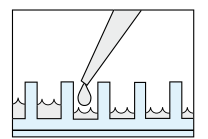
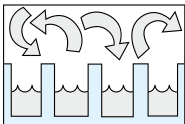
## **AVIOQ® HTLV-I/II**

## Microelisa System

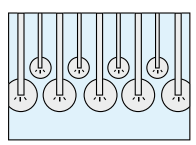
**Test Procedure** 



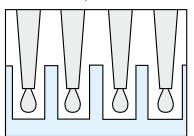
Dilute samples and controls with Sample Diluent in accordance with package insert. Add to HTLV coated microplate.



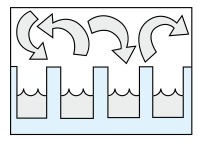
2 Cover the strips with adhesive plate sealer or equivalent. Incubate for 60 ± 5 minutes at 37°C ± 2°C.



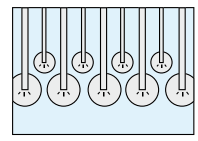
Aspirate and wash each well four times with diluted Wash Buffer. Use a soak cycle of at least 30+/-5 seconds.



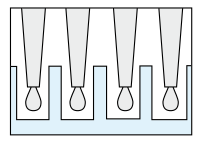
Add 100µl of reconstituted EnzAbody working solution into each well.



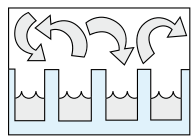
5 Cover the strips with a new plate sealer or equivalent and incubate for  $60 \pm 5$  minutes at  $37^{\circ}C \pm 2^{\circ}C$ .



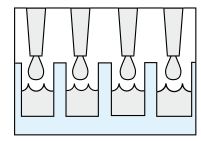
Aspirate and wash each well four times with diluted Wash Buffer. Use a soak cycle of at least 30+/-5 seconds.



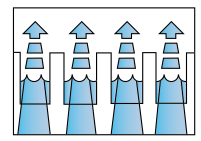
Add 100μl of TMB Substrate Solution into each well. Do not mix. Do not cover.



8 Incubate for 30 ± 5 minutes at room temperature (15°- 30°C). Do not cover.



Add 100µl of 2N Sulfuric Acid into each well.



Read absorbance at 450 nm ± 5 nm or 450 ± 5 nm and 620/630 nm ± 5 nm as reference.



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See package insert for additional information.