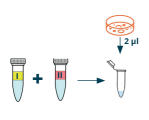


Quick Start Guide

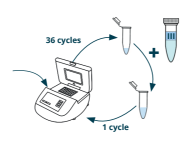
See **Instructions For Use (IFU)** included in kit for full details

Reaction Setup



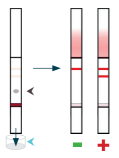
1. Mix 5 µl **Solution I** and 5 µl **Solution II** 2. Add 2 µl of cell culture supernatant

Reaction



- 3. Run the 36 cycle thermal cycler programme*
- 4. Add 10 µl **Solution III** and place back in the cycler for a final cycle*

Run Sample on Strip



- 5. Mix 20 µl of **Running Buffer** with the PCR reaction
- 6. Add 5 µl of this mixture to the **Test Strip** ≺
- 7. Place the **Test Strip** in a well or tube with 300 µl of **Running Buffer** \blacktriangleleft
- 8. Wait 10 minutes for the test to run and then read the result

^{*}See the IFU for full thermal cycler programme details



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