

## Certificate of Analysis

|            |                                              |              |           |
|------------|----------------------------------------------|--------------|-----------|
| Product:   | <b>METHIMAZOLE</b> (USP compendial name)     |              |           |
|            | <b>THIAMAZOLE</b> (Ph. Eur. compendial name) |              |           |
| CAS-No.:   | 60-56-0                                      | Ident.-No.:  | M 0400    |
| Batch-No.: | 0202019P                                     | Retest Date: | 05 / 2024 |

### *Additional tests according to Ph. Eur. Thiamazole monograph.*

| <u>Testing</u>         | <u>Requirement</u>                                       | <u>Result</u> |
|------------------------|----------------------------------------------------------|---------------|
| Melting point          | 143 - 146° C                                             | 145.5° C      |
| Appearance of solution | a) clear solution                                        | conforms      |
|                        | b) not more intensely colored than reference solution B6 | conforms      |
| Heavy metals           | nmt. 10 ppm                                              | passed        |

**Remark:** Product is in conformity with the requirements of the current USP and European Pharmacopoeia. The batch was manufactured, packed and tested at the above mentioned site. Batch records have been reviewed for accuracy, completeness, and compliance with established procedures, to determine compliance of the API with the registered manufacturing process, specifications and cGMP requirements.

Manufacture: May 21, 2019

QA release: August 27, 2019

QC release: August 27, 2019

Issue: October 28, 2019

Q-Manager:

*i.v. [Signature]*

Head of QC:

*A.N. [Signature]*

The above information is derived from our quality checks. It does not relieve the purchaser from examining the product upon delivery and gives no assurance of suitability of the product for any particular purpose.

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**Appearance:** A white to pale buff crystalline powder with a faint characteristic odour.

| <u>Testing</u>                                                 | <u>Requirement</u>                  | <u>Result</u> |
|----------------------------------------------------------------|-------------------------------------|---------------|
| <b>Identification</b>                                          |                                     |               |
| <b>IR</b>                                                      | identical versus reference spectrum | identical     |
| <b>Assay</b>                                                   | 98.0 - 101.0 %                      | 100.0 %       |
| <b>Loss on drying</b>                                          | nmt. 0.5 %                          | 0.1 %         |
| <b>Residue on ignition</b>                                     | nmt. 0.1 %                          | 0.0 %         |
| <b>Selenium</b>                                                | nmt. 30 ppm                         | < 30 ppm      |
| <b>Organic impurities (GC)</b>                                 |                                     |               |
| Impurity A                                                     | nmt. 0.1 %                          | < 0.02 % (DL) |
| Impurity B                                                     | nmt. 0.1 %                          | < 0.02 % (DL) |
| Impurity C                                                     | nmt. 0.1 %                          | 0.03 %        |
| Any other impurity                                             | nmt. 0.10 %                         | < 0.02 % (DL) |
| Sum of all impurities                                          | nmt. 0.5 %                          | passed        |
| <b>Ordinary impurities (TLC)</b>                               |                                     |               |
| 1-Methyl-2-(methylsulphonyl)-1 <i>H</i> -imidazolium rhodanide | nmt. 0.1 %                          | < 0.1 % (LoD) |
| <b>Residual solvents</b>                                       | complies with USP <467>             |               |
|                                                                | nmt. 1000 ppm acetone               | 188 ppm       |
|                                                                | nmt. 1000 ppm ethyl acetate         | < 20 ppm      |
|                                                                | nmt. 100 ppm methanol               | < 20 ppm      |

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