highQu professionally simple

Check the product label for actual catalog number, lot and expiry date.

ALLin™ RPH Polymerase

CAT.#	SIZE	COMPONENTS	COMPONENT COMPOSITION
HLE0101	250 u	250 u - ALLin™ RPH Polymerase, 5 u/μl 2 x 1 ml - 5X ALLin™ RPH Buffer	Enzyme in storage buffer. 1X ALLin™ RPH Buffer contains 0.25 mM dNTPs, 3 mM MgCl₂, enhancers, stabilizers.
HLE0105	1250 u	5 x 250 u - ALLin™ RPH Polymerase, 5 u/µl 10 x 1 ml - 5X ALLin™ RPH Buffer	Enzyme in storage buffer. 1X ALLin™ RPH Buffer contains 0.25 mM dNTPs, 3 mM MgCl ₂ , enhancers, stabilizers.
Storage	In the dark at -20°C.		

APPLICATIONS

- Amplification of difficult & complex (GC/AT rich) templates
- Long PCR (up to 35 kb) with higher fidelity
- Colony & crude sample PCR
- Multiplex PCR
- TA cloning

PRODUCT DETAILS

highQu ALLin™ RPH Polymerase (Robust, Proofreading, Hot-start Polymerase) is the versatile engineered enzyme combining best polymerase properties for excellence in most demanding PCR applications, like low copy detection, long or high fidelity PCR, amplification of complex templates, crude sample PCR and multiplexing.

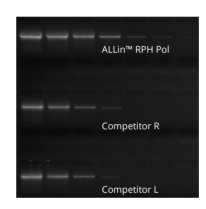
ALLin™ RPH Polymerase has 5 times higher fidelity than Taq DNA Polymerase and produces A-tailed products suitable for ligating into TA cloning vectors.

For the maximum convenience the 2X ALLin™ RPH Polymerase mastermix (HLM0101) is available.

BENEFITS

- RPH Robust, Proofreading, Hot-start Polymerase
- Low-copy number target detection ensured by small molecular inhibitor hot-start
- Long (up to 35 kb) high-fidelity (5X higher than Taq) amplification ensured by proofreading activity
- High yields under standard and fast cycling
- Robust amplification of GC or AT rich templates, crude samples
- 5X ALLin™ PCR Buffer contains optimal Mg²⁺ and dNTPs

PERFORMANCE



✓ Prepare a 50 µl reaction:

ALLin™ RPH
Polymerase ensures
higest sensitivity
amplification of 25 kb
target from lowest
amounts of human
genomic DNA.
The starting template
concentration is 200
ng of human genomic
DNA with 2x further
dilutions. 25kb
fragment of the p53
gene was amplified.

PROTOCOL

- Take typical measures to prevent PCR cross over contamination, keep your bench clean, wear gloves, use sterile tubes and filter pipet tips.
- Include a no-template control and positive control in parallel.
- Thaw and keep reagents on ice. Mix well before use.
- The longer the amplicon, the longer the extension time: Use 15 sec/kb extension for amplicons of <5 kb.
- $\bullet \quad$ Use 40-60 sec/kb extension for amplicons of 5-35 kb.
- Use 90 sec extension for multiplexing.
- Run an annealing temperature gradient from 55°C to 65°C to choose the best specificity conditions.
- Do not use fast cycling for multiplexing.

IN VITRO RESEARCH USE ONLY

Rev. & For. Primers	0.1-0.4 μM final each (≤ 2 μl of 10 μM)	
cDNA Template or	<100 ng or	
gDNA Template	5-500 ng	
5X ALLin™ RPH	10 μΙ	
Buffer		
Water (PCR Water	to 49 µl	
WAT0110)		
ALLin™ RPH	0.25 - 1 μl	
Polymerase, 5 u/μl		
Mix gently, avoid bubbles.		
✓ Place into the inst	trument set like:	
Initial denaturation	1 cycle: 95°C - 1 min	
Denaturation	25-35 cycles: 95°C - 15 sec	
Annealing	25-35 cycles: 55-65°C – 15 sec	
Extension	25-35 cycles: 72°C - ~10 min (for 10 kb)	
	15 sec/kb for <5 kb amplicons	
	40-60 sec/kb for >5 kb amplicons	
✓ Store probes for	short time on ice, for long at -20°C.	

ORDERING

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