Neonatal Medication Guideline

Clinical Guideline

Dobutamine

Policy developed by: SA Maternal, Neonatal & Gynaecology Community of Practice
Approved SA Health Safety & Quality Strategic Governance Committee on: 6 October 2017
Next review due: 6 October 2020

Summary
The purpose of this guideline is to guide nursing, midwifery, medical and pharmacy staff in the dosing and administration of dobutamine

Keywords
Dobutamine, neonatal medication guideline, inotrope, circulatory support, severe sepsis, septic shock, hypoperfusion, hypotension, tachycardia, arrhythmia, rapid atrial fibrillation, phaeochromocytoma

Policy history
Is this a new policy? N
Does this policy amend or update an existing policy? Y v1.0
Does this policy replace an existing policy? N
If so, which policies?

Applies to
All SA Health Portfolio
All Department for Health and Ageing Divisions
All Health Networks
CALHN, SALHN, NALHN, CHSALHN, WCHN, SAAS

Staff impact
All Clinical, Medical, Midwifery, Nursing, Students, Allied Health, Emergency, Mental Health, Pathology, Pharmacy

PDS reference
CG024

Version control and change history

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South Australian Neonatal Medication Guidelines

DOBUTamine
250mg/20mL injection

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Note

This guideline provides advice of a general nature. This statewide guideline has been prepared to promote and facilitate standardisation and consistency of practice, using a multidisciplinary approach. The guideline is based on a review of published evidence and expert opinion.

Information in this statewide guideline is current at the time of publication.

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Health practitioners in the South Australian public health sector are expected to review specific details of each patient and professionally assess the applicability of the relevant guideline to that clinical situation.

If for good clinical reasons, a decision is made to depart from the guideline, the responsible clinician must document in the patient's medical record, the decision made, by whom, and detailed reasons for the departure from the guideline.

This statewide guideline does not address all the elements of clinical practice and assumes that the individual clinicians are responsible for discussing care with consumers in an environment that is culturally appropriate and which enables respectful confidential discussion. This includes:

- The use of interpreter services where necessary,
- Advising consumers of their choice and ensuring informed consent is obtained,
- Providing care within scope of practice, meeting all legislative requirements and maintaining standards of professional conduct, and
- Documenting all care in accordance with mandatory and local requirements

This is a High Risk Medication

An overdose can be rapidly fatal.

Dose and Indications

Circulatory Support

Hypoperfusion, hypotension related to myocardial dysfunction

Severe sepsis and septic shock

Intravenous infusion

5 to 25 micrograms/kg/minute

Start at 5 micrograms/kg/minute and titrate dose every 10-20 minutes if required according to clinical response.
Preparation and Administration

Intravenous Infusion

Administer preferably via a central line but may be used peripherally in an emergency when central access is not available.

Select the strength required based on the weight of the infant in the context of any fluid restrictions. Maximum concentration for infusion is 4mg/mL.

DOBUTamine Concentration Selection Tables can be found on the following pages of this guideline to assist prescribers to gauge which strength is best for the patient.

Dilute the appropriate volume of the 12.5mg/mL DOBUTamine solution using compatible fluid; and administer by continuous infusion. Diluted preparation is stable for 24 hours at room temperature. Discard any remaining solution.

The three standard concentrations to select from are:

- DOBUTamine 1mg/mL (1000microgram/mL)
- DOBUTamine 2mg/mL (2000microgram/mL)
- DOBUTamine 4mg/mL (4000microgram/mL)

**Formulae**

To calculate infusion rate (mL/hr):

\[
\text{Rate (mL/hr)} = \frac{60 \times \text{Dose (micrograms/kg/min)} \times \text{Weight(kg)}}{\text{Strength (microgram/mL)}}
\]

To calculate the dose (micrograms/kg/min):

\[
\text{Dose (microgram/kg/min)} = \frac{\text{Rate(mL/hr)} \times \text{Strength (microgram/mL)}}{60 \times \text{Weight (kg)}}
\]
DOBUTamine
250mg/20mL injection

Dilution for DOBUTamine 1000microgram/mL

To make **25ml** syringe:
Dilute 2 mL DOBUTamine (12.5mg/mL) with 23mL of compatible fluid (total of 25mL). The resulting solution contains 1000micrograms/mL

To make **50ml** syringe:
Dilute 4mL DOBUTamine (12.5mg/mL) with 46mL of compatible fluid (total of 50mL). The resulting solution contains 1000micrograms/mL

Table ONE: Concentration selection table for DOBUTamine 1000micrograms/mL
Recommended for neonates weighing <1kg

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DOBUTamine
250mg/20mL injection

Dilution for DOBUTamine 2000microgram/mL

To make 25ml syringe:
Dilute 4 mL DOBUTamine (12.5mg/mL) with 21mL of compatible fluid (total of 25mL). The resulting solution contains 2000micrograms/mL

To make 50ml syringe:
Dilute 8mL DOBUTamine (12.5mg/mL) with 42mL of compatible fluid (total of 50mL). The resulting solution contains 2000micrograms/mL

Table 2: Concentration selection table for DOBUTamine 2000micrograms/mL
Generally used for neonates weighing 1kg to 3kg

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Dilution for DOBUTamine 4000microgram/mL

To make 25ml syringe:
Dilute 8 mL DOBUTamine (12.5mg/mL) with 17mL of compatible fluid (total of 25mL). The resulting solution contains 4000micrograms/mL.

To make 50ml syringe:
Dilute 16mL DOBUTamine (12.5mg/mL) with 34mL of compatible fluid (total of 50mL). The resulting solution contains 4000micrograms/mL.

Table 3: Concentration selection table for DOBUTamine 4000micrograms/mL
Generally used for neonates >3kg

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Compatible Fluids
Glucose 5%, glucose 10%, glucose / sodium chloride combinations, sodium chloride 0.9%

Adverse Effects

Common
Tachycardia, increased blood pressure, ventricular ectopic activity, hypotension (if patient is hypovolemic)

Infrequent
Phlebitis, rash, ventricular tachycardia or fibrillation, cutaneous vasodilation

Rare
Allergic reaction (due to sodium metabisulfite)

Monitoring
- Observe intravenous site for inflammation, extravasation and extreme vasoconstriction (tracking)
- Continuous heart rate
- Invasive blood pressure monitoring is recommended

Practice Points
- Contraindications include ventricular arrhythmias, rapid atrial fibrillation and phaeochromocytoma
- Hypovolaemia - should be corrected prior to DOBUTamine administration
- DOBUTamine is incompatible with alkaline solutions (eg sodium bicarbonate, phenytoin).
- Do not bolus other drugs via DOBUTamine infusion
- Caution when changing IV line (avoid bolus or prolonged interruption of drug infusion)
- Dobutamine solutions may show a pink discolouration which increases with time. This colour is due to a slight oxidation of the drug. However, there is no significant loss of drug within the recommended storage times for solutions of the drug.

Version control and change history

PDS reference: OCE use only

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