monitoring, evaluation of drugs present by LC-MS/MS, using oral fluid, reported as a comparison to an estimated steady-state range, per date of service including all drug compounds and metabolites
- Oncology (prostate), detection of 8 autoantibodies (ARF 6, NKX3-1, 5'- UTR-BMI1, CEP 164, 3'-UTRRopporin, Desmocollin, AURKAIP-1, CSNK2A2), multiplexed immunoassay and flow cytometry serum, algorithm reported as risk score



0024U	Glycosylated acute phase proteins (GlycA), nuclear magnetic resonance			
002.0	spectroscopy, quantitative			
0025U	1 17. 1			
	urine, quantitative			
0035U	Neurology (prion disease), cerebrospinal fluid, detection of prion protein by quaking			
000011	induced conformational conversion, qualitative			
0038U	Vitamin D, 25 hydroxy D2 and D3, by LCMS/MS, serum microsample, quantitative			
0039U	Deoxyribonucleic acid (DNA) antibody, double stranded, high avidity			
0041U	Borrelia burgdorferi, antibody detection of 5 recombinant protein groups, by immunoblot, IgM			
0042U	Borrelia burgdorferi, antibody detection of 12 recombinant protein groups, by immunoblot, IgG			
0043U	Tick-borne relapsing fever Borrelia group, antibody detection to 4 recombinant			
	protein groups, by immunoblot, IgM			
0044U	Tick-borne relapsing fever Borrelia group, antibody detection to 4 recombinant			
	protein groups, by immunoblot, IgG			
0051U	Prescription drug monitoring, evaluation of drugs present by LC-MS/MS, urine, 31			
	drug panel, reported as quantitative results, detected or not detected, per date of			
005011	service			
0052U	Lipoprotein, blood, high resolution fractionation and quantitation of lipoproteins,			
	including all five major lipoprotein classes and subclasses of HDL, LDL, and VLDL by vertical auto profile ultracentrifugation			
0054U	Prescription drug monitoring, 14 or more classes of drugs and substances,			
00540	definitive tandem mass spectrometry with chromatography, capillary blood,			
	quantitative report with therapeutic and toxic ranges, including steady-state range			
	for the prescribed dose when detected, per date of service			
0058U	U Oncology (Merkel cell carcinoma), detection of antibodies to the Merkel cell			
	polyoma virus oncoprotein (small T antigen), serum, quantitative			
0059U	Oncology (Merkel cell carcinoma), detection of antibodies to the Merkel cell			
	polyoma virus capsid protein (VP1), serum, reported as positive or negative			
0061U	Transcutaneous measurement of five biomarkers (tissue oxygenation [StO2],			
oxyhemoglobin [ctHbO2], deoxyhemoglobin [ctHbR], papillary and reticular dermal hemoglobin concentrations [ctHb1 and ctHb2]), using spatial frequency				
	dermal hemoglobin concentrations [ctHb1 and ctHb2]), using spatial frequency			
00/211	domain imaging (SFDI) and multi-spectral analysis			
0062U	Autoimmune (systemic lupus erythematosus) IgG and Igm analysis of 80 biomarkers, utilizing serum, and algorithm reported with a risk score			
0063U	Neurology (autism), 32 amines by LCMS/MS, using plasma, and algorithm			
00030	reported as metabolic signature associate with autism spectrum disorder			
0064U	Antibody, Treponema pallidum, total and rapid plasma regain (RPR),			
00010	immunoassay, qualitative			
0065U	Syphilis test, non-treponemal antibody, immunoassay, qualitative (RPR)			
0077U	Immunoglobulin paraprotein (M-protein), qualitative, immunoprecipitation and			
	mass spectrometry, blood or urine, including isotype			
0079U	Comparative DNA analysis using multiple selected single-nucleotide			
	polymorphisms			
0080U	Oncology (lung), mass spectrometric analysis of galectin-3-binding protein and			
	scavenger receptor cysteine-rich type 1 protein M130, with five clinical risk factors			
	(age, smoking status, nodule diameter, nodule-spiculation status and nodule			
	location), utilizing plasma, algorithm reported as a categorical probability of			
	malignancy (Covered for Medicare)			



0082U	Drug test(s), definitive, 90 or more drugs or substances, definitive chromatography with mass spectrometry, and presumptive, any number of drug classes, by
	instrument chemistry analyzer (utilizing immunoassay), urine, report of presence or absence of each drug, drug metabolite or substance with description and severity of significant interactions per date of service
0083U	Oncology, response to chemotherapy drugs using motility contrast tomography, fresh or frozen tissue, reported as likelihood of sensitivity or resistance to drugs or drug combinations
0091U	Oncology (colorectal) screening, cell enumeration of circulating tumor cells, utilizing whole blood, algorithm, for the presence of adenoma or cancer, reported as a positive or negative result
0092U	Oncology (lung), three protein biomarkers, immunoassay using magnetic nanosensor technology, plasma, algorithm reported as risk score for likelihood of malignancy
0093U	Prescription drug monitoring, evaluation of 65 common drugs by LC-MS/MS, urine, each drug reported detected or not detected
0095U	Eosinophilic esophagitis, 2 protein biomarkers (Eotaxin-3 [CCL26 {C-C motif chemokine ligand 26}] and Major Basic Protein [PRG2 {proteoglycan 2, pro eosinophil major basic protein}]), enzyme-linked immunosorbent assays (ELISA), specimen obtained by
	esophageal string test device, algorithm reported as probability of active or inactive eosinophilic esophagitis
0096U	Human papillomavirus (HPV), high-risk types (ie, 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, 68), male urine
0105U	Nephrology (chronic kidney disease), multiplex electrochemiluminescent immunoassay (ECLIA) of tumor necrosis factor receptor 1A, receptor superfamily 2 (TNFR1, TNFR2), and kidney injury molecule-1 (KIM-1) combined with
	longitudinal clinical data, including APOL1 genotype if available, and plasma (isolated fresh or frozen), algorithm reported as probability score for rapid kidney function decline (RKFD)
0106U	Gastric emptying, serial collection of 7 timed breath specimens, non-radioisotope carbon-13 (13C) spirulina substrate, analysis of each specimen by gas isotope ratio mass spectrometry, reported as rate of 13CO2 excretion
0107U	Clostridium difficile toxin(s) antigen detection by immunoassay technique, stool, qualitative, multiple-step method (Covered for Commercial & Medicare)
0108U	Gastroenterology (Barrett's esophagus), whole slide—digital imaging, including morphometric analysis, computer-assisted quantitative immunolabeling of 9 protein biomarkers (p16, AMACR, p53, CD68, COX-2, CD45RO, HIF1a, HER-2, K20) and morphology, formalin-fixed paraffin-embedded tissue, algorithm reported as risk of progression to high-grade dysplasia or cancer
0109U	Infectious disease (Aspergillus species), real-time PCR for detection of DNA from 4 species (A. fumigatus, A. terreus, A. niger, and A. flavus), blood, lavage fluid, or
0110U	tissue, qualitative reporting of presence or absence of each species Prescription drug monitoring, one or more oral oncology drug(s) and substances, definitive tandem mass spectrometry with chromatography, serum or plasma from capillary blood or venous blood, quantitative report with steady-state range for the
0116U	prescribed drug(s) when detected Prescription drug monitoring, enzyme immunoassay of 35 or more drugs confirmed with LC-MS/MS, oral fluid, algorithm results reported



0117U	Pain management, analysis of 11 endogenous analytes (methylmalonic acid, xanthurenic acid, homocysteine, pyroglutamic acid, vanilmandelate, 5-
	hydroxyindoleacetic acid, hydroxymethylglutarate, ethylmalonate, 3-
	hydroxypropyl mercapturic acid (3-HPMA), quinolinic acid, kynurenic acid), LC-
	MS/MS, urine, algorithm reported as a pain-index score with likelihood of atypical biochemical function associated with pain
0118U	Transplantation medicine, quantification of donor-derived cell-free DNA using
01100	whole genome next-generation sequencing, plasma, reported as percentage of
	donor-derived cell-free DNA in the total cell-free DNA (Covered for Medicare)
0119U	Cardiology, ceramides by liquid chromatography–tandem mass spectrometry,
	plasma, quantitative report with risk score for major cardiovascular events
0121U	Sickle cell disease, microfluidic flow adhesion (VCAM-1), whole blood
0122U	Sickle cell disease, microfluidic flow adhesion (P-Selectin), whole blood
0123U	Mechanical fragility, RBC, shear stress and spectral analysis profiling
0140U	Infectious disease (fungi), fungal pathogen identification, DNA (15 fungal targets),
	blood culture, amplified probe technique, each target reported as detected or not
01.4111	detected
0141U	Infectious disease (bacteria and fungi), gram-positive organism identification and
	drug resistance element detection, DNA (20 gram-positive bacterial targets, 4 resistance genes, 1 pan gram-negative bacterial target, 1 pan Candida target), blood
	culture, amplified probe technique, each target reported as detected or not detected
0142U	Infectious disease (bacteria and fungi), gram-negative bacterial identification and
01.20	drug resistance element detection, DNA (21 gram-negative bacterial targets, 6
	resistance genes, 1 pan gram-positive bacterial target, 1 pan Candida target),
	amplified probe technique, each target reported as detected or not detected
0174U	Oncology (solid tumor), mass spectrometric 30 protein targets, formalin-fixed
	paraffin-embedded tissue, prognostic and predictive algorithm reported as likely,
	unlikely, or uncertain benefit of 39 chemotherapy and targeted therapeutic
000611	oncology agents
0206U	Neurology (Alzheimer disease); cell aggregation using morphometric imaging and
	protein kinase C-epsilon (PKCe) concentration in response to amylospheroid treatment by ELISA, cultured skin fibroblasts, each reported as positive or negative
	for Alzheimer disease
0207U	Neurology (Alzheimer disease); quantitative imaging of phosphorylated ERK1 and
02070	ERK2 in response to bradykinin treatment by in situ immunofluorescence, using
	cultured skin fibroblasts, reported as a probability index for Alzheimer disease
	(List separately in addition to code for primary procedure)
0210U	Syphilis test, non-treponemal antibody, immunoassay, quantitative (RPR)
0303U	Hematology, red blood cell (RBC) adhesion to endothelial/subendothelial adhesion
	molecules, functional assessment, whole blood, with algorithmic analysis and
	result reported as an RBC adhesion index; hypoxic
0304U	Hematology, red blood cell (RBC) adhesion to endothelial/subendothelial adhesion
	molecules, functional assessment, whole blood, with algorithmic analysis and
0305U	result reported as an RBC adhesion index; normoxic Hematology, red blood cell (RBC) functionality and deformity as a function of
03030	shear stress, whole blood, reported as a maximum elongation index
0311U	Infectious disease (bacterial), quantitative antimicrobial susceptibility reported as
	phenotypic minimum inhibitory concentration (MIC)–based antimicrobial
	susceptibility for each organisms identified
	- · · · · · · · · · · · · · · · · · · ·



0312U	Autoimmune diseases (eg, systemic lupus erythematosus [SLE]), analysis of 8 IgG autoantibodies and 2 cell-bound complement activation products using enzymelinked immunosorbent immunoassay (ELISA), flow cytometry and indirect immunofluorescence, serum, or plasma and whole blood, individual components reported along with an algorithmic SLE-likelihood assessment
0316U 0321U	Borrelia burgdorferi (Lyme disease), OspA protein evaluation, urine Infectious agent detection by nucleic acid (DNA or RNA), genitourinary pathogens, identification of 20 bacterial and fungal organisms and identification of
0322U	16 associated antibiotic-resistance genes, multiplex amplified probe technique Neurology (autism spectrum disorder [ASD]), quantitative measurements of 14 acyl carnitines and microbiome-derived metabolites, liquid chromatography with tandem mass spectrometry (LC-MS/MS), plasma, results reported as negative or
0337U	positive for risk of metabolic subtypes associated with ASD Oncology (plasma cell disorders and myeloma), circulating plasma cell immunologic selection, identification, morphological characterization, and enumeration of plasma cells based on differential CD138, CD38, CD19, and CD45
0338U	protein biomarker expression, peripheral blood Oncology (solid tumor), circulating tumor cell selection, identification, morphological characterization, detection and enumeration based on differential EpCAM, cytokeratins 8, 18, and 19, and CD45 protein biomarkers, and
0342U	quantification of HER2 protein biomarker-expressing cells, peripheral blood Oncology (pancreatic cancer), multiplex immunoassay of C5, C4, cystatin C, factor B, osteoprotegerin (OPG), gelsolin, IGFBP3, CA125 and multiplex electrochemiluminescent immunoassay (ECLIA) for CA19-9, serum, diagnostic
0344U	algorithm reported qualitatively as positive, negative, or borderline Hepatology (nonalcoholic fatty liver disease [NAFLD]), semiquantitative evaluation of 28 lipid markers by liquid chromatography with tandem mass spectrometry (LC-MS/MS), serum, reported as at-risk for nonalcoholic steatohepatitis (NASH) or not NASH
0351U	Infectious disease (bacterial or viral), biochemical assays, tumor necrosis factor-related apoptosis-inducing ligand (TRAIL), interferon gamma-induced protein-10 (IP-10), and C-reactive protein, serum, or venous whole blood, algorithm reported as likelihood of bacterial infection
0358U	Neurology (mild cognitive impairment), analysis of B-amyloid 1-42 and 1-40, chemiluminescence enzyme immunoassay, cerebral spinal fluid, reported as positive, likely positive, or negative
0359U	Oncology (prostate cancer), analysis of all prostate-specific antigen (PSA) structural isoforms by phase separation and immunoassay, plasma, algorithm reports risk of cancer
0360U	Oncology (lung), enzyme-linked immunosorbent assay (ELISA) of 7 autoantibodies (p53, NY-ESO-1, CAGE, GBU4-5, SOX2, MAGE A4, and HuD),
0361U 0365U	plasma, algorithm reported as a categorical result for risk of malignancy Neurofilament light chain, digital immunoassay, plasma, quantitative Oncology (bladder), analysis of 10 protein biomarkers (A1AT, ANG, APOE, CA9, IL8, MMP9, MMP10, PAI1, SDC1 and VEGFA) by immunoassays, urine, diagnostic algorithm, including patient's age, race and gender, reported as a probability of harboring urothelial bladder cancer
0367U	Oncology (bladder), analysis of 10 protein biomarkers (A1AT, ANG, APOE, CA9, IL8, MMP9, MMP10, PAI1, SDC1 and VEGFA) by immunoassays, urine,



	diagnostic algorithm reported as a risk score for probability of rapid recurrence of recurrent or persistent cancer following transurethral resection
0385U	Nephrology (chronic kidney disease), apolipoprotein A4 (ApoA4), CD5 antigen-like (CD5L), and insulin-like growth factor binding protein 3 (IGFBP3) by
	enzyme-linked immunoassay (ELISA), plasma, algorithm combining results with
	HDL, estimated glomerular filtration rate (GFR) and clinical data reported as a risk
	score for developing diabetic kidney disease
0387U	Oncology (melanoma), autophagy and beclin 1 regulator 1 (AMBRA1) and loricrin
	(AMLo) by immunohistochemistry, formalin-fixed paraffin-embedded (FFPE) tissue, report for risk of progression
0390U	Obstetrics (preeclampsia), kinase insert domain receptor (KDR), Endoglin (ENG),
	and retinol binding protein 4 (RBP4), by immunoassay, serum, algorithm reported as a risk score
0393U	Neurology (eg, Parkinson disease, dementia with Lewy bodies), cerebrospinal fluid
	(CSF), detection of misfolded $\alpha$ -synuclein protein by seed amplification assay, qualitative
0399U	Neurology (cerebral folate deficiency), serum, detection of anti-human folate
	receptor IgG-binding antibody and blocking autoantibodies by enzyme-linked
	immunoassay (ELISA), qualitative, and blocking autoantibodies, using a functional
0.40.41	blocking assay for IgG or IgM, quantitative, reported as positive or not detected
0404U	Oncology (breast), semiquantitative measurement of thymidine kinase activity by
0406U	immunoassay, serum, results reported as risk of disease progression Oncology (lung), flow cytometry, sputum, 5 markers (meso-tetra [4-
04000	carboxyphenyl] porphyrin [TCPP], CD206, CD66b, CD3, CD19), algorithm
0412U	reported as likelihood of lung cancer Beta amyloid, AB42/40 ratio, immunoprecipitation with quantitation by liquid
04120	chromatography with tandem mass spectrometry (LC-MS/MS) and qualitative
	ApoE isoform-specific proteotyping, plasma combined with age, algorithm
	reported as presence or absence of brain amyloid pathology
0427U	Monocyte distribution width, whole blood (List separately in addition to code for
	primary procedure)
0431U	Glycine receptor alpha1 IgG, serum or cerebrospinal fluid (CSF), live cell-binding
042211	assay (LCBA), qualitative
0432U	Kelch-like protein 11 (KLHL11) antibody, serum or cerebrospinal fluid (CSF), cell-binding assay, qualitative
	assay, quantative
0435U	Oncology, chemotherapeutic drug cytotoxicity assay of cancer stem cells (CSCs),
	from cultured CSCs and primary tumor cells, categorical drug response reported
	based on cytotoxicity percentage observed, minimum of 14 drugs or drug
	combinations
0436U	Oncology (lung), plasma analysis of 388 proteins, using aptamer-based proteomics
	technology, predictive algorithm reported as clinical benefit from immune checkpoint inhibitor therapy
0441U	Infectious disease (bacterial, fungal, or viral infection), semiquantitative
01110	biomechanical assessment (via deformability cytometry), whole blood, with
	algorithmic analysis and result reported as an index
0442U	Infectious disease (respiratory infection), Myxovirus resistance protein A (MxA)
	and C-reactive protein (CRP), fingerstick whole blood specimen, each biomarker
	reported as present or absent



0445U	B-amyloid (Abeta42) and phospho tau (181P) (pTau181), electrochemiluminescent immunoassay (ECLIA), cerebral spinal fluid, ratio reported as positive or negative
0446U	for amyloid pathology Autoimmune diseases (systemic lupus erythematosus [SLE]), analysis of 10 cytokine soluble mediator biomarkers by immunoassay, plasma, individual components reported with an algorithmic risk score for current disease activity
0447U	Autoimmune diseases (systemic lupus erythematosus [SLE]), analysis of 11 cytokine soluble mediator biomarkers by immunoassay, plasma, individual components reported with an algorithmic prognostic risk score for developing a clinical flare
0450U	Oncology (multiple myeloma), liquid chromatography with tandem mass spectrometry (LCMS/MS), monoclonal paraprotein sequencing analysis, serum, results reported as baseline presence or absence of detectable clonotypic peptides
0451U	Oncology (multiple myeloma), LCMS/MS, peptide ion quantification, serum, results compared with baseline to determine monoclonal paraprotein abundance
0457U	Perfluoroalkyl substances (PFAS) (eg, perfluorooctanoic acid, perfluorooctane sulfonic acid), 9 PFAS compounds by LC-MS/MS, plasma or serum, quantitative
0458U	Oncology (breast cancer), S100A8 and S100A9, by enzymelinked immunosorbent assay (ELISA), tear fluid with age, algorithm reported as a risk score
0459U	β-amyloid (Abeta42) and total tau (tTau), electrochemiluminescent immunoassay (ECLIA), cerebral spinal fluid, ratio reported as positive or negative for amyloid pathology
0462U	Melatonin levels test, sleep study, 7 or 9 sample melatonin profile (cortisol optional), enzyme-linked immunosorbent assay (ELISA), saliva, screening/preliminary
0468U	Hepatology (nonalcoholic steatohepatitis [NASH]), miR-34a5p, alpha 2-macroglobulin, YKL40, HbA1c, serum and whole blood, algorithm reported as a single score for NASH activity and fibrosis
0472U	Carbonic anhydrase VI (CA VI), parotid specific/secretory protein (PSP) and salivary protein (SP1) IgG, IgM, and IgA antibodies, enzyme-linked immunosorbent assay (ELISA), semiqualitative, blood, reported as predictive evidence of early Sjögren syndrome
0479U	Tau, phosphorylated, pTau217
0482U	Obstetrics (preeclampsia), biochemical assay of soluble fmslike tyrosine kinase 1 (sFlt-1) and placental growth factor (PlGF), serum, ratio reported for sFlt1/PlGF, with risk of progression for preeclampsia with severe features within 2 weeks
0503U	Neurology (Alzheimer disease), beta amyloid (Aβ40, Aβ42, Aβ42/40 ratio) and tau-protein (ptau217, nptau217, ptau217/nptau217 ratio), blood, immunoprecipitation with quantitation by liquid chromatography with tandem mass spectrometry (LC-MS/MS), algorithm score reported as likelihood of positive or negative for amyloid plaques
0521U	Rheumatoid factor IgA and IgM, cyclic citrullinated peptide (CCP) antibodies, and scavenger receptor A (SR-A) by immunoassay, blood
0522U	Carbonic anhydrase VI, parotid specific/secretory protein and salivary protein 1 (SP1), IgG, IgM, and IgA antibodies, chemiluminescence, semiqualitative, blood
0524U	Obstetrics (preeclampsia), sFlt1/PIGF ratio, immunoassay, utilizing serum or plasma, reported as a value
0525U	Oncology, spheroid cell culture, 11-drug panel (carboplatin, docetaxel, doxorubicin, etoposide, gemcitabine, niraparib, olaparib, paclitaxel, rucaparib,



	topotecan, veliparib) ovarian, fallopian, or peritoneal response prediction for each drug
0526U	Nephrology (renal transplant), quantification of CXCL10 chemokines, flow cytometry, urine, reported as pg/mL creatinine baseline and monitoring over time
0535U	Perfluoroalkyl substances (PFAS) (eg, perfluorooctanoic acid, perfluorooctane sulfonic acid), by liquid chromatography with tandem mass spectrometry (LC-MS/MS), plasma or serum, quantitative
0545U	Acetylcholine receptor (AChR), antibody identification by immunofluorescence, using live cells, reported as positive or negative
0546U	Low-density lipoprotein receptor-related protein 4 (LRP4), antibody identification by immunofluorescence, using live cells, reported as positive or negative
0547U	Neurofilament light chain (NfL), chemiluminescent enzyme immunoassay, plasma, quantitative
0548U	Glial fibrillary acidic protein (GFAP), chemiluminescent enzyme immunoassay, using plasma
0550U	Oncology (prostate), enzyme-linked immunosorbent assays (ELISA) for total prostate-specific antigen (PSA) and free PSA, serum, combined with age, previous negative prostate biopsy status, digital rectal examination findings, prostate volume, and image and data reporting of the prostate, algorithm reported as a risk score for the presence of high-grade prostate cancer
0551U	Tau, phosphorylated, pTau217, by single-molecule array (ultrasensitive digital protein detection), using plasma



Experimental/Investigational/ Unproven Care/Benefit Exceptions

# APPENDIX A ADVANCE CARE PLANNING ASSESSMENT

I.	Medical history and reason for referral:
2.	Patient's understanding of current disease status and overall prognosis:
	Medical care options discussed with patient:
3.	Has patient completed an Advance Care Planning conversation, including designation of patient advocate as part of the advance directive, with a certified ACP facilitator*? Yes No If no, answer questions 4-9. If yes, this form is complete.
4.	What are patient's wishes/goals for remainder of life (quality of life vs. length of life; importance of physical comfort; how patient wishes to spend time, etc.)?
5.	How does patient describe their current physical/mental symptoms? What is quality of life rating using QOL, HR QOL scale, SF 36 (short-form health questionnaire)?
6.	Spiritual or cultural beliefs related to illness and death that would affect enrollment? Yes \[ \] No \[ \]
7.	Is advance directive complete? Yes No (i.e. Making Choices Michigan)
8.	Patient has designated a durable power of attorney for healthcare? Yes 🗌 No 🗍
9.	Does family/patient advocate support patient's preference for medical care as outlined in advance directive? Yes \bigcap No \bigcap
Tr	Certified ACP facilitators are trained through the Respecting Choices® curriculum. ained facilitators are available at health systems, Making Choices Michigan, and mmunity organizations.

#### APPENDIX B

#### CLINICAL TRIALS COVERAGE REFERENCE SHEET\*\*\*

	Commercial Fully- funded	Commercial Self-funded	Medicare
Clinical Trials	Routine services* only, use Clinical Trials Policy #91606	Non-grandfathered groups: routine services only, use Clinical Trials Policy #91606	Original Medicare covers routine services for those trials that are Medicare approved
		Grandfathered groups opting out of PPACA: use Clinical Trials for Self Funded groups opting out of PPACA #91448	If trial is not Medicare approved, there is no coverage under Original Medicare or Priority Health Medicare.
IDE (Investigational Device Exemption) Trial: Category A Device	Never covered. Device and all services, including routine services, are not covered. Use Experimental & Investigational Policy #91117	Never covered. Device and all services, including routine services, are not covered. Use Experimental & Investigational Policy #91117	Device is never covered. Routine care items and services in CMS-approved Category A IDE studies are covered by Priority Health Medicare
IDE Trial: Category B Device	Routine services only; device not covered.** Use Experimental & Investigational Policy #91117	Device and all services, including routine services, are not covered.** Use Experimental & Investigational Policy #91117	All services, including the device, are covered by Priority Health Medicare
Clinical Studies Approved Under Evidence Development (CED)	Use Experimental & Investigational Policy #91117 to determine coverage	Use Experimental & Investigational Policy #91117 and individual plan documents to determine coverage	All care and services are covered by Priority Health Medicare

<sup>\*</sup>Routine patient care costs are items or services that are typically covered benefits when provided outside a clinical trial. The clinical trial protocol may be needed to determine the specific services that are covered and excluded.

<sup>\*\*</sup>Priority Health may, at its discretion, choose to cover the experimental device if the cost of that device is less than the non-experimental arm of the trial.

<sup>\*\*\*</sup>For Medicaid/Healthy Michigan refer to Section III "Application to Products"



Experimental/Investigational/ Unproven Care/Benefit Exceptions

#### APPENDIX C

#### HUMANITARIAN USE DEVICE (HUD)/ HUMANITARIAN DEVICE EXEMPTION (HDE) REFERENCE SHEET

The following HUDs/HDEs may be covered when used in accordance with their FDA approval

	<b>HUD/HDE Covered Devices</b>	Medical Policy Supporting Coverage
1.	Activa Dystonia Therapy (Medtronic)	Stimulation Therapy and Devices medical policy #91468
2.	Impella circulatory assistance	Ventricular Assist Devices medical policy #91509
3.	Enterra Therapy System	Gastroparesis Testing and Treatment medical policy
		#91572
4.	Epicel (cultured epidermal autografts)	Skin Substitutes and Soft Tissue Grafts medical policy
		#91560
5.	INTACS for keratoconus	Vision Care medical policy #91538
6.	NeuRX diaphragmatic stimulator for	Stimulation Therapy and Devices medical policy #91468
	spinal cord injury	

**Note:** Devices that have FDA approval for humanitarian use or as HDEs are considered experimental and investigational and excluded from coverage unless they are listed above.

The FDA list of HDEs can be found at https://www.fda.gov/medical-devices/hde-approvals/listing-cdrh-humanitarian-device-exemptions (Accessed March 12, 2025)

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