



Bio-Rad
Laboratories

Diagnostics Group
9500 Jeronimo Road
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CERTIFICATE-OF-ANALYSIS

Product Name: Liquichek Hematology-16 Control

Master Batch Number: 82260

Product Base: Whole Blood

Physical Form: Liquid

Date of Manufacture: 2023-11-13

| Description | Material Number | Batch Number | Expiration Date |
|------------------|-----------------|--------------|-----------------|
| Trilevel | 760 | 82260 | 2024-04-21 |
| Low | 761 | 82261 | 2024-04-21 |
| Normal | 762 | 82262 | 2024-04-21 |
| High | 763 | 82263 | 2024-04-21 |
| Trilevel MiniPak | 760X | 82260 | 2024-04-21 |

Testing Results:

Each human donor unit used to manufacture this product was tested and found non-reactive at the donor level per current applicable FDA requirements using FDA-accepted methods including testing for HIV-1/HIV-2 antibody, HBsAg, HCV Antibody, HIV-1 and HCV by a nucleic acid test (NAT).


Homogeneity Claim:

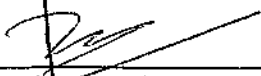
"Testing has been conducted to verify sufficient homogeneity in accordance with established requirements."

ADDITIONAL TESTING:

This product has been manufactured under applicable guidelines/standards and meets all established Bio-Rad quality requirements.

Approved By:


(QC Supervisor/Manager)


(QA Supervisor/Manager)

Date:

11-14-2023

11/14/2023

