



11 April 2023

**Shortage of DBL DOBUTAMINE HYDROCHLORIDE 250mg/20ml solution for injection and alternative supply arrangement under Section 19A of the *Therapeutic Goods Act, 1989***

Dear Healthcare Professional,

This communication is sent by ORSPEC Pharma to notify your organisation that due to the shortage of **DBL DOBUTAMINE HYDROCHLORIDE 250mg/20ml solution for injection (AUST R 46451)**. ORSPEC Pharma has arranged the supply of an alternative product on a temporary basis.

**Dobutamine 12.5 mg/ml (250mg/20mL) concentrate for solution for infusion (Hameln)**, is NOT registered in Australia and supply is granted under an exemption granted by the Therapeutic Goods Administration (TGA) under Section 19A of the *Therapeutic Goods Act, 1989* until **30 September 2023**.

**Dobutamine 12.5 mg/ml (250mg/20mL) concentrate for solution for infusion (Hameln)** is approved for use under Section 19A for the following indications:

*Dobutamine Hydrochloride Injection is indicated in adults who require short-term treatment of cardiac failure secondary to acute myocardial infarction, or cardiac surgery.*

**The s19A approved UK product is identical in active ingredient and strength to the Australian registered product. The differences between the products are noted below:**

	<b>ARTG product</b> DBL DOBUTAMINE HYDROCHLORIDE 250mg/20ml solution for injection (AUST R 46451)	<b>S19A product</b> Dobutamine 12.5 mg/ml (250mg/20mL) concentrate for solution for infusion (Hameln)
<b>Excipients</b>	Sodium metabisulfite Water for injections	Sodium Metabisulfite Sodium Hydroxide Hydrochloric Acid Water for Injections Carbon Dioxide
<b>Pack size</b>	single vial	pack of 10 vials

**Dobutamine 12.5 mg/ml (250mg/20mL) concentrate for solution for infusion (Hameln)** is registered in the United Kingdom and is packaged in the English language. For dosing and administration information, please refer to the Australian Product Information for **DBL DOBUTAMINE HYDROCHLORIDE 250mg/20ml** available at <https://www.ebs.tga.gov.au/>

#### **Adverse Event Reporting**

Reporting any suspected adverse event is important for the continued monitoring of the safety of all medicines. Any adverse events which are experienced with **Dobutamine 12.5 mg/ml (250mg/20mL) concentrate for solution for infusion (Hameln)**, should be reported by healthcare professionals and patients to ORSPEC Pharma on 024 339 4239 or email at [sas@orspecpharma.com](mailto:sas@orspecpharma.com). Alternatively, this information can be reported to the TGA at <https://www.tga.gov.au/reporting-problems>

Please forward this information to relevant staff members in your organisation.

For further information, please contact ORSPEC Pharma on 024 339 4239 or email [sas@orspecpharma.com](mailto:sas@orspecpharma.com).

Yours sincerely,



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ORSPEC Pharma