## Contractor Information

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<th>Contractor Name</th>
<th>Contract Type</th>
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<tr>
<td>CGS Administrators, LLC</td>
<td>DME MAC</td>
<td>17013 - DME MAC</td>
<td>J-B ILLINOIS, INDIANA, KENTUCKY, MICHIGAN, MINNESOTA, OHIO, WISCONSIN, ALABAMA, ARKANSAS, COLORADO, FLORIDA, GEORGIA, LOUISIANA, MISSISSIPPI, NORTH CAROLINA, NEW MEXICO, OKLAHOMA, PUERTO RICO, SOUTH CAROLINA, TENNESSEE, TEXAS, VIRGINIA, VIRGIN ISLANDS, WEST VIRGINIA, CONNECTICUT, DISTRICT OF COLUMBIA, DELAWARE, MASSACHUSETTS, MARYLAND, MAINE, NEW HAMPSHIRE, NEW JERSEY, NEW YORK - ENTIRE STATE, PENNSYLVANIA, RHODE ISLAND, VERMONT, ALASKA, AMERICAN SAMOA, ARIZONA, CALIFORNIA - ENTIRE STATE, GUAM, HAWAII, IOWA, IDAHO, KANSAS, MISSOURI - ENTIRE STATE, MONTANA, NORTH DAKOTA, NEBRASKA, NEVADA, OREGON, SOUTH DAKOTA</td>
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Contractor Name

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Article Information

General Information

Article ID
A52505

Original Article Effective Date
10/01/2015

Original ICD-9 Article ID
A47225
A17985
A17918
A17265

Revision Effective Date
01/01/2018

Revision Ending Date
N/A

Retirement Date
N/A

Article Title
Wheelchair Seating - Policy Article

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Article Guidance

Article Text:

NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:

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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”).

There is no separate payment for a solid insert (E0992) (see definition in Coding Guidelines) that is used with a seat or back cushion because a solid base is included in the allowance for a wheelchair seat or back cushion.

There is no separate payment for mounting hardware for a seat or back cushion.

There is no separate payment for a wheelchair seat or back cushion when it is used with a rollabout chair (E1031).

**REQUIREMENTS FOR SPECIFIC DMEPOS ITEMS PERSUANT TO 42 CFR 410.38(g)**

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(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.

**POLICY SPECIFIC DOCUMENTATION REQUIREMENTS**

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.

**MODIFIERS**

**KX MODIFIER:**

For a skin protection seat cushion (E2603, E2604, E2622, E2623), a KX modifier must be added to the code only if either criterion (a), (b), or (c) is met:

a. If there is a past history of or current pressure ulcer in the area of contact with the seating surface; or
b. If there is absent or impaired sensation in the area of contact with the seating surface due to one of the diagnoses listed as a covered diagnosis in the related LCD; or

KX modifier must be added to the code only if either criterion (a), (b), (c), or (d) is met:

a. If there is a past history of or current pressure ulcer in the area of contact with the seating surface; or
b. If there is absent or impaired sensation in the area of contact with the seating surface due to one of the diagnoses listed as a covered diagnosis for skin protection cushions (see diagnosis codes that support medical necessity section in the related LCD); or
If there is an inability to carry out a functional weight shift due one of the diagnoses listed as a covered diagnosis for skin protection cushions (see diagnosis codes that support medical necessity section in the related LCD); and

If the beneficiary has significant postural asymmetries due to one of the diagnoses listed as a covered diagnosis for positioning cushions (see diagnosis codes that support medical necessity section in the related LCD).

For a custom fabricated seat or back cushion (E2609, E2617), a KX modifier must be added to the code only if criterion (a) is met and criterion (b), (c), or (d) is met:

a. For E2609 or E2617, there is a comprehensive written evaluation by a licensed/certified medical professional, such as a PT or OT (who has no financial relationship with the supplier) which explains why a prefabricated seating system is not sufficient to meet the beneficiary’s seating and positioning needs; and
b. For E2609, there is a past history of or current pressure ulcer in the area of contact with the seating surface; or
c. For E2609, there is absent or impaired sensation in the area of contact with the seating surface or an inability to carry out a functional weight shift due to one of the diagnoses listed as a covered diagnosis for skin protection cushions (see diagnosis codes that support medical necessity section in the related LCD); or

d. For E2609 or E2617, the beneficiary has significant postural asymmetries due to one of the diagnoses listed as a covered diagnosis for positioning cushions (see diagnosis codes that support medical necessity section in the related LCD).

In addition to meeting the specific requirements listed above, for all seat and back cushions and positioning accessories, the KX modifier must be added to the code only if the item is being used with a wheelchair that meets coverage criteria specified in the Manual Wheelchair Bases or Power Mobility Devices LCD.

GA, GY, AND GZ MODIFIERS:

For a cushion or positioning accessory that is used with a power mobility device, if the requirements related to a 7-element order and face-to-face examination in the Power Mobility Devices Policy Article have not been met, the GY modifier must be added to the codes for all items.

For items provided with a manual wheelchair or power mobility device, if it is only needed for mobility outside the home, the GY modifier must be added to the codes for all items.

In all of the situations above describing use of the KX modifier, if all of the specific coverage criteria have not been met or if the wheelchair that it is being used with does not meet the coverage criteria in the Manual Wheelchair Bases or Power Mobility Devices LCD, the GA or GZ modifier must be added to a claim line for the seat or back cushion or positioning accessory.

When there is an expectation of a medical necessity denial, suppliers must enter the GA modifier on the claim line if they have obtained a properly executed Advance Beneficiary Notice (ABN) or the GZ modifier if they have not obtained a valid ABN.

If the GY modifier is used, the KX, GA, and GZ modifiers should not be used.

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

Miscellaneous

When billing for a custom fabricated cushion (E2609, E2617), the claim must include the manufacturer and model name/ number of the product (if applicable), or if not, a detailed description of the product that was provided.

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

CODING GUIDELINES:

The following definitions of seat cushions include results of simulation testing or human subject testing. See the Testing Methodologies section for technical information about the required testing.

A general use seat cushion (E2601, E2602) is a prefabricated cushion, which has the following characteristics:

1. It has the following minimum performance characteristics:
a. Simulation tests demonstrate a loaded contour depth of at least 25mm with an overload deflection of at least 5 mm, or
b. Human subject tests demonstrate an average peak pressure index that is less than 125% of that of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and

2. Following testing simulating 12 months of use:
   a. Simulation tests demonstrate an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 125% of those of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and

3. It has a removable vapor permeable or waterproof cover or it has a waterproof surface; and

4. The cushion and cover meet the minimum standards of the California Bulletin 117 or 133 for flame resistance; and

5. It has a permanent label indicating the model and the manufacturer; and

6. It has a warranty that provides for repair or full replacement if manufacturing defects are identified or the surface does not remain intact due to normal wear within 12 months.

A nonadjustable skin protection seat cushion (E2603, E2604) is a prefabricated cushion, which has the following characteristics:

1. It has the following minimum performance characteristics:
   a. Simulation tests demonstrate a loaded contour depth of at least 40 mm with an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 85% of that of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and

2. Following testing simulating 18 months of use:
   a. Simulation tests demonstrate an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 85% of those of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and

3. It has a removable vapor permeable or waterproof cover or it has a waterproof surface; and

4. The cushion and cover meet the minimum standards of the California Bulletin 117 or 133 for flame resistance; and

5. It has a permanent label indicating the model and the manufacturer; and

6. It has a warranty that provides for repair or full replacement if manufacturing defects are identified or the surface does not remain intact due to normal wear within 18 months.

An adjustable skin protection seat cushion (E2622, E2623) has all the characteristics of a E2603 or E2604 cushion and is determined to be adjustable by the PDAC.

A positioning seat cushion (E2605, E2606) is a prefabricated cushion that has the following characteristics:

1. It has the minimum structural features described in (a) or (b):
   a. The feature must be at least 25 mm in height in the pre-loaded state. It has two or more of the following:
      i. A pre-ischial bar or ridge which is placed anterior to the ischial tuberosities and prevents forward migration of the pelvis,
      ii. Two lateral pelvic supports which are placed posterior to the trochanters and are intended to maintain the pelvis in a centered position in the seat and/or provide lateral stability to the pelvis,
      iii. A medial thigh support which is placed in contact with the adductor region of the thigh and provides the prescribed amount of abduction and prevents adduction of the thighs,
      iv. Two lateral thigh supports which are placed anterior to the trochanters and provide lateral stability to the lower extremities and prevent unwanted abduction of the thighs.
   b. It has two or more air compartments located in areas which address postural asymmetries, each of which must have a cell height of at least 50 mm, must allow the user to add or remove air, and must have a valve which retains the desired air volume; and

2. It has the following minimum performance characteristics:
   a. Simulation tests demonstrate a loaded contour depth of at least 25mm with an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 125% of that of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and

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3. Following testing simulating 18 months of use:
   a. Simulation tests demonstrate an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 125% of those of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and
4. It has a removable vapor permeable or waterproof cover or it has a waterproof surface; and
5. The cushion and cover meet the minimum standards of the California Bulletin 117 or 133 for flame resistance; and
6. It has a permanent label indicating the model and the manufacturer; and
7. It has a warranty that provides for repair or full replacement if manufacturing defects are identified or the surface does not remain intact due to normal wear within 18 months.

A positioning cushion may have materials or components that can be added or removed to help address orthopedic deformities or postural asymmetries. This definition includes cushions which have a planar surface but have positioning features within the cushion which are made of a firmer material than the surface material.

A nonadjustable skin protection and positioning seat cushion (E2607, E2608) is a prefabricated cushion which has the following characteristics:

1. It has the minimum structural features described in (a) or (b):
   a. The feature must be at least 25 mm in height in the pre-loaded state. It has two or more of the following:
      i. A pre-ischial bar or ridge which is placed anterior to the ischial tuberosities and prevents forward migration of the pelvis,
      ii. Two lateral pelvic supports which are placed posterior to the trochanters and are intended to maintain the pelvis in a centered position in the seat and/or provide lateral stability to the pelvis,
      iii. A medial thigh support which is placed in contact with the adductor region of the thigh and provides the prescribed amount of abduction and prevents adduction of the thighs,
      iv. Two lateral thigh supports which are placed anterior to the trochanters and provide lateral stability to the lower extremities and prevent unwanted abduction of the thighs.
   b. It has two or more air compartments located in areas which address postural asymmetries, each of which must have a cell height of at least 50 mm, must allow the user to add or remove air, and must have a valve which retains the desired air volume; and
2. It has the following minimum performance characteristics:
   a. Simulation tests demonstrate a loaded contour depth of at least 40mm with an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 85% of that of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and
3. Following testing simulating 18 months of use:
   a. Simulation tests demonstrate an overload deflection of at least 5 mm, or
   b. Human subject tests demonstrate an average peak pressure index that is less than 85% of those of a standard reference cushion within the area of the ischial tuberosities and sacrum/coccyx; and
4. It has a removable vapor permeable or waterproof cover or it has a waterproof surface; and
5. The cushion and cover meet the minimum standards of the California Bulletin 117 or 133 for flame resistance; and
6. It has a permanent label indicating the model and the manufacturer; and
7. It has a warranty that provides for repair or full replacement if manufacturing defects are identified or the surface does not remain intact due to normal wear within 18 months.

This definition includes cushions which have a planar surface but have positioning features within the cushion which are made of a firmer material than the surface material.

An adjustable skin protection and positioning seat cushion (E2624, E2625) has all the characteristics of a E2607 or E2608 cushion and is determined to be adjustable by the PDAC. The adjustability feature only relates to the skin protection properties of the cushion.

Wheelchair cushions containing a fluid medium (air, gas, liquid, or gel) that have the capability for the immersion characteristics of the cushion to be altered by addition or removal of fluid will be considered adjustable. The adjustment may be in the manner of direct addition or removal of the fluid (e.g. add or remove air) or indirectly by addition or removal of packets of fluid.

Adjustment applies to the skin protection portion of the cushion's function only.
All cushions are considered to be adjustable up to the point of delivery to the beneficiary. Fitting of the cushion to the individual beneficiary may involve various forms of adjustment. Adjustable as applied here, requires that the procedure is capable of being performed by the beneficiary or caregiver using items supplied at the time of initial issue of the device in response to the beneficiary’s need for more or less skin protection because of weight loss or gain or muscle tone changes.

A general use back cushion (E2611, E2612) is a prefabricated cushion, which has the following characteristics:

1. It is planar or contoured; and
2. It has a removable vapor permeable or waterproof cover or it has a waterproof surface; and
3. The cushion and cover meet the minimum standards of the California Bulletin 117 or 133 for flame resistance; and
4. It has a permanent label indicating the model and the manufacturer; and
5. It has a warranty that provides for repair or full replacement if manufacturing defects are identified or the surface does not remain intact due to normal wear within 12 months.

A positioning back cushion (E2613-E2616, E2620, E2621) is a prefabricated cushion which has the following characteristics:

1. For codes E2613-E2616, there is at least 25 mm of posterior contour in the pre-loaded state. A posterior contour is a backward curve measured from a horizontal line in the midline of the cushion; and
2. For posterior-lateral cushions (E2615, E2616) and for planar cushions with lateral supports (E2620, E2621), there is at least 75 mm of lateral contour in the pre-loaded state. A lateral contour is a backward curve measured from a horizontal line connecting the lateral extensions of the cushion; and
3. For posterior pelvic cushions (E2613, E2614), there is mounting hardware that is adjustable for horizontal position, depth, and angle; and
4. It has a removable vapor permeable or waterproof cover or it has a waterproof surface; and
5. The cushion and cover meet the minimum standards of the California Bulletin 117 or 133 for flame resistance; and
6. It has a permanent label indicating the model and the manufacturer; and
7. It has a warranty that provides for repair or full replacement if manufacturing defects are identified or the surface does not remain intact due to normal wear within 18 months.

Included in this definition are cushions which have a planar surface but have positioning features within the cushion which are made of a firmer material than the surface material.

A positioning back cushion may have materials or components that may be added or removed to help address orthopedic deformities or postural asymmetries.

A custom fabricated seat cushion (E2609) and a custom fabricated back cushion (E2617) are cushions that are individually made for a specific beneficiary starting with basic materials including:

a. liquid foam or a block of foam and
b. sheets of fabric or liquid coating material.

The cushion must be fabricated using molded-to-beneficiary-model technique, direct molded-to-beneficiary technique, CAD-CAM technology, or detailed measurements of the beneficiary used to create a configured cushion. The cushion must have structural features that significantly exceed the minimum requirements for a seat or back positioning cushion. The cushion must have a removable vapor permeable or waterproof cover or it must have a waterproof surface. A custom fabricated cushion may include certain prefabricated components (e.g., gel or multi-cellular air inserts); these components must not be billed separately. If a custom fabricated seat and back are integrated into a one-piece cushion, code as E2609 plus E2617.

If foam-in-place or other material is used to fit a substantially prefabricated cushion to an individual beneficiary, the cushion must be billed as a prefabricated cushion, not custom fabricated.

A powered wheelchair seat cushion (E2610) is a battery-powered, prefabricated cushion in which an air pump provides either sequential inflation or deflation of the air cells or a low interface pressure throughout the cushion. One type of powered seat cushion is an alternating pressure cushion.
Pediatric seating system codes E2291-E2294 may only be billed with pediatric wheelchair base codes.

A headrest extension (E0966) is a sling support for the head. Code E0955 describes any type of cushioned headrest.

A headrest (E0955) describes any type of cushioned headrest which may contain one or more cushions to position the head and fixed mounting hardware.

Lateral positioning items are used to provide lateral thigh or knee support (E0953) or lateral trunk or hip support (E0956). These are provided in a variety of shapes and sizes to suit the needs of the user.

The code for a seat or back cushion includes any rigid or semi-rigid base or posterior panel, respectively, that is an integral part of the cushion.

A solid insert is a separate rigid piece of wood or plastic which is inserted in the cover of a cushion to provide additional support. If a supplier chooses to bill separately for a solid insert used with a seat cushion use code E0992 whether it is a manual or a power wheelchair. Code A9900 must be used for a solid insert used with a back cushion.

A solid support base for a seat cushion is a rigid piece of plastic or other material which is attached with hardware to the seat frame of a wheelchair in place of a sling seat. A cushion is placed on top of the support base. Use code E2231 for a solid support base that is used with a manual wheelchair. A solid support base is included in the allowance for the power wheelchair codes. There should be no separate billing with power wheelchairs.

If a supplier chooses to bill separately for mounting hardware, either nonadjustable or adjustable, for a seat or back cushion or solid support base, code A9900 must be used.

The only products which may be billed using codes E2601-E2608, E2611-E2616, E2620, E2621, and E2622-E2625 and the only brand name products that may be billed using codes E2609 or E2617 are those products for which a written coding verification 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 Request can be found on the PDAC web site or by contacting the PDAC. A Product Classification List with products which have received a coding verification can be found on the PDAC web site.

If a non-powered, prefabricated seat cushion, a prefabricated back cushion, or a brand name custom fabricated seat or back cushion has not received a written coding verification from the PDAC or if it is determined that the cushion does not meet the criteria for the code, it must be billed with code K0669.

Pediatric size positioning accessories are billed with the codes described in this policy. Codes E1025-E1027 (lateral thoracic and lateral/anterior supports) are invalid for claim submission.

Code E1028 (swingaway or removable mounting hardware upgrade) may be billed in addition to codes E0953, E0955-E0957. It must not be billed in addition to code E0960. It must not be used for mounting hardware related to a wheelchair seat cushion or back cushion code.

Wheelchair seat and back cushion codes are all-inclusive. Use of HCPCS code K0108 or any other HCPCS code to separately bill for added components such as the foam blocks, gel packs, air cells, or equivalent material is incorrect coding.

**TESTING METHODOLOGY**

There are two testing methods that may be used to document wheelchair seat cushion criteria: the simulation method and the human subject method. Simulation tests are used to measure loaded contour depth and bottoming out. Human subject tests are used to measure peak interface pressure.

**Simulation Test**

Simulation tests measure loaded contour depth and bottoming out. They use standardized models of the human buttocks known as cushion-loading indenters (CLIs). There are two CLIs that are used for simulation testing, a 25 mm CLI and a 40 mm CLI. Specific design features of acceptable CLIs can be found on the PDAC web site.

Test method for determining 25 mm and 40 mm of contour depth:

1. Place the test cushion on a flat, horizontal surface. Cushions with curved bases must be stable during contour measurement testing.
2. Align the CLI so that it is centered from the sides of the cushion and so that the ischial tuberosities of the models are 11-15 cm from the rear edge of the cushion. The ischial tuberosity portion of the CLI should be aligned with the analogous portion of the test cushion.

3. Load the CLI to 140 Newtons (31 pounds) & wait 5 minutes.

4. Contact of the lateral buttons with the cushion indicates that the cushion has contoured to 25 or 40 mm depending on the CLI used - i.e., that it has passed the test for that trial.

5. Repeat the test two times waiting 5 minutes between trials.

A cushion must pass the respective contour test during all trials to meet the minimum criteria specified in the cushion definition section.

Overload test method for measuring bottoming out:

1. Record the height of the CLI from the horizontal surface at the end of the loaded contour depth test described above.
2. Add 47 Newtons (10 pounds) to the CLI and record the height from the horizontal surface after 1 minute.
3. Subtract the height at overload (#2) from the height at standard load (#1).
4. Round the value in #3 to the nearest 5mm.
5. Remove the overload weight and repeat the test twice, waiting 5 minutes between tests and measuring the height in #1 and #2 each time.
6. Determine the median of the three values recorded in #4. This is the "overload deflection".

If the overload deflection is greater than or equal to 5mm, then the cushion is determined not to have bottomed out during the test.

Simulated use testing:

There must be simulation of 12 or 18 months of use of the cushion (depending on the cushion type - see Definitions section). Following simulated use, the measurements for loaded contour depth and overload as described above must be repeated.

Test report:

There must be a report of the tests which includes:

1. The name and address of the facility performing the tests and the date(s) of the tests; and
2. The manufacturer and brand name/number of the test cushion; and
3. The weight of the cushion to the nearest 250 gm; and
4. The width and length of the cushion; and
5. The temperature and relative humidity of the room where the tests are conducted; and
6. Identification of which CLI was used (25 mm or 40mm); and
7. The results of the three loaded contour depth tests and the overload deflection test prior to simulated used testing; and
8. A description of the method used to simulate cushion use; and
9. A statement specifying the number of months of use that were simulated; and
10. Measurements as described in #7 obtained after simulated use testing; and
11. A statement attesting that the testing methodology described in this policy was followed; and
12. The printed name and signature of the person performing or supervising the tests and the signature date.

Human Subject Tests

The ability to demonstrate that there is an important reduction in interface pressure in comparison with a standard reference cushion when tested with human subjects is the basis for this approach. Human subject tests must be performed by an entity that has received human subject testing approval from an Institutional Review Board approved by the US Department of Health and Human Services. Ten (10) wheelchair users must be studied, at least five of which must be clinically insensate on the body surface contacting the cushion.

Interface pressure measurements are taken with each subject seated on the cushion being tested as well as on a standardized reference cushion (see below). The measurements are obtained with a transducer placed on top of the cushion. Subjects must be seated on the cushion and interface pressure transducer for at least 60 seconds before data is collected. The subject should be positioned in their typical posture as determined by query and
The standard reference cushion must be an uncovered 75 mm (± 5mm) thick high resiliency foam with a rated 25% indentation force deflection (IFD) equal to 45 pounds (density range of 2.6-2.9 pounds/cubic ft and IFD range of 40-49 pounds).

There must be a report of the tests which includes:

1. The name and address of the facility performing the tests and the date(s) of the tests; and
2. The manufacturer and brand name/number of the test cushion; and
3. Information about the interface pressure measurement device utilized:
   a. Manufacturer and brand name
   b. Date of most recent calibration
   c. Percent error of measurement at 50 and 100 mm Hg pressure; and
4. Actual 25% IFD and density of the reference cushion (obtained from the foam manufacturer or supplier) and actual thickness of the reference cushion; and
5. Information on each subject (coding subjects to preserve confidentiality) including:
   a. Age
   b. Height
   c. Weight
   d. Disability
   e. Buttocks sensation status; and
6. Interface pressure measurements for each subject on the test cushion and on the reference cushion:
   a. If the transducer covers the entire seating area, the entire map showing the pressure in each cell must be submitted. The anatomical locations (as determined by palpation) of the right and left ischial tuberosities and the sacrum/coccyx must be identified on each map. (Data can be submitted as a hard copy map or utilizing the device software.) or,
   b. If the transducer only covers a portion of the seat surface, measurements must be taken at the following three locations (as determined by palpation): right and left ischial tuberosities and sacrum/coccyx. The report must identify the anatomical location of each set of measurements. The values for the three locations are considered a single test; and
7. The Peak Pressure Index (PPI) for each subject on the test cushion and on the reference cushion. The PPI is determined as follows:
   a. For each test, identify the cell in the sacro-ischial zone with the highest pressure; 
   b. Determine the greatest sum of pressures in the identified cell and the adjacent cells in a 9-10 square centimeter area. If there are multiple cells with the same "highest pressure", consider all of them in the determination of the "greatest sum". [Note: A 3 cm by 3 cm square or a 3.5 cm diameter circular area are examples of a 9-10 sq cm area. For example, if using an interface pressure sensing array with a cell size of 1 sq cm, 9 cells (a 3 by 3 array) are used and if using a sensing array with a cell size of 2.5 sq cm, 4 cells (a 2 by 2 array) are used.];
   c. For each test, calculate the average of the cells with the greatest sum of pressures;
   d. Calculate the average of the results obtained in step (c) for the 3 tests on the test cushion and the 3 tests on the reference cushion. These values are the PPIs for the subject on each cushion; and
8. A statement attesting that the testing methodology described in this policy was followed; and
9. The printed name and signature of the person performing or supervising the tests and the signature date.

To determine if the minimum performance characteristics specified in the Definitions section for a particular type of cushion have been met, calculate the average PPI for the 10 subjects on the test cushion and the average PPI for the 10 subjects on the reference cushion. Divide the average PPI on the test cushion by the average PPI on the reference cushion and multiply the value by 100 to give the percentage comparison of Peak Pressure Indexes. If the comparative pressures are less than the specified values (125% or 85% depending on the cushion), then the minimum performance characteristics with respect to pressure have been met.

**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 article does not apply to that Bill Type. Complete absence of all Bill Types indicates that coverage is not influenced by Bill Type and the article 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 article, 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 article should be assumed to apply equally to all Revenue Codes.

N/A

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### Revision History Information

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| 01/01/2018            | R8                      | Revision Effective Date: 01/01/2018  
CODING GUIDELINES:  
Revised: Positioning cushion language placement for clarification  
Added: Wheelchair seat and back cushion codes are all-inclusive  
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.

| 01/01/2018 | R7 | POLICY SPECIFIC DOCUMENTATION REQUIREMENTS  
Added: E0953 to modifier instructions  
12/21/2017: 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.  
Revision Effective Date: 01/01/2017  
NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:  
Added: 42 CFR 410.38(g) language, previously in Policy Specific Documentation Requirements section |

CODING GUIDELINES:
Revised: Coding guidelines for E0956 due to a narrative description error

POLICY SPECIFIC DOCUMENTATION REQUIREMENTS:
Added: 42 CFR 410.38(g) and Modifiers requirements

CODING GUIDELINES:
Changed: The third bullet for one characteristic of a positioning back cushion (E2613-E2616, E2620, E2621) from “vertical” to “horizontal”.
Revised: Description of headrest (E0955) and fixed mounting hardware
Added: Coding guideline for E0956 to specify that these items may be used on trunk, hip, thigh, and knee

RELATED LOCAL COVERAGE DOCUMENTS:
Added: LCD-related Standard Documentation Requirements Language Article

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.

NON-MEDICAL NECESSITY COVERAGE AND PAYMENT RULES:
Removed: Start date verbiage from Prescription Requirements

Revised: Face-to-Face Requirements for treating practitioner

Related Local Coverage Document(s) Article(s) A55426 - Standard Documentation Requirements for All Claims Submitted to DME MACs LCD(s) L33312 - Wheelchair Seating

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 04/04/2018 with effective dates 01/01/2018 - N/A Updated on 12/13/2017 with effective dates 01/01/2018 - N/A Updated on 08/17/2017 with effective dates 01/01/2017 - N/A Updated on 03/24/2017 with effective dates 01/01/2017 - N/A Updated on 03/17/2017 with effective dates 01/01/2017 - N/A

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Keywords

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