

## Xpert AmpliFi Mastermix

#GE27.0001 (1.25ml) | GE27.5001 (5x 1.25ml) | GE27s (trial size)  
(FOR RESEARCH ONLY)



**Product:** Xpert AmpliFi DNA Polymerase is an all-round and robust high-fidelity enzyme with enhanced DNA binding, resulting in improved processivity, shorter extension times, higher yield and capacity to amplify more difficult and longer targets. Combined with its extremely low error-rate, this makes Xpert AmpliFi ideal for applications such as high-fidelity PCR, site-directed mutagenesis, cloning, long-range PCR and DNA sequencing, when robustness and proof-reading are extremely important.

Xpert AmpliFi DNA polymerase has been derived from Pfu DNA polymerase for its proof-reading activity. Xpert AmpliFi has a very high accuracy resulting in fidelity that is about 100x higher than regular Taq DNA polymerase. DNA binding has been improved by protein engineering resulting in shorter extension times (2-6kb/min), making this also an ideal choice for consistent results in fast PCR amplifications.

**Applications:** High-Fidelity PCR, cloning, site-directed mutagenesis. Long-Range PCR, DNA sequencing

**Contents:** Xpert AmpliFi DNA Polymerase is supplied in vials of 1.25ml of a convenient 2X-MasterMix comprising 6mM MgCl<sub>2</sub>, 2mM dNTPs, enhancers and stabilizers, optimized to increased PCR success rates. Each vial is sufficient for 100 PCR reactions of 25µl. #GE27s is a trial sample (0.5ml), sufficient for 30 reactions.

Component	GE27.0001	GE27.5001	GE27s
Xpert AmpliFi Mastermix (2X)	1.25ml	5x1.25ml	0.5ml

**QC:** Functionally tested in PCR. Absence of endonucleases, exonucleases, and ribonucleases was confirmed by appropriate assays

**Properties:** Amplicon size: up to 20kb  
Extension rate: 2-6kb/min  
Hotstart: No  
A-overhang: No  
Error-rate: 100x lower than normal Taq

**Storage:** -20°C and protected from light for at least 1 year. No loss of performance is detected after 20 freeze/thaw cycles.

### Prior to use

Optimal PCR cycling conditions (incubation times, temperatures, and concentrations) depend on DNA target (GC-content, size, quantity, purity, etc.), specific primers, dNTPs, buffer composition and MgCl<sub>2</sub>-concentration, and need to be determined case by case. Xpert AmpliFi MasterMix already includes dNTPs and has been already been optimized with respect to the MgCl<sub>2</sub> concentration and other components to maximize success rates. Thus, it is not recommended to add additional MgCl<sub>2</sub> or dNTPs. We suggest starting with the following basic protocol and subsequently optimizing annealing temperature, incubation times and cycling number.

## Basic Protocol

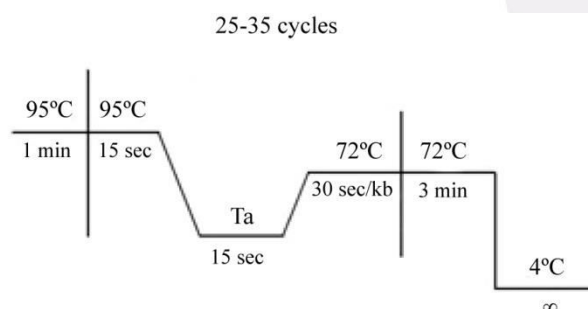
1. Mix for each PCR reaction:

Component	Volume (25 $\mu$ l)	Final Conc.
Xpert AmpliFi Mastermix (2x)	12.5 $\mu$ l	1X
Forward primer (5 pmol/ $\mu$ l)	2 $\mu$ l	0.4 $\mu$ M
Reverse primer (5 pmol/ $\mu$ l)	2 $\mu$ l	0.4 $\mu$ M
Template DNA*	0.5 – 5.0 $\mu$ l	1-100ng*
PCR-grade water	up to 25 $\mu$ l	

*\*) In case of gDNA <100ng (total amount) and in case of complex DNA <5ng  
For smaller/larger reaction volumes, scale it down/up proportionally.*

In order to minimize risk of contamination, reagent loss and improve pipetting accuracy, we recommend to prepare a mastermix for multiple samples (N), always including a negative control for the detection of possible contaminants, by mixing all components (N+1), except template DNA, dividing the mixture equally into each tube and then add template DNA or PCR grade water in case of the control to the individual PCR tubes.

2. Set-up initial PCR amplification as follows:



After an initial cycle of 1 min at 95°C (denaturation of template DNA including removal of all secondary DNA structures such as hairpins), cycle 25-35 times for 15 seconds at 95°C, 15 seconds at  $T_a$ , and 30 seconds per kb of target DNA at 72°C for extension. Set the annealing temperature ( $T_a$ ) as the melting temperature ( $T_m$ ) of the primer with the lowest  $T_m$ . After amplification, include a final extension step of 3 min at 72°C. Analyze PCR products by DNA Agarose gel electrophoresis.

## Optimization

### Denaturation

If amplification of GC-rich templates results in low yield, increasing temperature to 98°C-100°C may improve amount of product.

### Annealing Temperature ( $T_a$ ) and Primers

Optimizing the annealing temperature is crucial, as a too low temperature might result in non-specific amplification whereas a too high temperature results in no amplification. The melting temperature ( $T_m$ ) is defined as the temperature in which 50% of the primer and its complementary sequence of the target DNA are present as duplex DNA. By increasing the temperature above the melting temperature, this percentage decreases, however, primers will still anneal (up to a certain point) and initiate extension. PCR can therefore be performed at temperatures of several degrees higher than  $T_m$  and it is therefore recommended to optimize the  $T_a$  by performing a temperature gradient (e.g. starting at the lowest  $T_m$  and increasing with 2°C increments). Ideally, primers have melting temperatures of approximately 60°C and final concentration should be between 0.2 and 0.6 $\mu$ M (each).

### Incubation times.

Denaturation and annealing steps may require less time depending on the thermocycler apparatus, reaction volume and PCR tube (varies with the efficiency of heat-transfer). It might be worthwhile to optimize (reduce) times to as low as 10 seconds for both denaturation and annealing steps, which will greatly reduce overall PCR time. Optimal extension is carried out with a velocity of 2kb/min but may be reduced to as little as 6kb/min for less demanding templates.