

December 6, 2024

LEAH NOAEILL ABMRC LLC (DBA ABM RESPIRATORY CARE) 860 BLUE GENTIAN RD SUITE 200 EAGAN, MN 55121

Document Control Number (DCN): 24263003000000

Manufacturer Name	Product Name	Model Number	Assigned HCPCS Code(s)
ABMRC LLC	BIWAZE CLEAR	BK181	E0469
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - INFANT FACE MASK	BK902	A7021
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - CHILD FACE MASK	BK291	A7021
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - ADULT SMALL FACE MASK	BK289	A7021
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - ADULT MEDIUM FACE MASK	BK287	A7021



Manufacturer Name	Product Name	Model Number	Assigned HCPCS Code(s)
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - ADULT LARGE FACE MASK	BK285	A7021
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - MOUTHPIECE	BK280	A7021
ABMRC LLC	BIWAZE CLEAR DUAL LUMEN BREATHING CIRCUIT - TRACH	BK293	A7021

Dear LEAH NOAEILL,

The Pricing, Data Analysis, and Coding (PDAC) Contractor has reviewed the product(s) listed above and has approved the listed Healthcare Common Procedure Coding System (HCPCS) code(s) for billing the four Durable Medical Equipment Medicare Administrative Contractors (DME MACs).

The PDAC Contractor provides coding assistance to manufacturers to ensure proper coding of Durable Medical Equipment, Prosthetics, Orthotics, and Supplies (DMEPOS). The PDAC publishes coding decisions based on the coding guidelines established by the Local Coverage Determinations (LCDs) and associated Policy Articles and any related Advisory Articles established by the DME MACs. All products submitted to the PDAC for a coding verification review are examined by coders and professionals following a formal, standardized process.

Based on this review and application of DME MAC policy, the HCPCS code(s) listed below should be used when billing the DME MACs:

E0469 LUNG EXPANSION AIRWAY CLEARANCE, CONTINUOUS HIGH FREQUENCY OSCILLATION, AND NEBULIZATION DEVICE

A7021 SUPPLIES AND ACCESSORIES FOR LUNG EXPANSION AIRWAY CLEARANCE, CONTINUOUS HIGH FREQUENCY OSCILLATION, AND NEBULIZATION DEVICE (E.G., HANDSET, NEBULIZER KIT, BIOFILTER)

If you disagree with this decision, you may request a reconsideration within 45 calendar days of the Coding Verification letter's date and provide evidence to substantiate a reconsideration of PDAC's original coding determination. To request a reconsideration, complete the Reconsideration Request form located on the PDAC website at www.dmepdac.com. If your request for a reconsideration of PDAC's coding determination is made after the 45 calendar day time frame, it will require a new application and documentation to support the request.

It is the responsibility of manufacturers and distributors to notify the PDAC immediately of any changes involving their products, as listed on the Product Classification List (PCL) on the Durable Medical Equipment Coding System (DMECS). Further information for requesting updates to the PCL can be found on the PDAC website at www.dmepdac.com. It is also the responsibility of manufacturers and distributors to assure their websites and product marketing materials accurately reflect the product reviewed by the PDAC and the coding decision assigned.

An assignment of the HCPCS code(s) to product(s) is not an approval or endorsement of the product(s) by Medicare or Palmetto GBA; nor does it imply or guarantee claim reimbursement or coverage.

If you have questions, please contact the PDAC HCPCS Helpline at (877) 735-1326 during the hours of 9:30 a.m. to 5:00 p.m. ET, Monday through Friday. You may also visit our <u>website</u> to chat with one of our representatives or select the Contact Us button at the top of the page for email, FAX or postal mail information.

Sincerely,

Pricing, Data Analysis, and Coding (PDAC) Palmetto GBA, LLC www.dmepdac.com