TwistAmp® Basic Quick Guide

Part Number: TABASo3Guide | Revision E

Basic Information

RPA

- 1) Primers can be 18-35 bases1
- 2) Works best at constant temperature (37-39°C)
- a) Amplicons of 80-400bp are preferred

Set-up (single-plex)2

1) Prepare reaction mix in 1.5ml tube:

 Primer A (10μM)
 2.4 μl

 Primer B (10μM)
 2.4 μl

 Rehydration Buffer
 29.5 μl

Template and dH2O 13.2 μl (Total Volume 47.5 μl)

Vortex and spin briefly

- Add reaction mix to freeze-dried reaction. Pipette to mix.
- 3) Add 2.5 µl of 280mM MgAc (supplied) and mix well to start reaction.

WARNING: RPA REACTIONS START AT ROOM TEMPERATURE AS SOON AS MAGNESIUM IS ADDED.

PCR

- 1) Primers typically 18-25 bases
- 2) Thermal cycling required
- Amplicons of 50bp upwards are typical/optimal
- 4) Incubate at 37-39°C for 20-40 minutes.
- 4b) For low template copy number, remove strip after 4 minutes, vortex & spin briefly, replace in heating device.
- 5) After 20-40 minutes, clean amplicons before running on agarose gels.

WARNING: IF TUBES ARE OPENED AFTER AMPLIFICATION THERE IS A GREAT RISK OF CONTAMINATION OF WORK SURFACES WITH AMPLICON! ENSURE THAT APPROPRIATE AVOIDANCE MEASURES ARE TAKEN!

WARNING: SWITCH OFF HEATED LIDS BEFORE STARTING REACTIONS!

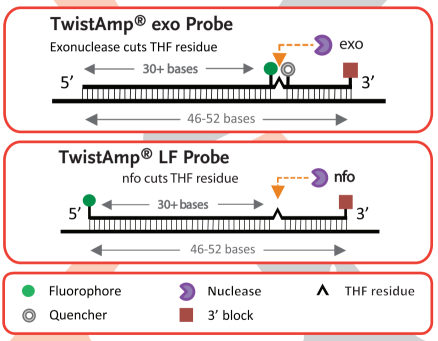
- 1 For rapid amplification 30-35 bases are optimal
- ² See manual for multiplexing



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RPA uses TwistDx's proprietary probe systems

RPA does NOT use PCR probe systems



refer to manual for details of probe design



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