



Australian Government
Department of Health
Therapeutic Goods Administration

TGA use only

This form, when completed, will be classified as 'For official use only'.

For guidance on how your information will be treated by the TGA see: Treatment of information provided to the TGA at <https://www.tga.gov.au/treatment-information-provided-tga>.

Special Access Scheme – Category B

Important information

Email completed form to SAS@tga.gov.au (preferred) or fax to 02 6232 6112.

The SAS Category B application form should be completed if guidance for use of an unapproved good will be met and the SAS Category A or SAS Category C pathways are not applicable.

Privacy information

For general privacy information, go to <https://www.tga.gov.au/privacy>.

The TGA is collecting personal information in this form in order to:

- Assess the application.
- Contact the medical practitioner and discuss the application where necessary.
- The personal information of the medical practitioner may be disclosed to State and Territory authorities with responsibility for therapeutic goods or medical practitioner registration.

Do not provide the name of the patient. Only provide the patient initials and other information as required on the form.

Please complete the form clearly and fully. Applications for special access will be processed in the order in which they are received.

Patient details (minimum of 3 (three) identifiers required)

Diagnosis(es) or Medical Condition(s):	Nonketotic hypoglycaemia
Indication:	Nonketotic hypoglycaemia
Clinical justification for use of product: (e.g. include seriousness of condition, details of previous treatment including reasons why a therapeutic good currently listed on the ARTG cannot be used for the treatment of this patient in this circumstance)	requiring combination of meds for m of seizures

Product details (attach efficacy and safety data to support proposed use of the product and details of intended monitoring)

Therapeutic good type			
Medicine <input checked="" type="checkbox"/>	Biological <input type="checkbox"/>	Medical device <input type="checkbox"/>	
Medicine/biological		Medical device	
Trade Name (if known)	Sponsor / Supplier	Trade name	
Sodium Benzoate			
Active Ingredient(s)		Product description (including variant ²)	
Sodium Benzoate			
Dosage form (e.g. tablet)	Strength (e.g. 1 mg/ml)	No of units	Sponsor / Supplier
Tablet	500mg		
Route of administration (e.g., IV)	Dose & frequency (1 lds)	Proposed duration of treatment	Intended date of use
PO	600mg @ TDS		
Quantity ¹ required for treatment or duration			
ongoing (100 tablets, 2 repeats)			

¹ For substances captured by the Customs (Prohibited Imports) Regulations 1956 the quantity must be provided.

² Variant means a medical device the design of which has been varied to accommodate different patient anatomical requirements (for example, relating to the shape, size, length, diameter or gauge of the device).

