

Durable Medical Equipment, Orthotics, Medical Supplies, and Repairs/Replacements (for Louisiana Only)

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[➔ Instructions for Use](#)

Certain Content mandated by Louisiana Department of Health

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Application

This Medical Policy only applies to the state of Louisiana. Portions of the coverage rationale contained in this policy represent Louisiana Medicaid coverage policy and is set forth below in accordance with state requirements.

Coverage Rationale

When determining medical necessity, clinical guidelines will be applied in the following order:

1. Federal, state, and contractual requirements
2. UnitedHealthcare Community Plan Medical Policy
3. InterQual® CP Durable Medical Equipment
4. InterQual® Medicare Durable Medical Equipment
5. CMS DME MAC

Durable Medical Equipment (DME), related supplies, and orthotics are Medically Necessary when:

- Consistent with the state definition of DME and/or Orthotic; and
- The item(s) meets the plans Medically Necessary definition (refer to the federal, state, or contractual requirements); and
- Ordered by a physician, or ordered by a nurse practitioner, clinical nurse specialist, or physician assistant acting within the scope of practice under state law; and
- The item is not otherwise excluded from coverage

Electric Breast Pump

An electric breast pump is a mechanical device powered by batteries or electricity that nursing mothers use to extract milk from their breasts. Medicaid considers personal-use, double, electric breast pumps a coverable item for nursing mothers. A new breast pump is covered for every delivery.

Note: Single, manual, and hospital-grade breast pumps are not covered items under Louisiana Medicaid.

Equipment Criteria

Electric breast pumps dispensed to Medicaid beneficiaries must meet, at a minimum, the below criteria:

- Have an adjustable suction pressure rate with either written instructions or an automatic mechanism to prevent a suction greater than 250 mm Hg;
- Be adaptable for simultaneous pumping of both breasts (double collection);
- Automatically cycle with an adjustable variable cycling rate, typically 30 to 60 or more cycles per minute;
- Include a battery option and adapter to be used as an alternate power source when electricity is not immediately available;
- Breast shields (flanges) that are adjustable and flexible, or flanges that are available in several different sizes if rigid, including larger sizes;
- All accessories necessary for pumping two breasts simultaneously for electric pumps;
- At least two collection bottles with spill-proof standard size caps, which are bisphenol-A (BPA) and DEHP-free; and
- Accessories and supplies must be compatible with the pump provided; materials must be of durable quality for withstanding repeated boiling, washing, and pumping use

Replacement Criteria

Medicaid will allow replacement of a breast pump older than three years and after expiration of manufacturer's warranty. Replacement and warranty are subject to policy in the Section 18.2 of the provider manual.

Electric Breast Pump Supplies

Electric breast pump supplies will be available to the nursing mother once every 180 days. DME providers must obtain a prior authorization for replacement supplies. The prior authorization request must include a prescription and baby's date of birth.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Human Milk Storage Bags

Human milk storage bags are designed to safely store and protect expressed human milk for feeding a child. Medicaid covers 100 human milk storage bags per month for lactating beneficiaries. The Medicaid reimbursement rate on file covers a one month supply of storage bags.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Enteral Infusion Pump

A standard enteral infusion pump will be approved only with documented evidence that the pump is medically necessary and that syringe or gravity feedings are not satisfactory due to complications such as aspiration, diarrhea, dumping syndrome, etc. Medicaid will pay for the rental of a standard enteral infusion pump and accessories.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Orthotic Devices

Orthotic devices include leg braces, neck braces, knee braces and supports, spinal supports, splints, brace attachments and repairs. The request for approval should include the following:

- A complete description of special type brace;
- The beneficiaries mental and physical ability to use the device;
- Whether the device is a replacement;
- Whether training is indicated; and
- The plan of training, when indicated

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Equipment Maintenance and Repair

Medicaid will reimburse for the maintenance and repair of equipment only when the following conditions are met:

- Equipment is covered by Medicaid;
- Equipment is the personal property of the beneficiary;
- Item is still medically necessary;
- The equipment is used exclusively by the beneficiary;
- No other payment source is available to pay for the needed repairs;
- Equipment damage is not due to misuse, abuse, neglect, loss or wrongful disposition by the beneficiary, the beneficiary's caregiver, or the provider;
- Equipment maintenance is performed by a qualified technician;
- Maintenance is not currently covered under a manufacturer's or provider's warranty agreement; and
- Maintenance is not performed on a duplicate type of item already being maintained for the beneficiary during the maximum limit period

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Limitations for Replacement of Equipment

Medicaid will not replace equipment that is lost, destroyed, or damaged as a result of misuse, abuse, neglect, loss, or wrongful disposition of equipment by the beneficiary, the beneficiary's caregiver(s), or the provider. At a minimum, examples of equipment misuse, abuse, neglect, loss or wrongful disposition by the beneficiary, the beneficiary's caregiver, or the provider include, but are not limited to the following:

- Failure to clean and maintain the equipment as recommended by the equipment manufacturer;
- Failure to store the equipment in a secure and covered area when not in use; and
- Loss, destruction, or damage to the equipment caused by the malicious, intentional, or negligent acts of the beneficiary, the beneficiary's caregiver, or the provider

If equipment is stolen or destroyed in a fire, the provider must obtain, in a timely manner, a completed police or insurance report that describes the specific medical equipment that was stolen or destroyed. The police or insurance report must be submitted with the new PA request.

Medicaid may replace equipment when the beneficiary's medical necessity changes. The provider must submit the documentation required to justify the purchase of the replacement equipment.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.1: Services and Limitations)

Artificial Larynxes

An artificial larynx is approved only if the larynx is removed and the beneficiary is unable to use an esophageal voice. Repairs and batteries are included.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Ventilators (Applies for 2 Years of Age and Older)

Ventilators are covered to treat neuromuscular diseases, thoracic restrictive diseases, and chronic respiratory failure consequent to chronic obstructive pulmonary disease.

For members 2 years of age and older, ventilators are not covered when used only to deliver continuous or intermittent positive airway pressure for adults and children. Any type of ventilator would not be Medically Necessary when:

- The ventilator is used only in a bi-level PAP (HCPCS codes E0470 and E0471) mode
- Used for conditions that qualify for use of a Respiratory Assistance Device (RAD) that are not life-threatening conditions where interruption of respiratory support would quickly lead to serious harm or death

- Ventilators, such as Trilogy mechanical ventilators (HCPCS codes E0465 and E0466), used for the treatment of conditions that deliver continuous or intermittent positive airway pressure are not Medically Necessary

Mechanical ventilators (HCPCS codes E0465 and E0466) are considered medically necessary in certain clinical scenarios. For medical necessity clinical coverage criteria, refer to the InterQual® Medicare: Durable Medical Equipment, Ventilators.

Click [here](#) to view the InterQual® criteria.

Note:

- Ventilators must not be billed using codes for CPAP (HCPCS code E0601) or bi-level PAP (HCPCS codes E0470, E0471, and E0472). The use of CPAP or bi-level PAP HCPCS codes to bill a ventilator is incorrect coding, even if the ventilator is only being used in CPAP or bi-level mode.

Ventilator Assist Devices

PAP Therapy

For the evaluation of PAP therapy, Hypopnea is defined as an abnormal respiratory event lasting at least 20 seconds associated with at least a 30 percent reduction in thoracoabdominal movement or airflow as compared to baseline, and with at least a 4 percent decrease in oxygen saturation.

The apnea-hypopnea index (AHI) is defined as the average number of episodes of apnea and hypopnea per hour and must be based on a minimum of two hours of sleep without the use of a positive airway pressure device, reported by Polysomnography using actual recorded hours of sleep (i.e., the AHI may not be extrapolated or projected).

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Bi-level Positive Airway Pressure

The following policy guidelines apply to all ventilator assist devices:

- All equipment needs, including emergency equipment, must be prior authorized. The PAU will act on emergency requests and give a decision within two working days. If not an emergency, the PAU will act on written requests and give a decision within 25 days. Unless the physician can clearly justify purchase of the equipment, a rental trial period of up to three months can be requested to have an adequate trial period to document appropriateness;
- Other equipment, such as low pressure alarms, must be separately documented to show medical necessity. Low pressure alarms will be approved for beneficiaries who are ventilator dependent or at risk for a life threatening event. Pulse oximetry, due to its technology limitations, is not reimbursable for home use;
- These guidelines exist to assist the physician and the fiscal intermediary to efficiently approve most applications but allow physicians to request consideration for beneficiaries which for unique reasons fall outside criteria. All medical providers are expected to preserve pertinent information which may periodically be surveyed to evaluate these criteria in the future;
- Non-disposable, reusable supplies should be prescribed, if appropriate, for medical care and economical reasons. Periodic exacerbations may increase supply needs; therefore, an extra prescription should be written. The prescription should be written out "As needed" and not by using the acronym "prn" so it can be used anytime during a several month span; and
- The use of oxygen must be considered for those beneficiaries where these devices fail to adequately improve the beneficiary's condition. There must be documentation of satisfactory clinical improvement such that mechanical ventilation through a tracheotomy tube is justifiably avoided.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Continuous Positive Airway Pressure

A continuous positive airway pressure (CPAP) machine is used to treat beneficiaries who have moderate to severe obstructive sleep apnea.

A respiratory cycle is defined as an inspiration, followed by expiration. Polysomnography is the continuous and simultaneous monitoring and recording of various physiological and pathophysiological parameters of sleep with physician review, interpretation, and report. It must include sleep staging, which is defined to include a 1-4 lead electroencephalogram (EEG), and electrooculogram (EOG), and a submental electromyogram (EMG).

Polysomnography must also include at least the following additional parameters of sleep: airflow, respiratory effort, and oxygen saturation by oximetry. It may be performed as either a whole night study for diagnosis only or as a split night study to diagnose and initially evaluate treatment. Apnea is defined as the cessation of airflow for at least 10 seconds documented on a polysomnogram.

Criteria for Adults

A single level CPAP device is covered if the beneficiary has a diagnosis of obstructive sleep apnea (OSA), documented by an attended facility-based polysomnogram, and meets either of the following criteria:

- The AHI is greater than or equal to 15 events per hour; or
- The AHI is from 5 to 14 events per hour with documented symptoms of:
 - Excessive daytime sleepiness, impaired cognition, mood disorders, or insomnia; or
 - Hypertension, ischemic heart disease, or history of stroke

Polysomnographic studies must be performed in a facility based sleep study laboratory and not in the home or in a mobile facility. These labs must be qualified providers of Medicare or Medicaid services and comply with all applicable state regulatory requirements. Polysomnographic studies may not be performed by a DME provider.

Pediatric Criteria (Under 21 Years of Age)

- A single level CPAP device is covered if the beneficiary has a diagnosis of OSA documented by an attended, facility-based polysomnogram and there is:
- Documentation of physical exam (including airway) and of any other medical condition, which may be correctable (e.g., tonsillectomy and/or adenoidectomy) prior to the institution of assisted ventilation;
- Documentation of how sleep disturbance reduces the quality of life and affects the activities of daily living;
- Prescription by a physician with training and expertise in pediatric respiratory sleep disorders;
- Documentation of the medical diagnosis, which is known to cause respiratory/sleep disorders;
- Sleep or respiratory study documenting two or more of the following:
 - Oxygen saturation of less than 90 percent pulse oximetry or partial pressure of transcutaneous or arterial of less than 60mm. Hg.;
 - Carbon dioxide greater than 55 mm. Hg. By end tidal, transcutaneous, arterial, or capillary blood measurement; and
 - Apnea of 10 to 20 seconds duration on the average of one per hour
- A follow up plan should be submitted identifying the responsible physician or facility, giving data collected to demonstrate the success or failure of intervention, and showing a visit within the first month of use and a second assessment within the first three months of use;
- Indication of a responsible, committed home environment and of caregivers properly trained in appropriate respiratory care; and
- A written plan for home health follow up care

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria)

Medical Supplies

Durable Medical Equipment (DME)/supplies are covered when medical necessity criteria are met for use as part of the medical care of a beneficiary. Refer to the Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.2: Specific Coverage Criteria.

Non-Covered DME Services and Items

A non-covered service, item or supply is not available for reimbursement. Listed below are items and services that are not reimbursed by Medicaid through the DME program.

- Clinically unproven equipment;
- Comfort or convenience equipment;
- Dentures;
- Disposable supplies customarily provided as part of a nursing or personal care service or a medical diagnostic or monitoring procedure;
- Electric lifts (manual lifts are covered);
- Emergency and non-emergency alert devices;
- Environmental modifications (e.g., home, bathroom, ramps, etc.);
- Equipment designed for use by a physician or trained medical personnel;
- Experimental equipment;
- Facilitated communications (FC);
- Furniture and other items which do not serve a medical purpose;
- Hand Held Showers;
- Investigational equipment;
- Items used for cosmetic purposes;
- Personal comfort, convenience, or general sanitation items;
- Physical fitness equipment;
- Precautionary-type equipment (e.g., power generators, backup oxygen equipment);
- Rehabilitation Equipment;
- Reimbursement for delivery or delivery mileage of Medical Supplies;
- Routine and first aid items;
- Safety alarms and alert systems/buttons;
- Scooters;
- Seat lifts and recliner lifts;
- Standard car seats;
- Supplies or equipment covered by Medicaid per diem rates (nursing home residents maybe approved for orthotics and prosthetics, but not for DME and supplies);
- Televisions, telephones, VCR machines and devices designed to produce music or provide entertainment;
- Training equipment or self-help equipment;
- Van lifts;
- Wheelchair Lifts; and
- Wheelchair Ramps

Note: This list is not all inclusive.

(Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, Section 18.1: Services and Limitations)

Definitions

Check the federal, state, or contractual definitions that supersede the definitions below.

Durable Medical Equipment (DME): Medical equipment that is all of the following:

- Suitable for use in any setting in which normal life activities take place
- Can withstand repeated use
- Generally not useful to an individual in the absence of a disability, illness, or injury
- Can be reusable or removable
- Is not implantable within the body
- Primarily and customarily used to serve a medical purpose
- Meets the state definition of DME

Medical Supplies: Supplies are health care related items that are consumable or disposable, or cannot withstand repeated use by more than one individual, that are required to address an individual medial disability, illness, or injury (CFR § 440.70).

Reasonable Useful Lifetime: RUL is the expected minimum lifespan for the item. It starts on the initial date of service and runs for the defined length of time. The default RUL for Durable Medical Equipment is 5 years. RUL is also applied to other non-DME items such as orthoses and prostheses. RUL is not applied to supply items.

Applicable Codes

UnitedHealthcare has adopted the requirements and intent of the National Correct Coding Initiative. The Centers for Medicare & Medicaid Services (CMS) has contracted with Palmetto to manage Pricing, Data and Coding (PDAC) for Durable Medical Equipment, Prosthetics, Orthotics and Supplies (DMEPOS). This notice is to confirm UnitedHealthcare has established the PDAC as a source for correct coding and coding clarification.

Benefit Considerations

Contact Lenses Scleral Bandages (Shells)

Contact lenses or scleral shells that are used to treat an injury or disease (e.g., corneal abrasion, keratoconus, or severe dry eye) are not considered DME and may be covered as a therapeutic service. Please check the federal, state, or contractual requirements for coverage.

Cranial Remolding Orthosis

Cranial molding helmets (cranial remolding orthosis, billed with S1040) used to facilitate a successful post-surgical outcome are covered. For all indications, refer to the Medical Policy titled Plagiocephaly and Craniosynostosis Treatment (for Louisiana Only).

Note: A protective helmet (HCPCS code A8000–A8004) is not a cranial remolding device. It is considered a safety device worn to prevent injury to the head rather than a device needed for active treatment.

Implanted Devices

Any device, appliance, pump, machine, stimulator, or monitor that is fully implanted into the body is not covered as DME. (If covered, the device is covered as part of the surgical service.)

Cochlear Implant Benefit Clarification: The external components (i.e., speech processor, microphone, and transmitter coil) are considered under the DME benefit, and the implantable components are considered under the medical-surgical benefit. Reference the federal, state, or contractual requirements to determine if there are DME benefits for repair or replacement of external components.

Insulin Pumps

Insulin pumps, disposable and durable, are covered. For state specific information on mandated coverage of diabetes supplies, reference the federal, state, or contractual requirements. Refer to the Medical Policy titled Continuous Glucose Monitoring and Insulin Delivery for Managing Diabetes (for Louisiana Only).

Lymphedema Stockings for the Arm

Post-mastectomy lymphedema stockings for the arm are considered DME. For state specific information on mandated coverage, reference the state or contractual requirements.

References

Centers for Disease Control and Prevention. https://www.cdc.gov/growthcharts/clinical_charts.htm. Accessed May 24, 2023.

Code of Federal Regulations (CFR). Home health services. 42 CFR 440.70. Available at: <https://www.ecfr.gov/current/title-42/chapter-IV/subchapter-C/part-440/subpart-A/section-440.70>. Accessed July 23, 2023.

Louisiana Department of Health Durable Medical Equipment Provider Manual. Chapter Eighteen of the Medicaid Services Manual. Issued September 1, 2010. <https://www.lamedicaid.com/provweb1/Providermanuals/manuals/DME/DME.pdf>. Accessed October 30, 2023.

Noridian Healthcare Solutions. Reasonable Useful Lifetime and Duplicate Items – Billing Reminder.(April 2011) <https://med.noridianmedicare.com/web/jddme/article-detail/-/view/2230703/reasonable-useful-lifetime-and-duplicate-items-billing-reminder#main-content>. Accessed June 13, 2023.

State Medicaid contracts.

Policy History/Revision Information

Date	Summary of Changes
05/01/2024	<p>Coverage Rationale</p> <p><i>Electric Breast Pump</i></p> <p>Human Milk Storage Bags</p> <ul style="list-style-type: none"> ● Added language to indicate human milk storage bags are designed to safely store and protect expressed human milk for feeding a child <ul style="list-style-type: none"> ○ Medicaid covers 100 human milk storage bags per month for lactating beneficiaries ○ The Medicaid reimbursement rate on file covers a one month supply of storage bags <p><i>Equipment Maintenance and Repair</i></p> <ul style="list-style-type: none"> ● Revised language to indicate Medicaid will reimburse for the maintenance and repair of equipment only when the following conditions are met: <ul style="list-style-type: none"> ○ Equipment is covered by Medicaid ○ Equipment is the personal property of the beneficiary ○ Item is still medically necessary ○ The equipment is used exclusively by the beneficiary ○ No other payment source is available to pay for the needed repairs ○ Equipment damage is not due to misuse, abuse, neglect, loss or wrongful disposition by the beneficiary, the beneficiary’s caregiver, or the provider ○ Equipment maintenance is performed by a qualified technician ○ Maintenance is not currently covered under a manufacturer’s or provider’s warranty agreement ○ Maintenance is not performed on a duplicate type of item already being maintained for the beneficiary during the maximum limit period <p><i>Artificial Larynxes</i></p> <ul style="list-style-type: none"> ● Revised language to indicate an artificial larynx is approved only if the larynx is removed and the beneficiary is unable to use an esophageal voice; repairs and batteries are included <p><i>Ventilator Assist Devices</i></p> <p>Bi-level Positive Airway Pressure</p> <ul style="list-style-type: none"> ● Removed language indicating: <ul style="list-style-type: none"> ○ Bi-level PAP devices (HCPCS codes E0470 and E0471) are considered medically necessary in certain clinical scenarios; ○ For medical necessity criteria, refer to the InterQual® CP: Durable Medical Equipment, Noninvasive Airway Assistive Devices ● Added language to indicate the following policy guidelines apply to all ventilator assist devices: <ul style="list-style-type: none"> ○ All equipment needs, including emergency equipment, must be prior authorized <ul style="list-style-type: none"> ▪ The PAU will act on emergency requests and give a decision within two working days; if not an emergency, the PAU will act on written requests and give a decision within 25 days ▪ Unless the physician can clearly justify purchase of the equipment, a rental trial period of up to three months can be requested to have an adequate trial period to document appropriateness ○ Other equipment, such as low pressure alarms, must be separately documented to show medical necessity

Date	Summary of Changes
	<ul style="list-style-type: none"> ▪ Low pressure alarms will be approved for beneficiaries who are ventilator dependent or at risk for a life threatening event ▪ Pulse oximetry, due to its technology limitations, is not reimbursable for home use ○ These guidelines exist to assist the physician and the fiscal intermediary to efficiently approve most applications but allow physicians to request consideration for beneficiaries which for unique reasons fall outside criteria; all medical providers are expected to preserve pertinent information which may periodically be surveyed to evaluate these criteria in the future ○ Non-disposable, reusable supplies should be prescribed, if appropriate, for medical care and economical reasons <ul style="list-style-type: none"> ▪ Periodic exacerbations may increase supply needs; therefore, an extra prescription should be written ▪ The prescription should be written out “as needed” and not by using the acronym “prn” so it can be used anytime during a several month span ○ The use of oxygen must be considered for those beneficiaries where these devices fail to adequately improve the beneficiary’s condition; there must be documentation of satisfactory clinical improvement such that mechanical ventilation through a tracheotomy tube is justifiably avoided <p>Continuous Positive Airway Pressure</p> <ul style="list-style-type: none"> ● Added language to indicate: <ul style="list-style-type: none"> ○ A continuous positive airway pressure (CPAP) machine is used to treat beneficiaries who have moderate to severe obstructive sleep apnea ○ A respiratory cycle is defined as an inspiration, followed by expiration ○ Polysomnography is the continuous and simultaneous monitoring and recording of various physiological and pathophysiological parameters of sleep with physician review, interpretation, and report; it must include sleep staging, which is defined to include a 1-4 lead electroencephalogram (EEG), and electrooculogram (EOG), and a submental electromyogram (EMG) ○ Polysomnography must also include at least the following additional parameters of sleep: airflow, respiratory effort, and oxygen saturation by oximetry <ul style="list-style-type: none"> ▪ It may be performed as either a whole night study for diagnosis only or as a split night study to diagnose and initially evaluate treatment ▪ Apnea is defined as the cessation of airflow for at least 10 seconds documented on a polysomnogram <p>Criteria for Adults</p> <ul style="list-style-type: none"> ○ A single level CPAP device is covered if the beneficiary has a diagnosis of obstructive sleep apnea (OSA), documented by an attended facility-based polysomnogram, and meets either of the following criteria: <ul style="list-style-type: none"> ▪ The AHI is greater than or equal to 15 events per hour ▪ The AHI is from 5 to 14 events per hour with documented symptoms of: <ul style="list-style-type: none"> – Excessive daytime sleepiness, impaired cognition, mood disorders, or insomnia – Hypertension, ischemic heart disease, or history of stroke ○ Polysomnographic studies must be performed in a facility based sleep study laboratory and not in the home or in a mobile facility <ul style="list-style-type: none"> ▪ These labs must be qualified providers of Medicare or Medicaid services and comply with all applicable state regulatory requirements ▪ Polysomnographic studies may not be performed by a DME provider <p>Pediatric Criteria (Under 21 years of Age)</p> <ul style="list-style-type: none"> ○ A single level CPAP device is covered if the beneficiary has a diagnosis of OSA documented by an attended, facility-based polysomnogram, and there is: <ul style="list-style-type: none"> ▪ Documentation of physical exam (including airway) and of any other medical condition, which may be correctable (e.g., tonsillectomy and/or adenoidectomy) prior to the institution of assisted ventilation

Date	Summary of Changes
	<ul style="list-style-type: none"> ▪ Documentation of how sleep disturbance reduces the quality of life and affects the activities of daily living ▪ Prescription by a physician with training and expertise in pediatric respiratory sleep disorders ▪ Documentation of the medical diagnosis, which is known to cause respiratory/sleep disorders ▪ Sleep or respiratory study documenting two or more of the following: <ul style="list-style-type: none"> – Oxygen saturation of less than 90 percent pulse oximetry or partial pressure of transcutaneous or arterial of less than 60mmHg – Carbon dioxide greater than 55 mmHg by end tidal, transcutaneous, arterial, or capillary blood measurement – Apnea of 10 to 20 seconds duration on the average of one per hour ▪ A follow up plan should be submitted identifying the responsible physician or facility, giving data collected to demonstrate the success or failure of intervention, and showing a visit within the first month of use and a second assessment within the first three months of use ▪ Indication of a responsible, committed home environment and of caregivers properly trained in appropriate respiratory care ▪ A written plan for home health follow up care <p>Medical Supplies</p> <ul style="list-style-type: none"> ● Removed language indicating: <ul style="list-style-type: none"> ○ The covered items and services include medical supplies ○ Urinary catheters are approved only if the beneficiary’s medical condition necessitates the use of a catheter ● Replaced reference to the “Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, <i>Section 18.1: Services and Limitations</i>” with “Louisiana Durable Medical Equipment Provider Manual, Chapter 18: Durable Medical Equipment, <i>Section 18.2: Specific Coverage Criteria</i>” <p>Definitions</p> <ul style="list-style-type: none"> ● Removed definition of “Injury” <p>Benefit Considerations</p> <ul style="list-style-type: none"> ● Added content/language (relocated from the <i>Coverage Rationale</i> section of the policy) pertaining to: <ul style="list-style-type: none"> ○ Contact lenses and scleral bandages (shells) ○ Cranial remolding orthosis ○ Implanted devices ○ Insulin pumps ○ Lymphedema stockings for the arm <p>Supporting Information</p> <ul style="list-style-type: none"> ● Updated <i>References</i> section to reflect the most current information ● Archived previous policy version CS032LA.R

Instructions for Use

This Medical Policy provides assistance in interpreting UnitedHealthcare standard benefit plans. When deciding coverage, the federal, state or contractual requirements for benefit plan coverage must be referenced as the terms of the federal, state or contractual requirements for benefit plan coverage may differ from the standard benefit plan. In the event of a conflict, the federal, state or contractual requirements for benefit plan coverage govern. Before using this policy, please check the federal, state or contractual requirements for benefit plan coverage. UnitedHealthcare reserves the right to modify its Policies and Guidelines as necessary. This Medical Policy is provided for informational purposes. It does not constitute medical advice.

UnitedHealthcare may also use tools developed by third parties, such as the InterQual® criteria, to assist us in administering health benefits. The UnitedHealthcare Medical Policies are intended to be used in connection with the independent

professional medical judgment of a qualified health care provider and do not constitute the practice of medicine or medical advice.