### Contractor Information

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<th>CONTRACTOR NAME</th>
<th>CONTRACT TYPE</th>
<th>CONTRACT NUMBER</th>
<th>JURISDICTION</th>
<th>STATE(S)</th>
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<td>DME MAC</td>
<td>17013 - DME MAC</td>
<td>J-B</td>
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<td>18003 - DME MAC</td>
<td>J-C</td>
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<td>CONTRACTOR NAME</td>
<td>CONTRACT TYPE</td>
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**LCD Information**

**Document Information**

**LCD ID**
L33789

**Original ICD-9 LCD ID**
L27239
L23613
L21271
L23598

**LCD Title**
Power Mobility Devices

**Original Effective Date**
For services performed on or after 10/01/2015

**Revision Effective Date**
For services performed on or after 01/01/2019

**Revision Ending Date**
N/A

**Retirement Date**
N/A

**Proposed LCD in Comment Period**
N/A

**Notice Period Start Date**
N/A

**Source Proposed LCD**
N/A

**Notice Period End Date**
N/A
CMS National Coverage Policy

CMS Pub. 100-3, Medicare National Coverage Determinations Manual, Chapter 1, Sections 280.3

Coverage Guidance

Coverage Indications, Limitations, and/or Medical Necessity

For any item to be covered by Medicare, it must 1) be eligible for a defined Medicare benefit category, 2) be reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member, and 3) meet all other applicable Medicare statutory and regulatory requirements.

The purpose of a Local Coverage Determination (LCD) is to provide information regarding “reasonable and necessary” criteria based on Social Security Act § 1862(a)(1)(A) provisions.
In addition to the "reasonable and necessary" criteria contained in this LCD there are other payment rules, which are discussed in the following documents, that must also be met prior to Medicare reimbursement:

- The LCD-related Standard Documentation Requirements Article, located at the bottom of this policy under the Related Local Coverage Documents section.
- The LCD-related Policy Article, located at the bottom of this policy under the Related Local Coverage Documents section.
- Refer to the Supplier Manual for additional information on documentation requirements.
- Refer to the DME MAC web sites for additional bulletin articles and other publications related to this LCD.

For the items addressed in this LCD, the "reasonable and necessary" criteria, based on Social Security Act § 1862(a)(1)(A) provisions, are defined by the following coverage indications, limitations and/or medical necessity.

Power Mobility Device bases require a 7-element order. Refer to this LCD’s related Policy Article for this statutory requirement. Other separately billable wheelchair options and accessories require a detailed product description (DPD). Refer to the Standard Documentation Requirements for All Claims Submitted to DME MACs Policy Article for information about the DPD.

Refer to this related Policy Article for information on the face-to-face examination.

The term power mobility device (PMD) includes power operated vehicles (POVs) and power wheelchairs (PWCs).

**GENERAL COVERAGE CRITERIA:**

All of the following basic criteria (A-C) must be met for a power mobility device (K0800-K0898) or a push-rim activated power assist device (E0986) to be covered. Additional coverage criteria for specific devices are listed below.

A. The beneficiary has a mobility limitation that significantly impairs his/her ability to participate in one or more mobility-related activities of daily living (MRADLs) such as toileting, feeding, dressing, grooming, and bathing in customary locations in the home. A mobility limitation is one that:
   - Prevents the beneficiary from accomplishing an MRADL entirely, or
   - Places the beneficiary at reasonably determined heightened risk of morbidity or mortality secondary to the attempts to perform an MRADL; or
   - Prevents the beneficiary from completing an MRADL within a reasonable time frame.

B. The beneficiary’s mobility limitation cannot be sufficiently and safely resolved by the use of an appropriately fitted cane or walker.

C. The beneficiary does not have sufficient upper extremity function to self-propel an optimally-configured manual wheelchair in the home to perform MRADLs during a typical day.
   - Limitations of strength, endurance, range of motion, or coordination, presence of pain, or deformity or absence of one or both upper extremities are relevant to the assessment of upper extremity function.
   - An optimally-configured manual wheelchair is one with an appropriate wheelbase, device weight, seating options, and other appropriate nonpowered accessories.

**POWER OPERATED VEHICLES (K0800-K0808, K0812):**

A POV is covered if all of the basic coverage criteria (A-C) have been met and if criteria D-I are also met.

D. The beneficiary is able to:
   - Safely transfer to and from a POV, and
Operate the tiller steering system, and
Maintain postural stability and position while operating the POV in the home.

E. The beneficiary’s mental capabilities (e.g., cognition, judgment) and physical capabilities (e.g., vision) are sufficient for safe mobility using a POV in the home.

F. The beneficiary’s home provides adequate access between rooms, maneuvering space, and surfaces for the operation of the POV that is provided.

G. The beneficiary’s weight is less than or equal to the weight capacity of the POV that is provided and greater than or equal to 95% of the weight capacity of the next lower weight class POV – i.e., a Heavy Duty POV is covered for a beneficiary weighing 285 – 450 pounds; a Very Heavy Duty POV is covered for a beneficiary weighing 428 – 600 pounds.

H. Use of a POV will significantly improve the beneficiary’s ability to participate in MRADLs and the beneficiary will use it in the home.

I. The beneficiary has not expressed an unwillingness to use a POV in the home.

If a POV will be used inside the home and coverage criteria A-I are not met, it will be denied as not reasonable and necessary.

Group 2 POVs (K0806-K0808) have added capabilities that are not needed for use in the home. Therefore, if a Group 2 POV is provided it will be denied as not reasonable and necessary.

If a POV will only be used outside the home, see related Policy Article for information concerning noncoverage.

POWER WHEELCHAIRS (K0013, K0813-K0891, K0898):

A power wheelchair is covered if:

a. All of the basic coverage criteria (A-C) are met; and
b. The beneficiary does not meet coverage criterion D, E, or F for a POV; and
c. Either criterion J or K is met; and
d. Criteria L, M, N, and O are met; and
e. Any coverage criteria pertaining to the specific wheelchair type (see below) are met.

J. The beneficiary has the mental and physical capabilities to safely operate the power wheelchair that is provided; or
K. If the beneficiary is unable to safely operate the power wheelchair, the beneficiary has a caregiver who is unable to adequately propel an optimally configured manual wheelchair, but is available, willing, and able to safely operate the power wheelchair that is provided; and
L. The beneficiary’s weight is less than or equal to the weight capacity of the power wheelchair that is provided and greater than or equal to 95% of the weight capacity of the next lower weight class PWC – i.e., a Heavy Duty PWC is covered for a beneficiary weighing 285 – 450 pounds; a Very Heavy Duty PWC is covered for a beneficiary weighing 428 – 600 pounds; an Extra Heavy Duty PWC is covered for a beneficiary weighing 570 pounds or more.
M. The beneficiary’s home provides adequate access between rooms, maneuvering space, and surfaces for the operation of the power wheelchair that is provided.
N. Use of a power wheelchair will significantly improve the beneficiary’s ability to participate in MRADLs and the beneficiary will use it in the home. For beneficiaries with severe cognitive and/or physical impairments, participation in MRADLs may require the assistance of a caregiver.
O. The beneficiary has not expressed an unwillingness to use a power wheelchair in the home.

If a PWC will be used inside the home and if coverage criteria (a)-(e) are not met, it will be denied as not reasonable and necessary.
If a PWC will only be used outside the home, see related Policy Article for information concerning noncoverage.

**ADDITIONAL CRITERIA FOR SPECIFIC TYPES OF POWER WHEELCHAIRS:**

I. A Group 1 PWC (K0813-K0816) or a Group 2 PWC (K0820-K0829) is covered if all of the coverage criteria (a)-(e) for a PWC are met and the wheelchair is appropriate for the beneficiary’s weight.

II. A Group 2 Single Power Option PWC (K0835 – K0840) is covered if all of the coverage criteria (a)-(e) for a PWC are met and if:
   A. Criterion 1 or 2 is met; and
   B. Criteria 3 and 4 are met.
      1. The beneficiary requires a drive control interface other than a hand or chin-operated standard proportional joystick (examples include but are not limited to head control, sip and puff, switch control).
      2. The beneficiary meets coverage criteria for a power tilt or a power recline seating system (see Wheelchair Options and Accessories policy for coverage criteria) and the system is being used on the wheelchair.
      3. The beneficiary has had a specialty evaluation that was performed by a licensed/certified medical professional, such as a physical therapist (PT) or occupational therapist (OT), or practitioner who has specific training and experience in rehabilitation wheelchair evaluations and that documents the medical necessity for the wheelchair and its special features (see Documentation Requirements section). The PT, OT, or practitioner may have no financial relationship with the supplier.
      4. The wheelchair is provided by a supplier that employs a RESNA-certified Assistive Technology Professional (ATP) who specializes in wheelchairs and who has direct, in-person involvement in the wheelchair selection for the beneficiary.

   If a Group 2 Single Power Option PWC is provided and if criterion II(A) or II(B) is not met (including but not limited to situations in which it is only provided to accommodate a power seat elevation feature, a power standing feature, or power elevating legrests), it will be denied as not reasonable and necessary.

III. A Group 2 Multiple Power Option PWC (K0841-K0843) is covered if all of the coverage criteria (a)-(e) for a PWC are met and if:
   A. Criterion 1 or 2 is met; and
   B. Criteria 3 and 4 are met.
      1. The beneficiary meets coverage criteria for a power tilt and recline seating system (see Wheelchair Options and Accessories policy) and the system is being used on the wheelchair.
      2. The beneficiary uses a ventilator which is mounted on the wheelchair.
      3. The beneficiary has had a specialty evaluation that was performed by a licensed/certified medical professional, such as a PT or OT, or practitioner who has specific training and experience in rehabilitation wheelchair evaluations and that documents the medical necessity for the wheelchair and its special features (see Documentation Requirements section). The PT, OT, or practitioner may have no financial relationship with the supplier.
      4. The wheelchair is provided by a supplier that employs a RESNA-certified Assistive Technology Professional (ATP) who specializes in wheelchairs and who has direct, in-person involvement in the wheelchair selection for the beneficiary.

   If a Group 2 Multiple Power Option PWC is provided and if criterion III(A) or III(B) is not met, it will be denied as not reasonable and necessary.

IV. A Group 3 PWC with no power options (K0848-K0855) is covered if:
   A. All of the coverage criteria (a)-(e) for a PWC are met; and
   B. The beneficiary’s mobility limitation is due to a neurological condition, myopathy, or congenital skeletal deformity; and
   C. The beneficiary has had a specialty evaluation that was performed by a licensed/certified medical professional, such as a PT or OT, or practitioner who has specific training and experience in rehabilitation wheelchair evaluations and that documents the medical necessity for the wheelchair and its special features (see Wheelchair Options and Accessories policy for coverage criteria) and the system is being used on the wheelchair.
features (see Documentation Requirements section). The PT, OT, or practitioner may have no financial relationship with the supplier; and

D. The wheelchair is provided by a supplier that employs a RESNA-certified Assistive Technology Professional (ATP) who specializes in wheelchairs and who has direct, in-person involvement in the wheelchair selection for the beneficiary.

If a Group 3 PWC is provided and if criteria (IV)(A) – (IV)(D) are not met, it will be denied as not reasonable and necessary.

V. A Group 3 PWC with Single Power Option (K0856-K0860) or with Multiple Power Options (K0861-K0864) is covered if:
   A. The Group 3 criteria IV(A) and IV(B) are met; and
   B. The Group 2 Single Power Option (criteria II[A] and II[B]) or Multiple Power Options (criteria III[A] and III[B]) (respectively) are met.

If a Group 3 Single Power Option or Multiple Power Options PWC is provided and if criterion V(A) or (V)(B) is not met, it will be denied as not reasonable and necessary.

VI. Group 4 PWCs (K0868-K0886) have added capabilities that are not needed for use in the home. Therefore, if these wheelchairs are provided they will be denied as not reasonable and necessary.

VII. A Group 5 (Pediatric) PWC with Single Power Option (K0890) or with Multiple Power Options (K0891) is covered if:
   A. The beneficiary is expected to grow in height; and
   B. The Group 2 Single Power Option (criteria II[A] and II[B]) or Multiple Power Options (criteria III[A] and III[B]) (respectively) are met.

If a Group 5 PWC is provided and if criteria (VII)(A) – (VII)(C) are not met, it will be denied as not reasonable and necessary.

VIII. A push-rim activated power assist device (E0986) for a manual wheelchair is covered if all of the following criteria are met:
   A. All of the criteria for a power mobility device listed in the Basic Coverage Criteria section are met; and
   B. The beneficiary has been self-propelling in a manual wheelchair for at least one year; and
   C. The beneficiary has had a specialty evaluation that was performed by a licensed/certified medical professional, such as a PT or OT, or practitioner who has specific training and experience in rehabilitation wheelchair evaluations and that documents the need for the device in the beneficiary’s home. The PT, OT, or practitioner may have no financial relationship with the supplier; and
   D. The wheelchair is provided by a supplier that employs a RESNA-certified Assistive Technology Professional (ATP) who specializes in wheelchairs and who has direct, in-person involvement in the wheelchair selection for the beneficiary.

If all of the coverage criteria are not met, it will be denied as not reasonable and necessary.

A custom motorized/power wheelchair base (K0013) will be covered if:

1. The beneficiary meets the general coverage criteria for a power wheelchair; and
2. The specific configurational needs of the beneficiary are not able to be met using wheelchair cushions, or options or accessories (prefabricated or custom fabricated), which may be added to another power wheelchair base.

If coverage criterion 1 for K0013 is not met, the claim will be denied as not reasonable and necessary.

If coverage criterion 2 for K0013 is not met, the claim will be denied for incorrect coding (see related Policy Article for additional information).
A custom motorized/power wheelchair base is not reasonable and necessary if the expected duration of need for the chair is less than three months (e.g., post-operative recovery).

If the PWC base is not covered, then related accessories will be denied.

**MISCELLANEOUS:**

A POV or power wheelchair with Captain's Chair is not appropriate for a beneficiary who needs a separate wheelchair seat and/or back cushion. If a skin protection and/or positioning seat or back cushion that meets coverage criteria (see Wheelchair Seating LCD) is provided with a POV or a power wheelchair with Captain's Chair, the POV or PWC will be denied as not reasonable and necessary. (Refer to Wheelchair Seating LCD and Policy Article for information concerning coverage of general use, skin protection, or positioning cushions when they are provided with a POV or power wheelchair with Captain's Chair.)

For beneficiaries who do not have special skin protection or positioning needs, a power wheelchair with Captain's Chair provides appropriate support. Therefore, if a general use cushion is provided with a power wheelchair with a sling/solid seat/back instead of Captain's Chair, the wheelchair and the cushion(s) will be covered only if either criterion 1 or criterion 2 is met:

1. The cushion is provided with a covered power wheelchair base that is not available in a Captain's Chair model – i.e., codes K0839, K0840, K0843, K0860 – K0864, K0890, K0891; or
2. A skin protection and/or positioning seat or back cushion that meets coverage criteria is provided.

If one of these criteria is not met, both the power wheelchair with a sling/solid seat and the general use cushion will be denied as not reasonable and necessary.

If a heavy duty, very heavy duty, or extra heavy duty PWC or POV is provided and if the beneficiary’s weight is outside the range listed in criterion G or L above (i.e., for heavy duty – 285 – 400 pounds, for very heavy duty – 428 – 600 pounds, for extra heavy duty – 570 pounds or more), it will be denied as not reasonable and necessary.

Refer to the related Policy Article for information concerning coverage of Group 2 PWCs with seat elevators (K0830, K0831).

The delivery of the PMD must be within 120 days following completion of the face-to-face examination. (Exception: For PWCs that go through Advance Determination of Medicare Coverage (ADMC) or Prior Authorization (PA) and receive an affirmative determination, the delivery must be within 6 months following the determination.)

An add-on to convert a manual wheelchair to a joystick-controlled power mobility device (E0983) or to a tiller-controlled power mobility device (E0984) will be denied as not reasonable and necessary.

Payment is made for only one wheelchair at a time. Backup chairs are denied as not reasonable and necessary.

One month's rental of a PWC or POV (K0462) is covered if a beneficiary-owned wheelchair is being repaired. Payment is based on the type of replacement device that is provided but will not exceed the rental allowance for the power mobility device that is being repaired.

A power mobility device will be denied as not reasonable and necessary if the underlying condition is reversible and
the length of need is less than 3 months (e.g., following lower extremity surgery which limits ambulation).

A POV or PWC which has not been reviewed by the Pricing, Data Analysis, and Coding (PDAC) contractor or which has been reviewed by the PDAC and found not to meet the definition of a specific POV/PWC will be denied as not reasonable and necessary and should be coded as K0899.

GENERAL

A Detailed Written Order (DWO) (if applicable) must be received by the supplier before a claim is submitted. If the supplier bills for an item addressed in this policy without first receiving a completed DWO, the claim shall be denied as not reasonable and necessary.

For Durable Medical Equipment, Prosthetics, Orthotics and Supplies (DMEPOS) base items that require a Written Order Prior to Delivery (WOPD), the supplier must also obtain a DWO before submitting a claim for any associated options, accessories, and/or supplies that are separately billed. In this scenario, if the supplier bills for associated options, accessories, and/or supplies without first receiving a completed DWO, the claim shall be denied as not reasonable and necessary.

A WOPD (if applicable) must be received by the supplier before a DMEPOS item is delivered to a beneficiary. If a supplier delivers a DMEPOS item without first receiving a completed WOPD, the claim shall be statutorily denied. Refer to the LCD-related Policy Article, located at the bottom of this policy under the Related Local Coverage Documents section.

An item/service is correctly coded when it meets all the coding guidelines listed in CMS HCPCS guidelines, LCDs, LCD-related Policy Articles, or DME MAC articles. Claims that do not meet coding guidelines shall be denied as not reasonable and necessary/incorrectly coded.

Proof of delivery (POD) is a Supplier Standard and DMEPOS suppliers are required to maintain POD documentation in their files. Proof of delivery documentation must be made available to the Medicare contractor upon request. All services that do not have appropriate proof of delivery from the supplier shall be denied as not reasonable and necessary.

Summary of Evidence

N/A

Analysis of Evidence
(Rationale for Determination)

N/A

Coding Information
Bill Type Codes:
Contractors may specify Bill Types to help providers identify those Bill Types typically used to report this service. Absence of a Bill Type does not guarantee that the policy does not apply to that Bill Type. Complete absence of all Bill Types indicates that coverage is not influenced by Bill Type and the policy should be assumed to apply equally to all claims.

N/A

Revenue Codes:
Contractors may specify Revenue Codes to help providers identify those Revenue Codes typically used to report this service. In most instances Revenue Codes are purely advisory. Unless specified in the policy, services reported under other Revenue Codes are equally subject to this coverage determination. Complete absence of all Revenue Codes indicates that coverage is not influenced by Revenue Code and the policy should be assumed to apply equally to all Revenue Codes.

N/A

CPT/HCPCS Codes

Group 1 Paragraph:
The appearance of a code in this section does not necessarily indicate coverage.

HCPCS MODIFIERS:

EY – No physician or other licensed health care provider order for this item or service

GA - Waiver of liability statement issued as required by payer policy, individual case

GY – Item or service statutorily excluded or doesn’t meet the definition of any Medicare benefit category

GZ - Item or service expected to be denied as not reasonable and necessary

KX – Requirements specified in the medical policy have been met

HCPCS CODES:
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<th>DESCRIPTION</th>
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**ICD-10 Codes that Support Medical Necessity**

N/A

**ICD-10 Codes that DO NOT Support Medical Necessity**

**Group 1 Paragraph:**
Not specified

**Group 1 Codes: N/A**
General Information

Associated Information

DOCUMENTATION REQUIREMENTS

Section 1833(e) of the Social Security Act precludes payment to any provider of services unless “there has been furnished such information as may be necessary in order to determine the amounts due such provider.” It is expected that the beneficiary’s medical records will reflect the need for the care provided. The beneficiary’s medical records include the treating practitioner’s office records, hospital records, nursing home records, home health agency records, records from other healthcare professionals and test reports. This documentation must be available upon request.

GENERAL DOCUMENTATION REQUIREMENTS

In order to justify payment for DMEPOS items, suppliers must meet the following requirements:

- Prescription (orders)
- Medical Record Information (including continued need/use if applicable)
- Correct Coding
- Proof of Delivery

Refer to the LCD-related Standard Documentation Requirements article, located at the bottom of this policy under the Related Local Coverage Documents section for additional information regarding these requirements.

Refer to the Supplier Manual for additional information on documentation requirements.

Refer to the DME MAC web sites for additional bulletin articles and other publications related to this LCD.

POLICY SPECIFIC DOCUMENTATION REQUIREMENTS

Items covered in this LCD have additional policy-specific requirements that must be met prior to Medicare reimbursement.

Refer to the LCD-related Policy article, located at the bottom of this policy under the Related Local Coverage Documents section for additional information.

PRESCRIPTION (ORDER) REQUIREMENTS

7-ELEMENT ORDERS (PIM 5.2.4)

The order, referred to as the 7-element order, that the supplier must receive within 45 days after completion of the
face-to-face examination (see Policy Article) must contain all of the following elements:

1. Beneficiary’s name
2. Description of the item that is ordered. This may be general – e.g., “power operated vehicle”, “power wheelchair”, or “power mobility device”– or may be more specific.
3. Date of the face-to-face examination
4. Pertinent diagnoses/conditions that relate to the need for the PMD
5. Length of need
6. Prescribing physician/practitioner’s signature
7. Date the prescription was written

The supplier may provide a template order listing the seven required elements but is prohibited from completing any part of it. The treating practitioner completing the face-to-face requirements must write the 7-element order. The 7-element order may only be written after the completion of the face-to-face exam requirements. Refer to the Policy Article NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES section for information regarding the statutory requirements for PMDs.

A date stamp or equivalent must be used to document receipt date.

DETAILED PRODUCT DESCRIPTION

Once the supplier has determined the specific power mobility device that is appropriate for the beneficiary based on the practitioner’s 7-element order, the supplier must prepare a written document (termed a detailed product description). This detailed product description (DPD) must comply with the requirements for a detailed written order as outlined in the Supplier Manual and CMS’ Program Integrity Manual (Internet-Only Manual, Pub. 100-8), Chapter 5. Regardless of the form of the description, there must be sufficient detail to identify the item(s) in order to determine that the item(s) dispensed is properly coded.

The practitioner must sign and date the detailed product description and the supplier must receive it prior to delivery of the PWC or POV. Signature and date stamps are not allowed. Signatures must comply with the CMS signature requirements outlined in PIM 3.3.2.4. A date stamp or equivalent must be used to document the supplier receipt date. The detailed product description must be available on request.

A prescription is not considered as part of the medical record. Medical information intended to demonstrate compliance with coverage criteria may be included on the prescription but must be corroborated by information contained in the medical record. (PIM 5.7)

FACE-TO-FACE EXAMINATION:

The report of the face-to-face examination (see Policy Article) should provide information relating to the following questions.

For POVs and PWCs  What is this beneficiary’s mobility limitation and how does it interfere with the performance of activities of daily living?
For POVs and PWCs  Why can’t a cane or walker meet this beneficiary’s mobility needs in the home?
For POVs and PWCs  Why can’t a manual wheelchair meet this beneficiary’s mobility needs in the home?
For POVs  Does this beneficiary have the physical and mental abilities to transfer into a POV and to operate it safely in the home?
For PWCs  Why can’t a POV (scooter) meet this beneficiary’s mobility needs in the home?
For PWCs  Does this beneficiary have the physical and mental abilities to operate a power wheelchair safely?
The report should provide pertinent information about the following elements, but may include other details. Each element would not have to be addressed in every evaluation.

- History of the present condition(s) and past medical history that is relevant to mobility needs
  - Symptoms that limit ambulation
  - Diagnoses that are responsible for these symptoms
  - Medications or other treatment for these symptoms
  - Progression of ambulation difficulty over time
  - Other diagnoses that may relate to ambulatory problems
  - How far the beneficiary can walk without stopping
  - Pace of ambulation
  - What ambulatory assistance (cane, walker, wheelchair, caregiver) is currently used
  - What has changed to now require use of a power mobility device
  - Ability to stand up from a seated position without assistance
  - Description of the home setting and the ability to perform activities of daily living in the home

- Physical examination that is relevant to mobility needs
  - Weight and height
  - Cardiopulmonary examination
  - Musculoskeletal examination
    - Arm and leg strength and range of motion
  - Neurological examination
    - Gait
    - Balance and coordination

The evaluation should be tailored to the individual beneficiary’s conditions. The history should paint a picture of the beneficiary’s functional abilities and limitations on a typical day. It should contain as much objective data as possible. The physical examination should be focused on the body systems that are responsible for the beneficiary’s ambulatory difficulty or impact on the beneficiary’s ambulatory ability.

A date stamp or equivalent must be used to document the date that the supplier receives the report of the face-to-face examination. The written report of this examination must be available upon request.

Practitioners shall document the examination in a detailed narrative note in their charts in the format that they use for other entries. The note must clearly indicate that a major reason for the visit was a mobility examination.

Many suppliers have created forms which have not been approved by CMS which they send to practitioners and ask them to complete. Even if the practitioner completes this type of form and puts it in his/her chart, this supplier-generated form is not a substitute for the comprehensive medical record as noted above. Suppliers are encouraged to help educate practitioners on the type of information that is needed to document a beneficiary's mobility needs.

Practitioners shall also provide reports of pertinent laboratory tests, x-rays, and/or other diagnostic tests (e.g., pulmonary function tests, cardiac stress test, electromyogram, etc.) performed in the course of management of the beneficiary. Upon request, suppliers shall provide notes from prior visits to give a historical perspective of the progression of disease over time and to corroborate the information in the face-to-face examination.

If the report of a licensed/certified medical professional (LCMP) examination is to be considered as part of the face-to-face examination (see Policy Article), there must be a signed and dated attestation by the supplier or LCMP that the LCMP has no financial relationship with the supplier. (Note: Evaluations performed by an LCMP who has a financial relationship with the supplier may be submitted to provide additional clinical information, but will not be considered as part of the face-to-face examination by the practitioner.)
Although beneficiaries who qualify for coverage of a power mobility device may use that device outside the home, because Medicare’s coverage of a wheelchair or POV is determined solely by the beneficiary’s mobility needs within the home, the examination must clearly distinguish the beneficiary’s abilities and needs within the home from any additional needs for use outside the home.

**SPECIALTY EVALUATION:**

The specialty evaluation that is required for beneficiary’s who receive a Group 2 Single Power Option or Multiple Power Options PWC, any Group 3 PWC, or a push-rim activated power assist device is in addition to the requirement for the face-to-face examination. The specialty evaluation provides detailed information explaining why each specific option or accessory – i.e., power seating system, alternate drive control interface, or push-rim activated power assist – is needed to address the beneficiary’s mobility limitation. There must be a written report of this evaluation available on request.

**HOME ASSESSMENT:**

Prior to or at the time of delivery of a POV or PWC, the supplier or practitioner must perform an on-site evaluation of the beneficiary’s home to verify that the beneficiary can adequately maneuver the device that is provided considering physical layout, doorway width, doorway thresholds, and surfaces. There must be a written report of this evaluation available on request.

**Miscellaneous**

**Appendices**

**Utilization Guidelines**

Refer to Coverage Indications, Limitations, and/or Medical Necessity

**Sources of Information**

CMS Decision Memorandum on Mobility Assistive Equipment.

Information received from multiple sources during the comment period.

**Bibliography**

N/A

**Revision History Information**

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**COVERAGE INDICATIONS, LIMITATIONS AND/OR MEDICAL NECESSITY:**  
Revised: Standard Documentation Language to add covered prior to a beneficiary’s Medicare eligibility  
HCPCS CODES:  
Revised: HCPCS Narrative for E0986  
**DOCUMENTATION REQUIREMENTS:**  
Revised: Standard Documentation Language to add who can enter date of delivery date on the POD  
Added: Instructions for equipment retained from a prior payer and repair/replacement verbiage | • Provider Education/Guidance  
• Revisions Due To CPT/HCPCS Code Changes |

**Associated Documents**

**Attachments**

N/A

**Related Local Coverage Documents**

Article(s)

A52498 - Power Mobility Devices - Policy Article
A55426 - Standard Documentation Requirements for All Claims Submitted to DME MACs

**Related National Coverage Documents**

N/A

**Public Version(s)**

Updated on 03/29/2019 with effective dates 01/01/2019 - N/A

Updated on 04/13/2017 with effective dates 01/01/2017 - 12/31/2018

Some older versions have been archived. Please visit the MCD Archive Site to retrieve them.

**Keywords**

N/A
END OF LOCAL COVERAGE DETERMINATION
Per the Code of Federal Regulations, 42 C.F.R § 426.325, only those portions of the currently effective Local Coverage Determination (LCD) that are based on section 1862(a)(1)(A) of the Social Security Act, may be challenged through an acceptable complaint as described in 42 C.F.R § 426.400. Also, per 42 C.F.R § 426.325 items that are not reviewable, and therefore cannot be challenged, include the Policy Article. Please note the distinction of the documents when reviewing the materials.
## Contractor Information

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### Article Information

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Article Guidance

Article Text:

**NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES**

For any item to be covered by Medicare, it must 1) be eligible for a defined Medicare benefit category, 2) be reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member, and 3) meet all other applicable Medicare statutory and regulatory requirements. Information provided in this policy article relates to determinations other than those based on Social Security Act §1862(a)(1)(A) provisions (i.e. “reasonable and necessary”).

Power mobility devices are covered under the Durable Medical Equipment benefit (Social Security Act §1861(s)(6)). In order for a beneficiary’s equipment to be eligible for reimbursement the reasonable and necessary (R&N) requirements set out in the related Local Coverage Determination must be met. In addition, there are specific statutory payment policy requirements, discussed below, that also must be met.

The Medicare Prescription Drug, Improvement, and Modernization Act of 2003, added section 1834(a)(1)(E)(iv) which provides that payment may not be made for a motorized or power wheelchair unless the practitioner who has conducted the face-to-face examination him or herself writes the 7-element order. It is a statutory requirement that
all items of the 7-element order be entered specifically by and only by the practitioner who has conducted the face-to-face requirements. (See below).

For a power operated vehicle (POV) or power wheelchair (PWC) to be covered, the supplier must receive from the prescribing practitioner a written order, termed the 7-element order, containing all the elements specified in the Documentation Requirements section of the Local Coverage Determination within 45 days after completion of the practitioner’s face-to-face examination and prior to delivery of the device. (Exception: If the examination is performed during a hospital or nursing home stay, the supplier must receive the order within 45 days after discharge.) If these requirements are not met, the claim will be denied as noncovered.

If the detailed product description for the specific device is not obtained prior to delivery, payment will not be made for the item even if the documentation is subsequently obtained. If a similar item is provided by an unrelated supplier who has obtained the required documentation prior to delivery, it will be eligible for coverage.

A power mobility device may not be ordered by a podiatrist. If it is, it will be denied as noncovered.

REQUIREMENTS FOR SPECIFIC DMEPOS ITEMS PURSUANT TO 42 CFR 410.38(c) and 42 CFR 410.38(g)

42 CFR 410.38(c) requires a face-to-face evaluation and a specific written order prior to delivery for specified PMD codes.

42 CFR 410.38(g) requires a face-to-face evaluation and a specific written order prior to delivery for specified HCPCS codes. CMS provides a list of the specified codes, which is periodically updated, located [here](#).

Claims for the specified items subject to 42 CFR 410.38(c) or 42 CFR 410.38(g) that do not meet the requirements specified in the LCD-related Standard Documentation Requirements Article will be denied as statutorily noncovered – failed to meet statutory requirements.

If the supplier delivers the item prior to receipt of a written order, it will be denied as statutorily noncovered. If the written order is not obtained prior to delivery, payment will not be made for that item even if a written order is subsequently obtained. If a similar item is subsequently provided by an unrelated supplier who has obtained a written order prior to delivery, it will be eligible for coverage.

For the specified items that must meet this 42 CFR 410.38(g) requirement, the 7-element order with the NPI applied to it will fulfill both the 42 CFR 410.38(g) and the DWO prescription requirements.

FACE-TO-FACE EXAMINATION:

For a POV or PWC to be covered, the treating practitioner must conduct a face-to-face examination of the beneficiary before writing the order and the supplier must receive a written report of this examination within 45 days after completion of the face-to-face examination and prior to delivery of the device. If this requirement is not met, the claim will be denied as noncovered. (Exceptions: If this examination is performed during a hospital or nursing home stay, the supplier must receive the report of the examination within 45 days after discharge. If the POV or PWC is a replacement during the 5 year useful lifetime of an item in the same performance group that was previously covered by Medicare, a face-to-face examination is not required. Note: Replacement during an item’s useful lifetime is limited to situations involving loss or irreparable damage from a specific accident or natural disaster [e.g., fire, flood, etc.]. )

The practitioner may refer the beneficiary to a licensed/certified medical professional, such as a physical therapist...
(PT) or occupational therapist (OT), who has experience and training in mobility evaluations to perform part of the face-to-face examination. This person may have no financial relationship with the supplier. (Exception: If the supplier is owned by a hospital, PT or OT working in the inpatient or outpatient hospital setting may perform part of the face-to-face examination.)

If the beneficiary was referred before being seen by the practitioner, then once the practitioner has received and reviewed the written report of this examination, the practitioner must see the beneficiary and perform any additional examination that is needed. The report of the practitioner’s visit shall state concurrence or any disagreement with the LCMP examination. In this situation, the practitioner must provide the supplier with a copy of both examinations within 45 days after the face-to-face examination with the practitioner.

If the practitioner saw the beneficiary to begin the examination before referring the beneficiary to an LCMP, then if the practitioner sees the beneficiary again in person after receiving the report of the LCMP examination, the 45-day period begins on the date of that second practitioner visit. However, it is also acceptable for the practitioner to review the written report of the LCMP examination, to sign and date that report, and to state concurrence or any disagreement with that examination. In this situation, the practitioner must send a copy of the note from his/her initial visit to evaluate the beneficiary plus the annotated, signed, and dated copy of the LCMP examination to the supplier. The 45-day period begins when the practitioner signs and dates the LCMP examination.

For an item addressed in this policy to be covered by Medicare, a written signed and dated order must be received by the supplier prior to delivery of the item. If the supplier delivers the item prior to receipt of a written order, it will be denied as noncovered. If the written order is not obtained prior to delivery, payment will not be made for that item even if a written order is subsequently obtained. If a similar item is subsequently provided by an unrelated supplier who has obtained a written order prior to delivery, it will be eligible for coverage.

A custom motorized/power wheelchair base (K0013) must be uniquely constructed or substantially modified for a specific beneficiary according to the description and orders of the beneficiary’s treating practitioner. The beneficiary’s needs must not be able to be accommodated by any other existing PMD and accessories, including customized seating arrangements. See 42 CFR Section 414.224 and Internet-Only Manual, Publication 100-04, Medicare Claims Processing Manual, Chapter 20, Section 30.3 for more information on customized DME.

MISCELLANEOUS:

A seat elevator is a statutorily noncovered option on a power wheelchair. If a PWC with a seat elevator (K0830, K0831) is provided, it will be denied as noncovered.

If any POV or PWC is only for use outside the home, it will be denied as noncovered.

Reimbursement for the wheelchair codes includes all labor charges involved in the assembly of the wheelchair. Reimbursement also includes support services, such as delivery, set-up, and education about the use of the PMD.

Upgrades that are beneficial primarily in allowing the beneficiary to perform leisure or recreational activities are noncovered.

POLICY SPECIFIC DOCUMENTATION

In addition to policy specific documentation requirements, there are general documentation requirements that are applicable to all DMEPOS policies. These general requirements are located in the DOCUMENTATION REQUIREMENTS section of the LCD.

Refer to the LCD-related Standard Documentation Requirements article, located at the bottom of this Policy Article
under the Related Local Coverage Documents section for additional information regarding GENERAL DOCUMENTATION REQUIREMENTS and the POLICY SPECIFIC DOCUMENTATION REQUIREMENTS discussed below.

If documentation of the medical necessity for a K0013 wheelchair is requested, contractors must be able to determine that the item delivered is a customized item. Documentation must include a description of the beneficiary’s unique physical and functional characteristics that require a custom motorized/power wheelchair base. This must include a detailed description of the manufacturing of the wheelchair base, including types of materials used in custom fabricating or substantially modifying it, and the construction process and labor skills required to modify it. The record must document that the needs of the beneficiary cannot be met using another power wheelchair base that incorporates seating modifications or other options or accessories (prefabricated and/or custom). The documentation must demonstrate that the K0013 is so different from another power wheelchair base that the two items cannot be grouped together for pricing purposes.

MODIFIERS

KX, GA, GY, AND GZ MODIFIERS:

If the requirements related to a face-to-face examination have not been met, the GY modifier must be added to the codes for the power mobility device and all accessories.

If the power mobility device or push-rim activated power assist device that is provided is only needed for mobility outside the home, the GY modifier must be added to the codes for the item and all accessories.

A KX modifier may be added to the code for a power mobility device and all accessories only if one of the following conditions is met:

1. If all of the coverage criteria specified in the related LCD have been met for the product that is provided; or
2. If there is an affirmative Advance Determination of Medicare Coverage (ADMC) for the product that is provided.

If the requirements for use of the KX modifier or GY modifier are not met, the GA or GZ modifier must be added to the code. When there is an expectation of a medical necessity denial, suppliers must enter GA on the claim line if they have obtained a properly executed Advance Beneficiary Notice (ABN) or GZ if they have not obtained a valid ABN.

Claim lines billed without a KX, GA, GY, or GZ modifier will be rejected as missing information.

MISCELLANEOUS:

Programs exist within Medicare for certain power wheelchair codes addressed in the Power Mobility Devices LCD and related PA:

1. Advanced Determination of Medicare Coverage (ADMC) – This is a voluntary program that allows suppliers and beneficiaries to request prior approval and determine, in advance of submitting a claim, whether the purchase of a DME item would likely be covered. Currently it extends to certain wheelchair bases, options and accessories.

2. Condition of Payment Prior Authorization (PA) Program – This is program requires obtaining prior authorization as a condition of payment for certain DMEPOS items. This program was initially required for HCPCS codes
K0856 and K0861, in 4 states, beginning March 20, 2017. It was expanded nationwide for dates of services on or after July 17, 2017. The program was expanded, for dates of service on or after September 1, 2018, to include power mobility devices K0813 - K0829, K0835 - K0843, K0848 - K0856, and K0861.

Please refer to the Supplier Manual for further details about these programs.

**CODING GUIDELINES**

**DEFINITIONS:**

Power Mobility Device (PMD) - Base codes include both integral frame and modular construction type power wheelchairs (PWCs) and power operated vehicles (POVs).

Power Wheelchair - Chair-like battery powered mobility device for people with difficulty walking due to illness or disability, with integrated or modular seating system, electronic steering, and four or more wheel non-highway construction.

Power Operated Vehicle - Chair-like battery powered mobility device for people with difficulty walking due to illness or disability, with integrated seating system, tiller steering, and three or four-wheel non-highway construction.

Beneficiary Weight Capacity – The terms Standard Duty, Heavy Duty, etc., refer to weight capacity, not performance. For example, the term Group 3 heavy duty power wheelchair denotes that the PWC has Group 3 performance characteristics and beneficiary weight handling capacity between 301 and 450 pounds. A device is not required to carry all the weight listed in the class of devices, but must have a beneficiary weight capacity within the range to be included. For example, a PMD that has a weight capacity of 400 pounds is coded as a Heavy Duty device.

Portable - A category of devices with lightweight construction or ability to disassemble into lightweight components that allows easy placement into a vehicle for use in a distant location.

Performance Testing - Term used to denote the RESNA based test parameters used to test PMDs. The PMD is expected to meet or exceed the listed performance and durability figures for the category in which it is to be used when tested. There is no requirement to test the PMD with all possible accessories.

Test Standards - Performance and durability acceptance criteria defined by ANSI/RESNA standard testing protocols.

Crash Testing - Successful completion of WC-19 testing.

Top End Speed - Minimum speed acceptable for a given category of devices. It is to be determined by the RESNA test for maximum speed on a flat hard surface.

Range - Minimum distance acceptable for a given category of devices on a single charge of the batteries. It is to be determined by the appropriate RESNA test for range.

Obstacle Climb - Vertical height of a solid obstruction that can be climbed using the standing and/or 0.5 meter run-up RESNA test.

Dynamic Stability Incline - The minimum degree of slope at which the PMD in the most common seating and positioning configuration(s) remains stable at the required beneficiary weight capacity. If the PMD is stable at only one configuration, the PMD may have protective mechanisms that prevent climbing inclines in configurations that may be unstable.
Radius Pivot Turn – The distance required for the smallest turning radius of the PMD base. This measurement is equivalent to the “minimum turning radius” specified in the ANSI/RESNA bulletins.

PWC Basic Equipment Package - Each power wheelchair code is required to include all these items on initial issue (i.e., no separate billing/payment at the time of initial issue, unless otherwise noted). The statement that an item may be separately billed does not necessarily indicate coverage.

- Lap belt or safety belt. Shoulder harness/straps or chest straps/vest may be billed separately.
- Battery charger, single mode
- Complete set of tires and casters, any type
- Legrests. There is no separate billing/payment if fixed, swingaway, or detachable non-elevating legrests with or without calf pad are provided. Elevating legrests may be billed separately.
- Footrests/foot platform. There is no separate billing/payment if fixed, swingaway, or detachable footrests or a foot platform without angle adjustment are provided. There is no separate billing for angle adjustable footplates with Group 1 or 2 PWCs. Angle adjustable footplates may be billed separately with Group 3, 4 and 5 PWCs.
- Armrests. There is no separate billing/payment if fixed, swingaway, or detachable non-adjustable height armrests with arm pad are provided. Adjustable height armrests may be billed separately.
- Any weight specific components (braces, bars, upholstery, brackets, motors, gears, etc.) as required by beneficiary weight capacity.
- Any seat width and depth. Exception: For Group 3 and 4 PWCs with a sling/solid seat/back, the following may be billed separately:
  - For Standard Duty, seat width and/or depth greater than 20 inches;
  - For Heavy Duty, seat width and/or depth greater than 22 inches;
  - For Very Heavy Duty, seat width and/or depth greater than 24 inches;
  - For Extra Heavy Duty, no separate billing
- Any back width. Exception: For Group 3 and 4 PWCs with a sling/solid seat/back, the following may be billed separately:
  - For Standard Duty, seat back width greater than 20 inches;
  - For Heavy Duty, back width greater than 22 inches;
  - For Very Heavy Duty, back width greater than 24 inches;
  - For Extra Heavy Duty, no separate billing
- Controller and Input Device

There is no separate billing/payment if a non-expandable controller and a standard proportional joystick (integrated or remote) is provided. An expandable controller, a nonstandard joystick (i.e., nonproportional or mini, compact or short throw proportional), or other alternative control device may be billed separately.

Refer to the bundling table in the Wheelchair Options and Accessories Policy Article for a list of codes that are not
separately billable at the time of initial issue of a PWC.

POV Basic Equipment Package - Each POV is to include all these items on initial issue (i.e., no separate billing/payment at the time of initial issue):

- Battery or batteries required for operation
- Battery charger, single mode
- Weight appropriate upholstery and seating system
- Tiller steering
- Non-expandable controller with proportional response to input
- Complete set of tires
- All accessories needed for safe operation

Cross Brace Chair - A type of construction for a power wheelchair in which opposing rigid braces hinge on pivot points to allow the device to fold.

Power Options - Tilt, recline, elevating legrests, seat elevators, or standing systems that may be added to a PWC to accommodate a beneficiary’s specific need for seating assistance.

No Power Options – A category of PWCs that is incapable of accommodating a power tilt, recline, seat elevation, or standing system. If a PWC can only accept power elevating legrests, it is considered to be a No Power Option chair.

Single Power Option - A category of PWCs with the capability to accept and operate a power tilt or power recline or power standing or, for Groups 3, 4, and 5, a power seat elevation system, but not a combination power tilt and recline seating system. It may be able to accommodate power elevating legrests, seat elevator, and/or standing system in combination with a power tilt or power recline. A PMD does not have to be able to accommodate all features to qualify for this code. For example, a power wheelchair that can only accommodate a power tilt could qualify for this code.

Multiple Power Options - A category of PWCs with the capability to accept and operate a combination power tilt and recline seating system. It may also be able to accommodate power elevating legrests, a power seat elevator, and/or a power standing system. A PWC does not have to accommodate all features to qualify for this code.

Actuator – A motor that operates a specific function of a power seating system – i.e., tilt, back recline, power sliding back, elevating legrest(s), seat elevation, or standing.

Proportional Control Input Device - A device that transforms a user's drive command (a physical action initiated by the wheelchair user) into a corresponding and comparative movement, both in direction and in speed, of the wheelchair. The input device shall be considered proportional if it allows for both a non-discrete directional command and a non-discrete speed command from a single drive command movement. (Note: In the Wheelchair Options and Accessories policy, the term “interface” is used instead of “control input device”.)

Non-Proportional Control Input Device - A device that transforms a user's discrete drive command (a physical action initiated by the wheelchair user, such as activation of a switch) into perceptually discrete changes in the wheelchair's speed, direction, or both.

Alternative Control Device - A device that transforms a user’s drive commands by physical actions initiated by the
user to input control directions to a power wheelchair that replaces a standard proportional joystick. Includes mini-
proportional, compact, or short throw joysticks, head arrays, sip and puff and other types of different input control
deVICES.

Non-Expandable Controller - An electronic system that controls the speed and direction of the power wheelchair drive
mechanism. Only a standard proportional joystick (used for hand or chin control) can be used as the input device. This system may be in the form of an integral controller or a remotely placed controller. The non-expandable controller:

a. May have the ability to control up to 2 power seating actuators through the drive control (for example, seat
elevator and single actuator power elevating legrests). (Note: Control of the power seating actuators though
the Control Input Device would require the use of an additional component, E2310 or E2311.)
b. May allow for the incorporation of an attendant control.

Expandable Controller - An electronic system that is capable of accommodating one or more of the following
additional functions:

a. Proportional input devices (e.g., mini, compact, or short throw joysticks, touchpads, chin control, head control,
etc.) other than a standard proportional joystick.
b. Non-proportional input devices (e.g., sip and puff, head array, etc.)
c. Operate 3 or more powered seating actuators through the drive control. (Note: Control of the power seating
actuators though the Control Input Device would require the use of an additional component, E2310 or E2311.)

d. A separate display (i.e., for alternate control devices)
e. Other electronic devices (e.g., control of an augmentative speech device or computer through the chair’s drive
control)
f. An attendant control

Integral Control System - Non-expandable wheelchair control system where the joystick is housed in the same box
as the controller. The entire unit is located and mounted near the hand of the user. A direct electrical connection is
made from the Integral Control box to the motors and batteries through a high power wire harness.

Remotely Placed Controller - Non-expandable or expandable wheelchair control system where the joystick (or
alternative control device) and the controller box are housed in separate locations. The joystick (or alternative
control device) is connected to the controller through a low power wire harness. The separate controller connects
directly to the motors and batteries through a high power wire harness.

Codes E2310 and E2311 describe electronic components that allow the patient to control two or more of the following
motors from a single interface, e.g., proportional joystick, touchpad, or nonproportional interface:

- Power tilt
- Power recline, with or without shear reduction
- Combination power tilt and recline, with or without shear reduction
• Power leg elevation with or without articulation, power center mount elevating foot platform with or without articulating properties.

The interface includes a function selection switch that allows the patient to select the motor that is being controlled and an indicator feature to visually show which function has been selected. When the wheelchair drive function has been selected, the indicator feature may also show the direction that has been selected (forward, reverse, left, right). This indicator feature may be in a separate display box or may be integrated into the wheelchair interface. Payment for the interface code includes an allowance for fixed mounting hardware for the control box and the display box, if present.

A harness (E2313) describes all the wires, fuse boxes, fuses, circuits, switches, etc. that are required for the operation of an expandable controller (E2377). It also includes all the necessary fasteners, connectors, and mounting hardware.

Sling Seat/Back - Flexible cloth, vinyl, leather or equal material designed to serve as the support for buttocks or back of the user respectively. They may or may not have thin padding but are not intended to provide cushioning or positioning for the user.

Solid Seat/Back - Rigid metal or plastic material usually covered with cloth, vinyl, leather or equal material, with or without some padding material designed to serve as the support for the buttocks or back of the user respectively. They may or may not have thin padding but are not intended to provide cushioning or positioning for the user. PWCs with an automotive-style back and a solid seat pan are considered as a solid seat/back system, not a captains chair.

Captains chair - A one or two-piece automotive-style seat with rigid frame, cushioning material in both seat and back sections, covered in cloth, vinyl, leather or equal as upholstery, and designed to serve as a complete seating, support, and cushioning system for the user. It may have armrests that can be fixed, swingaway, or detachable. It may or may not have a headrest, either integrated or separate.

Stadium Style Seat - A one or two piece stadium-style seat with rigid frame and cushioning material in both seat and back sections, covered in cloth, vinyl, leather or equal as upholstery, and designed to serve as a complete seating, support, and cushioning system for the user. It may have armrests that can be fixed, swingaway, or detachable. It will not have a headrest. Chairs with stadium style seats are billed using the captains chair codes.

Highway Use - Mobility devices that are powered and configured to operate legally on public streets.

Push-rim activated power assist (E0986) – An option for a manual wheelchair in which sensors in specially designed wheels determines the force that is exerted by the beneficiary upon the wheel. Additional propulsive and/or braking force is then provided by motors in each wheel. E0986 is all-inclusive. All components, e.g., drive wheels, batteries, chargers, controls, mounting hardware, etc, for a manual wheel chair conversion are considered as included in 1 UOS of the code.

CODE-SPECIFIC REQUIREMENTS:

There are five PWC Groups and two POV Groups. Groups are divided based on performance. Each group of PMDs has subdivisions based on beneficiary weight capacity, seat type, portability, and/or power seating system capability.

All POVs (K0800 – K0808, K0812) must have the specified components and meet the following requirements:

• Have all components in the POV Basic Equipment Package
• Seat Width: Any width appropriate to weight group
• Seat Depth: Any depth appropriate to weight group
• Seat Height: Any height (adjustment requirements-none)
• Back Height: Any height (minimum back height requirement-none)
• Seat to Back Angle: Fixed or adjustable (adjustment requirements – none)
• Meet the following testing requirements:
  □ Fatigue test – 200,000 cycles
  □ Drop test – 6,666 cycles

Group 1 POVs (K0800 – K0802) must meet the following requirements:

• Length - less than or equal to 48 inches
• Width - less than or equal to 28 inches
• Minimum Top End Speed - 3 MPH
• Minimum Range - 5 miles
• Minimum Obstacle Climb - 20 mm
• Radius Pivot Turn - less than or equal to 54 inches
• Dynamic Stability Incline - 6 degrees

Group 2 POVs (K0806 – K0808) must meet the following requirements:

• Length - less than or equal to 48 inches
• Width - less than or equal to 28 inches
• Minimum Top End Speed - 4 MPH
• Minimum Range - 10 miles
• Minimum Obstacle Climb - 50 mm
• Radius Pivot Turn - less than or equal to 54 inches
• Dynamic Stability Incline - 7.5 degrees

The following requirements describe the configurations of power wheelchairs as they are coded by the Pricing, Data Analysis, and Coding (PDAC) contractor. Items provided to the beneficiary may include upgraded components which are substituted for the basic component and are billed separately. One example is a power seating system. When this is provided, the base code used should be that with a sling/solid seat/back. Another example is the provision of an expandable controller when the base code includes a non-expandable controller but is capable of an upgrade.

All PWCs (K0813 – K0891, K0898) must have the specified components and meet the following requirements:

• Have all components in the PWC Basic Equipment Package
- Have the seat option listed in the code descriptor
- Seat Width: Any width appropriate to weight group
- Seat Depth: Any depth appropriate to weight group
- Seat Height: Any height (adjustment requirements - none)
- Back Height: Any height (minimum back height requirement - none)
- Seat to Back Angle: Fixed or adjustable (adjustment requirements – none)
- May include semi-reclining back
- Meet the following testing requirements:
  - Fatigue test – 200,000 cycles
  - Drop test – 6,666 cycles

All Group 1 PWCs (K0813 – K0816) must have the specified components and meet the following requirements:

- Standard integrated or remote proportional joystick
- Non-expandable controller
- Incapable of upgrade to expandable controller
- Incapable of upgrade to alternative control devices
- May have crossbrace construction
- Accommodates non-powered options and seating systems (e.g., recline-only backs, manually elevating legrests) (except captains chairs)
- Length - less than or equal to 40 inches
- Width - less than or equal to 24 inches
- Minimum Top End Speed - 3 MPH
- Minimum Range - 5 miles
- Minimum Obstacle Climb - 20 mm
- Dynamic Stability Incline - 6 degrees

For Group 1 portable wheelchairs (K0813, K0814), the largest single component may not exceed 55 pounds.

All Group 2 PWCs (K0820 – K0843) must have the specified components and meet the following requirements:

- Standard integrated or remote proportional joystick
- May have crossbrace construction
- Accommodates seating and positioning items (e.g., seat and back cushions, headrests, lateral trunk supports, lateral hip supports, medial thigh supports) (except captains chairs)
- Length - less than or equal to 48 inches
- Width - less than or equal to 34 inches
- Minimum Top End Speed - 3 MPH
- Minimum Range - 7 miles
- Minimum Obstacle Climb - 40 mm
- Dynamic Stability Incline - 6 degrees

For Group 2 portable PWCs (K0820, K0821), the largest single component may not exceed 55 pounds.

Group 2 no power option PWCs (K0820 – K0829) must have the specified components and meet the following requirements:
Non-expandable controller

- Incapable upgrade to expandable controller
- Incapable of upgrade to alternative control devices
- Incapable of accommodating a power tilt, recline, seat elevation, standing system
- Accommodates non-powered options and seating systems (e.g., recline-only backs, manually elevating legrests) (except captains chairs)

Group 2 seat elevator PWCs (K0830, K0831) must have the specified components and meet the following requirements:

- Non-expandable controller
- Incapable of upgrade to expandable controller
- Incapable of upgrade to alternative control devices
- Accommodates only a power seat elevating system

Group 2 single power option PWCs (K0835 – K0840) must have the specified components and meet the following requirements:

- Non-expandable controller
- Capable of upgrade to expandable controller
- Capable of upgrade to alternative control devices
- See Single Power Option definition for seating system capability

Group 2 multiple power option PWCs (K0841 – K0843) must have the specified components and meet the following requirements:

- Non-expandable controller
• Capable of upgrade to expandable controller
• Capable of upgrade to alternative control devices
• See Multiple Power Options definition for seating system capability
• Accommodates a ventilator

All Group 3 PWCs (K0848 – K0864) must have the specified components and meet the following requirements:

• Standard integrated or remote proportional joystick
• Non-expandable controller
• Capable of upgrade to expandable controller
• Capable of upgrade to alternative control devices
• May not have crossbrace construction
• Accommodates seating and positioning items (e.g., seat and back cushions, headrests, lateral trunk supports, lateral hip supports, medial thigh supports) (except captains chairs)
• Drive wheel suspension to reduce vibration
• Length - less than or equal to 48 inches
• Width - less than or equal to 34 inches
• Minimum Top End Speed - 4.5 MPH
• Minimum Range - 12 miles
• Minimum Obstacle Climb - 60 mm
• Dynamic Stability Incline - 7.5 degrees

All Group 4 PWCs (K0868 – K0886) must have the specified components and meet the following requirements:

• Standard integrated or remote proportional joystick
• Non-expandable controller
• Capable of upgrade to expandable controller
• Capable of upgrade to alternative control devices
• May not have crossbrace construction
• Accommodates seating and positioning items (e.g., seat and back cushions, headrests, lateral trunk supports, lateral hip supports, medial thigh supports) (except captains chairs)
• Drive wheel suspension to reduce vibration
• Length - less than or equal to 48 inches
• Width - less than or equal to 34 inches
Minimum Top End Speed - 6 MPH
Minimum Range - 16 miles
Minimum Obstacle Climb - 75 mm
Dynamic Stability Incline - 9 degrees

Group 3 and 4 no power option PWCs (K0848 – K0855, K0868 – K0871) must have the specified components and meet the following requirements:

- Incapable of accommodating a power tilt, recline, seat elevation, standing system
- Accommodates non-powered options and seating systems (e.g., recline-only backs, manually elevating legrests)

Group 3 and 4 single power option PWCs (K0856 – K0860, K0877 – K0880) must have the specified components and meet the following requirements:

- See Single Power Option definition for seating system capability

Group 3 and 4 multiple power option PWCs (K0861 – K0864, K0884 – K0886) must have the specified components and meet the following requirements:

- See Multiple Power Options definition for seating system capability
- Accommodates a ventilator

All Group 5 PWCs (K0890, K0891) must have the specified components and meet the following requirements:

- Standard integrated or remote proportional joystick
- Non-expandable controller
- Capable of upgrade to expandable controller
- Capable of upgrade to alternative control devices
- Seat Width: minimum of 5 one-inch options
- Seat Depth: minimum of 3 one-inch options
- Seat Height: adjustment requirements - ≥ 3 inches
- Back Height: adjustment requirements minimum of 3 options
- Seat to Back Angle: range of adjustment - minimum of 12 degrees
- Accommodates non-powered options and seating systems
- Accommodates seating and positioning items (e.g., seat and back cushions, headrests, lateral trunk supports, lateral hip supports, medial thigh supports)
- Adjustability for growth (minimum of 3 inches for width, depth and back height adjustment)
- Special developmental capability (i.e., seat to floor, standing, etc.)
- Drive wheel suspension to reduce vibration
- Length - less than or equal to 48 inches
- Width - less than or equal to 34 inches
- Minimum Top End Speed - 4 MPH
- Minimum Range - 12 miles
- Minimum Obstacle Climb - 60 mm
- Dynamic Stability Incline - 9 degrees
- Crash testing - Passed

Group 5 single power option PWC (K0890) must have the specified components and meet the following requirements:

- See Single Power Option definition for seating system capability

Group 5 multiple power option PWC (K0891) must have the specified components and meet the following requirements:

- See Multiple Power Options definition for seating system capability
- Accommodates a ventilator

MISCELLANEOUS:

The only products that may be billed using codes K0800-K0898 are those products for which a written coding verification determination has been made by the Pricing, Data Analysis, and Coding (PDAC) contractor. Information concerning the documentation that must be submitted to the PDAC for a Coding Verification Review can be found on the PDAC web site or by contacting the PDAC. A Product Classification List with devices which have received a coding verification determination can be found on the PDAC web site. Note that code K0013 is not included in the list of products requiring Coding Verification Review.

Manufacturers and suppliers should refer to the PDAC web site or contact the PDAC for information concerning testing requirements.

If a power mobility device has not received a written coding verification determination from the PDAC or if the PDAC determines that the product does not meet the requirements of any code, it must be billed with code K0899.

Suppliers should contact the Pricing, Data Analysis, and Coding (PDAC) contractor for guidance on the correct coding of these items.

Coding Information
Bill Type Codes:

Contractors may specify Bill Types to help providers identify those Bill Types typically used to report this service. Absence of a Bill Type does not guarantee that the policy does not apply to that Bill Type. Complete absence of all Bill Types indicates that coverage is not influenced by Bill Type and the policy should be assumed to apply equally to all claims.

N/A

Revenue Codes:

Contractors may specify Revenue Codes to help providers identify those Revenue Codes typically used to report this service. In most instances Revenue Codes are purely advisory. Unless specified in the policy, services reported under other Revenue Codes are equally subject to this coverage determination. Complete absence of all Revenue Codes indicates that coverage is not influenced by Revenue Code and the policy should be assumed to apply equally to all Revenue Codes.

N/A

CPT/HCPCS Codes

N/A

ICD-10 Codes that are Covered

N/A

ICD-10 Codes that are Not Covered

N/A

Revision History Information

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| 01/01/2019            | R8                      | Revision Effective Date: 01/01/2019  
POLICY SPECIFIC DOCUMENTATION:  
CODING GUIDELINES:  
Added: Coding guidelines for power wheelchair electronics |
<p>| 04/04/2019            |                         | 04/04/2019: At this time 21st Century Cures Act applies to new and revised |</p>
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<td>09/01/2018</td>
<td>R7</td>
<td>LCDs which require comment and notice. This revision is to an article that is not a local coverage determination.</td>
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| 09/01/2018            | R7                      | Revision Effective Date: 09/01/2018  
POLICY SPECIFIC DOCUMENTATION:  
Revised: ADMC, Prior Authorization of Power Mobility Devices (PMD) Demonstration and Condition of Payment Prior Authorization Program information  
09/13/2018: At this time 21st Century Cures Act applies to new and revised LCDs that restrict coverage, which require comment and notice. This revision is to an article that is not a local coverage determination. |
| 01/01/2017            | R6                      | Revision Effective Date: 01/01/2018  
NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:  
Added: 42 CFR(c) and (g) language, previously in Policy Specific Documentation section  
POLICY SPECIFIC DOCUMENTATION:  
Revised: ADMC eligible base codes to conform to the Condition of Payment PA Program  
Added: Prior Authorization of Power Mobility Devices (PMD) Demonstration and Condition of Payment PA Program information  
04/12/2018: At this time 21st Century Cures Act applies to new and revised LCDs that restrict coverage, which require comment and notice. This revision is to an article that is not a local coverage determination. |
| 01/01/2017            | R5                      | Revision Effective Date: 01/01/2017  
POLICY SPECIFIC DOCUMENTATION REQUIREMENTS:  
42 CFR 410.38(c) and 42 CFR 410.38(g) language, K0013 billing instructions, Modifier instructions and ADMC eligible codes  
RELATED LOCAL COVERAGE DOCUMENTS:  
Added: LCD-related Standard Documentation Requirements Language Article |
| 07/01/2016            | R4                      | Revision Effective Date: 07/01/2016  
NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:  
Revised Standard Language to add Statutory Prescription (Order) Requirements, revised Face to Face and ACA requirements - Effective 04/28/2016 |
<p>| 07/01/2016            | R3                      | Effective July 1, 2016 oversight for DME MAC Articles is the responsibility of CGS Administrators, LLC 18003 and 17013 and Noridian Healthcare Solutions, LLC 19003 and 16013. No other changes have been made to the Articles. |</p>
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| 10/01/2015            | R2                     | **Revision Effective Date: 10/01/2015**  
NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:  
Revised: HCPCS Narrative for E0986 and updated standard language documentation |
| 10/01/2015            | R1                     | **Revision Effective Date: 10/01/2014**  
NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:  
Clarification: The face to face treating physician and prescribing physician requirements under ACA 6407 (Requirements effective 07/01/2013)  
CODING GUIDELINES:  
Clarification: E0986 is all inclusive |

**Associated Documents**

**Related Local Coverage Document(s)**
- Article(s)
- [A55426 - Standard Documentation Requirements for All Claims Submitted to DME MACs](#)
- LCD(s)
- [L33789 - Power Mobility Devices](#)

**Related National Coverage Document(s)**
- N/A

**Statutory Requirements URL(s)**
- N/A

**Rules and Regulations URL(s)**
- N/A

**CMS Manual Explanations URL(s)**
- N/A

**Other URL(s)**
- N/A

**Public Version(s)**
- Updated on 03/29/2019 with effective dates 01/01/2019 - N/A
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- Updated on 04/04/2018 with effective dates 01/01/2017 - N/A
- Updated on 04/13/2017 with effective dates 01/01/2017 - N/A

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**Keywords**
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