

Patient Name : Mr MR.DUMMY
DOB/Age/Gender : 23 Y/Male
Patient ID / UHID : 8052379/RCL7248930
Referred By : Dr. Dr. X
Sample Type : Serum

Sample Collected : Apr 26, 2024, 01:00 PM
Report Date : May 25, 2024, 12:32 PM.
Barcode No : SI484720
Report Status : Final Report

Test Description	Value(s)	Unit(s)	Reference Range
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HTLV I & II Total Antibody

HTLV I & II Total Antibody (Serum, CMIA)	0.66	S/CO	Non reactive: < 1.0 Reactive: >= 1.0
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Interpretation:

Human T cell lymphotropic virus type I & II (HTLV 1 & II) antibodies indicate infection due to these viruses, which belong to the Retroviridae family.

HTLV I : Infection may progress to infective dermatitis, uveitis syndrome, HTLV I associated myelopathy (HAM), Adult T cell Leukemia/ Lymphoma (ATL).

HTLV II: Plays a key role in certain neurological, hematological and dermatological diseases. Association with Hairy cell Leukemia is not clearly established. Presence of these antibodies (HTLV I & II) indicate infection and the individual is considered unfit to donate blood.

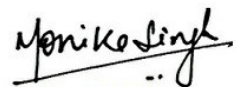
*** End Of Report ***

Disclaimer: Method given in report are only indicative and can be changed depending upon type of machine and kit available at time of testing.

Not all tests at all locations are under NABL scope. Availability of tests under NABL scope varies from lab to lab.



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Booking Centre :- DEMO PARTNER CHENNAI, DEMO PARTNER CHENNAI
Processing Lab :-

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