

**Notice Date:** December 2021  
**To:** Ohio Providers  
**From:** CareSource  
**Subject:** Notice of Philips Respironics Voluntary Medical Device Recall  
**Effective Date:** June 14, 2021

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**Please Note:** This communication is an update from the [network notification](#) dated Sept. 13, 2021.

### **Summary**

As part of CareSource's commitment to continuous improvement, we are focused on ensuring quality care for all our members through active provider engagement. On June 14, 2021, Philips Respironics ("Philips") announced a voluntary recall pertaining to certain ventilators and devices used to treat obstructive sleep apnea, including continuous positive airway pressure ("CPAP") devices and bi-level positive airway pressure ("BiPAP") devices (collectively referred to as "Devices").

The Devices are being recalled due to two issues related to the polyester-based polyurethane ("PE-PUR") sound abatement foam used in the Devices:

- PE-PUR foam may degrade into particles that may enter the Device's air pathway and be ingested or inhaled by the user; and
- The PE-PUR foam may off-gas certain chemicals.

### **Impact**

The FDA issued a Safety Communication on June 30, 2021 regarding Philips' voluntary recall which summarizes major issues and considerations related to the recall.

### ***Please note this section is new***

CareSource wants to ensure members have necessary tools to improve their overall health. The

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