



**Mallinckrodt™**  
Pharmaceuticals

## Technical Bulletin

Ref: **TB-17002-ENG**

Product: INOmax DS<sub>IR</sub>®, INOmax DS<sub>IR</sub>® Plus,  
INOmax DS<sub>IR</sub>® Plus MRI and INOflo DS®

Affected parts: INOmax DS<sub>IR</sub>, INOmax DS<sub>IR</sub> Plus,  
INOmax DS<sub>IR</sub> Plus MRI and INOflo DS

Subject: **Purging the Regulator Supply Line and/or the Delivery System**

Issue date: November 2017

Priority: Low

Classification: Information for Distribution

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This technical bulletin is written to advise clinicians that testing has been completed which demonstrates that the INOmax DS<sub>IR</sub>, INOmax DS<sub>IR</sub> Plus, INOmax DS<sub>IR</sub> Plus MRI and INOflo DS may stand idle for 24 hours after a successful Pre-Use procedure and depressurization of the regulator(s) has been completed.

If the device is not placed into use within these 24 hours repeat the Pre-Use procedure.



**WARNING: If the INOmax DS<sub>IR</sub>, INOmax DS<sub>IR</sub> Plus, INOmax DS<sub>IR</sub> Plus MRI or INOflo DS is depressurized and is not used on a patient within 24 hours, repeat the pre-use procedure.**

For technical assistance regarding any of the above devices, please contact Technical Support at 1-877-566-9466 (North America) or your specific country manager.

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