

Product/sample name : Midazolam hydrochloridum: 502530
 Client-code : 129
 Project-code : 0104213
 Batch number : 17K21-B05
 Analysis : DAC/NRF 2015/1
 Version : 1

| Test | Test method | Specification | Result | Complies |
|------------------------|--------------|---|--------------------------|----------|
| Characters | Appearance | White or almost white, crystalline powder | White crystalline powder | Yes |
| Solubility | Client spec. | Sparingly soluble, clear after 2 days | Conform | Yes |
| Identity B | 2.2.24 | Compared to reference | Conform | Yes |
| Identity D | 2.3.1 | Reaction of chloride (a) | Conform | Yes |
| Appearance of solution | 2.2.1 | Clear | Clear | Yes |
| | 2.2.2 (II) | $\leq Y6$ | $< Y6$ | Yes |
| pH | 2.2.3 | 3.7 – 4.7 | 4.3 | Yes |
| Related substances | 2.2.29 | Impurity B: ≤ 0.2 % | Not detected | Yes |
| | | Impurity A: ≤ 0.1 % | Not detected | Yes |
| | | Impurity G: ≤ 0.1 % | Not detected | Yes |
| | | Any other imp.: ≤ 0.10 % | 0.09 % | Yes |
| | | Sum: ≤ 0.3 % | 0.1 % | Yes |
| Impurity C | 2.2.27 (TLC) | ≤ 0.1 % | < 0.1 % | Yes |
| Loss on drying | 2.2.32 | < 0.5 % | 0.2 % | Yes |
| Sulphated ash | 2.4.14 | ≤ 0.1 % | < 0.1 % | Yes |
| Assay | Monograph | 98.5 - 101.5 % | 99.4 % | Yes |
| Endotoxin | 2.6.14 | ≤ 0.25 EU / mg | < 0.05 EU / mg | Yes |

QA-statement:

This study met the criteria for a valid test and was performed in compliance with the Good control Laboratory Practice as defined in the Guide to Good Manufacturing Practice for Medicinal Products in the European Community.

The reported results adequately reflect the raw data of the study.

Conclusion:

The results do comply with the specifications

This certificate is approved by Manager QA on 18 December 2017

Oath 18dec17