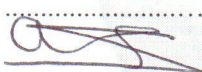


DEPARTMENT OF HEALTH
MEDICINE REGISTRATION CERTIFICATE

It is hereby certified that registration of the medicine as described below has been approved by the Medicines Control Council in terms of section 15 (3) (a) of the Medicines and Related Substances Act, 1965 (Act 101 of 1965), subject to the conditions indicated:

1. Registered name **STERIPREP 2 %**
2. Registration number **50/13.1/0841**
3. Approved name of every active ingredient and quantities thereof per dosage unit or per suitable mass or volume or unit of the medicine
**EACH 100,0 ml SOLUTION CONTAINS:
CHLORHEXIDINE GLUCONATE 2,0 % m/v**
4. Dosage form **SOLUTION**
5. Conditions under which the medicine is registered
see annexure
6. Registered in the name of (applicant)
BARRS PHARMACEUTICAL INDUSTRIES (PTY) LTD
7. Original date of registration **30 SEPTEMBER 2016**
8. Manufacturer, packer, final product release control (FPRC)/final product responsibility (FPRR)
**BARRS PHARMACEUTICAL INDUSTRIES (PTY) LTD, N'DABENI, CAPE TOWN,
RSA – MANUFACTURER, PACKER, FPRC, FPRR**

Issued at Pretoria on

 2016 -09- 30

REGISTRAR OF MEDICINES

