



Australian Government

Department of Health and Aged Care  
Therapeutic Goods Administration

# Australian Register of Therapeutic Goods Certificate

Issued to

**Prescribing Biochemists Pty Ltd**

for approval to supply

## MACROMOLECULAR SYSTEM HEPAMIN B Tablet (Cancelled)

**ARTG Identifier**                      AUST R 19378  
**ARTG Start Date**                      2/10/1991  
**Product Type**                      Registered Medicine

| Manufacturer Details | Address                                                | Manufacturing Steps |
|----------------------|--------------------------------------------------------|---------------------|
| Manucom Pty Ltd      | 270 Lahrs Road<br>ORMEAU , QLD , 42<br>08<br>Australia | Release for supply  |
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### ARTG Standard Conditions

The above Medicine Registered has been entered on the Register subject to the following conditions:

- Conditions applicable to the relevant category and class of therapeutic goods as specified in the document "Conditions Applying to Therapeutic Goods Accepted for Registration or Listing as Goods Currently Supplied at the Commencement of the Therapeutic Goods Act 1989"

### Products Covered by This Entry

#### 1. MACROMOLECULAR SYSTEM HEPAMIN B Tablet (Cancelled)

| Container Type | Container Material | Container Condition | Container Closure | Shelf Life Time | Shelf Life Temperature | Shelf Life Conditions |
|----------------|--------------------|---------------------|-------------------|-----------------|------------------------|-----------------------|
| Bottle         | Not recorded       | Not recorded        | Not recorded      | Not recorded    | Not recorded           |                       |

### Product Specific Conditions

No specific conditions have been recorded against this entry.

### Product Permitted Indications

No permitted indications have been recorded against this entry.

### Product Indication Requirements

No indication requirements have been recorded against this entry.

### Product Standard Indications

- This product accepted for registration/listing as 'currently supplied' at the time of commencement of the Act. Indications held in ARTG paper records. (Old code)

### Product Specific Indications

No specific indications have been recorded against this entry.

### Warnings

No warnings have been recorded against this entry.

### Dosage Form

- Tablet, film coated

### Route of Administration

- Oral

**Visual Identification**

- Yellow, speckled, film coated 7/16" (11.2mm) diameter round, biconvex tablet.

**Additional Product information**

**Product Formulation(s)**

**Active Ingredients**

|                          | Quantity | Units     |
|--------------------------|----------|-----------|
| Liver extract            | 100      | mg        |
| pyridoxine hydrochloride | 8.3      | mg        |
| thiamine nitrate         | 20       | mg        |
| calcium pantothenate     | 10       | mg        |
| lysine hydrochloride     | 10       | mg        |
| nicotinamide             | 10       | mg        |
| riboflavine              | 10       | mg        |
| folic acid               | 60       | microgram |
| cyanocobalamin           | 25       | microgram |

Therapeutic Goods Administration  
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