





ISO 9001:2015

ISO <u>13485:</u>2016

HCV ELISA 3rd Gen

PRODUCT FEATURES



- Accurate

- The test kit has a high sensitivity rate, typically over 99.5%.
- High Specificity, accurately distinguish HCV antigens from other proteins.
- Validated through extensive clinical trials.



User Friendly

- Self Contained kit requires minimum equipment.
- Requires minimal technical expertise.
- kit includes clear, step-by-step instructions.



Reliable Result

- Delivers accurate results across multiple and diverse conditions.
- Quality control features that help verify the correct functioning of the kit.
- Designed to withstand typical handling and environmental factors.

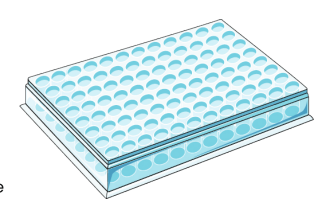
HCV ELISA 4th Gen

Intended Use

Qualitative detection of antibodies against HCV in human serum or plasma, based on 3rd Genetarion ELISA technique.

Salient Features

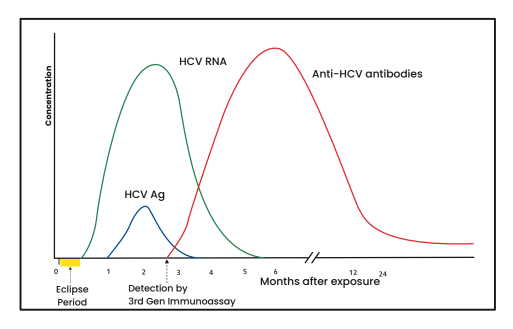
- Detection of all genotypes of HCV
- Microplates coated with Core, NS3, Ns4, Ns5 Ag of HCV
- Convenient microwell sample dilution
- Sensitivity 100% Specificity > 99.5%
- Duration of assay 90 mins + process time



Ordering Information

96 Tests

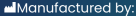
PT/MVM/100



Performace Characteristics

The performance of 3rd generation HCV Prolisa with reference to sensitivity and specificity has been determined by National Institute of Biologicals, Noida. The samples included in the panels comprises of confirmed HCV strong positives and weak positives and confirmed HCV negative serum/plasma.





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