**CERTIFICATE OF ANALYSIS**

**CERTIFICATE No:** CAN/2015-16/509

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| **PRODUCT** | TIGERON 750 mg TABLETS | | REF. NO: NDQD201511538 |
| **DATE RECEIVED:** 10.11.2015 | **LABEL CLAIM:** | Each film coated tablet contains; Levofloxacin hemihydrate equivalent to Levofloxacin 750 mg. | |
| **BATCH NO.:** TC5002 | **PRESENTATION:** | Pink coloured, caplet shaped, biconvex faced tablets, embossed '750' on one face and plain on the other, packed in a blister strip of 10 tablets in a unit box. | |
| **MGF. DATE:** Apr.2015 | **MANUFACTURER:** | KUSUM Healthcare Pvt. Ltd. | |
| **EXP. DATE:**  Mar.2018 | **ADDRESS:** | SP 289 (A), RIICO, Indl. Area, Chopanki, Bhiwadi (Rajasthan), INDIA. | |
| **CLIENT REF NO:** | **CLIENT:** | Highridge Pharmaceuticals Ltd. P. O. Box 32982 - 00600, Nairobi, KENYA. | |
|  | **TEST(S) REQUESTED:** | Uniformity of Weight, Identification, Dissolution, Assay | |

**RESULTS**

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| **TEST** | **METHOD** | **COMPEDIA** | **SPECIFICATION** | **DETERMINED** | **REMARKS** |
| **Uniformity of Weight** | Weight | BP 2016 Vol. V App. XII C. | NMT 2 tablets deviate by more than 5% from mean tablet weight | None Deviates | **COMPLIES** |
| **Identification** | HPLC | Adopted In-House Method | RT of the major peak in the assay sample preparation corresponds to that in the assay standard preparation | Super-imposable peak at RT 8.7 -/+ 10% min. present in both the assay sample and standard preparations | **COMPLIES** |
| **Dissolution** | UV | Adopted In-House Method | No tablet less than 85.0% [n=6] | 102.3% (RSD=0.2%; n=6) | **COMPLIES** |
| **Assay** | HPLC | Adopted In-House Method | 95.0 - 105.0% | 99.4% (RSD=0.3%; n=6) | **COMPLIES** |

**CONCLUSION:** The product complies with the specifications for the tests performed.

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| ANALYST: | DR. E. MUTUA |  | DATE:31-03-2016 |
| ANALYST: | DR. S. MUTERU |  | DATE:31-03-2016 |
| ANALYST: | ERNEST MBAE |  | DATE:04-04-2016 |
| DIRECTOR: | HEZEKIAH CHEPKWONY |  | DATE:04-14-2016 |