



# Directorate of AYUSH

(Ayurveda, Yoga & Naturopathy, Unani, Siddha and Homoeopathy)

Department of Health & Family Welfare, Govt. of Odisha

Heads of Department Building, 3<sup>rd</sup> Floor (Annex), Unit-5, Bhubaneswar-751001, Tel:0674-2394577, FAX:06742391180, Email:directorayushodisha@gmail.com



F. No. VII-DCC (AY)-19/2011

1245

/DAYUSH, dt. 17.02.2022

To

The Principal Sri Sri Nrusinghanath Ayurvedic College  
At- Nrusinghanath, P.O. - Paikmal,  
Dist - Bargarh, PIN- 768039

**Sub:** Renewal of Licence to manufacture for sale of Ayurvedic, Siddha or Unani drugs and Certificate of GMP.

**Ref:** Your application no. 1690, dt. 25.12.2021 and compliance no. 134, dt. 29.01.2022

Sir/ Madam,

Considering your application cited above and pursuant to Ministry of AYUSH, Government of India Notification no. G.S.R. 716(E), dt. 01.10.2021, the Drug Licence bearing no. OR-82/Ayur issued in favour of your firm is hereby renewed and re-issued in form 25-D which shall take effect from 01.01.2022 and shall remain valid perpetually unless sooner suspended or cancelled. The certificate of GMP (Good Manufacturing Practices) is issued alongside in form 26 E-I which shall remain valid till 31.12.2026 unless it is cancelled by the Licensing Authority subject to deposit of a certificate retention fee of rupees one thousand before the expiry of a period of every succeeding five years from the date of its issue. The total items approved under this licence remains unchanged.

You are further requested to comply the conditions as laid down in the D/L and the said Act and Rules made thereunder and with such further requirements, if any, as may be specified in the rules amended from time to time.

Special attention should be given towards prohibition of Advertisements of Certain Drugs for treatment of certain disease conditions as provided under the Schedule of the Drugs and Magic Remedies (Objectionable Advertisements) Act 1954 and Rules made thereunder.

In the event of any objection received from any source and/or any deviation detected at any point of time, action shall follow as per Rules.

Receipt of this order along with certificate in form 25-D, Certificate of GMP in form 26-E-I may please be acknowledged.

## List of Enclosures

1. Drug Licence in form 25 D
2. Certificate of GMP in form 26 E-I

Yours faithfully,

*[Signature]*  
DIRECTOR

(Licensing Authority for ASU Drugs)

*Forwarded to Secretary for information 2.03.22*

*22/03/2022*

*Forwarded to Pharmacy Superintendent 2.03.22*





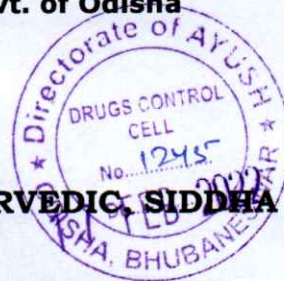
# Directorate of AYUSH

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Department of Health & Family Welfare, Govt. of Odisha  
(DRUGS CONTROLL CELL)

FORM 25-D

(See Rule -154)



## LICENCE TO MANUFACTURE FOR SALE OF AYURVEDIC, SIDDEHA OR UNANI DRUGS

No. of Licence and date of issue – OR- 82/AYUR, dt. 17<sup>th</sup> May 1988

1. **Nagarjuna Rasashala of Sri Sri Nrusinghanath Ayurveda College And Research Institute** is/are hereby licensed to manufacture the following Ayurvedic, Siddha or Unani drugs on the premises situated At – Nrusinghanath, P.O.- Paikmal, Dist – Bargarh, PIN – 768039 in the State of Odisha, under the direction and supervision of the following technical staff :-

**a) Technical staff**

- |   |                         |
|---|-------------------------|
| i) Dr. Akhilesh Sahu, MD (Ay.), Regd. No. -6319 | Manufacturing & Testing |
| ii) Sri Jashobanta Sahu, BSc (Chemistry)        | For Testing             |
| iii) Sri Prem Ranjan Majhi, B.Sc.(Botany)       | For Testing             |

**b) Name of drugs** (each item to be separately specified)

Total no. of Items is **60** (Sixty) only [60 Shastriya/ Classical] as approved earlier.

2. The licence shall be in force from **01 . 01 . 2022**

3. The licence is subject to the conditions stated below and to such other conditions as may be specified in the rules for the time being in force under the Drugs and Cosmetics Act, 1940.

Date of issue:- **17 FEB 2022**  
Bhubaneswar

**DIRECTOR**  
(Licensing Authority)

**Director AYUSH**  
**Odisha, Bhubaneswar**

### Conditions of Licence

1. Any change in the Technical staff named in the licence shall be forthwith reported to the Licensing Authority.
2. This licence shall be deemed to extend to such additional items as the licensee may intimate to the Licensing Authority from time to time, and as may be endorsed by the Licensing Authority.
3. The licensee shall inform the Licensing Authority in writing in the event of any change in the constitution of the firm operating under the licence. Where any change in the constitution of the firm takes place, the current licence shall be deemed to be valid for a maximum period of three months from the date on which the change takes place unless, in the meantime, a fresh licence has been taken from the Licensing Authority in the name of the firm with the changed constitution.
4. The licence unless sooner suspended or cancelled shall remain valid perpetually. However, the compliance with the conditions of licence and the provisions of the Drugs and Cosmetics Act 1940 (23 of 1940) and the Drugs Rules, 1945 shall be assessed not less than once in five years or as needed as per risk based approach.
5. The licence is issued only after fulfilment of the requirements of Good Manufacturing Practices (GMP) of Ayurveda, Siddha or Unani drugs as laid down in Schedule T of the Drugs Rules, 1945."





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Department of Health & Family Welfare, Govt. of Odisha

(DRUGS CONTROLL CELL)

**FORM 26 - E - I**

(See Rule -155 - B / 157)



### **CERTIFICATE OF GOOD MANUFACTURING PRACTICES (GMP) TO MANUFACTURER OF AYURVEDA, SIDDHA OR UNANI DRUGS**

Certified that manufacturing unit licensee, namely **Nagarjuna Rasashala** of **Sri Sri Nrusinghanath Ayurveda College and Research Institute**, At - Nrusinghanath, P.O.- Paikmal, Dist - Bargarh, PIN - 768039, Licence no. OR - 82/ Ayur comply with the requirements of Good Manufacturing Practices (GMP) of Ayurveda Drugs as laid down in Schedule - T of the Drugs and Cosmetics Rules, 1945.

This certificate is valid till **31 - 12 - 2026** and the Good Manufacturing Practices (GMP) is valid for the various dosage forms or Rasausadhis as follows:

**(60 items as approved earlier)**

Bhubaneswar

Dated : **17 FEB 2022**

**DIRECTOR**

(State Licensing Authority for ASU Drugs)

**Director AYUSH**  
**Odisha, Bhubaneswar**