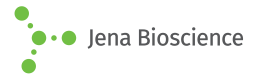
DATA SHEET

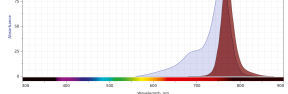




HighFidelity IR750 PCR Labeling Kit

Preparation of IR750-labeled DNA probes by PCR

Cat. No.	Amount		
APP-101-IR750-S	10 reactions x 20 μl		
APP-101-IR750-L	50 reactions x 20 μl		
100			
75 -			
8			



excitation and emission spectrum of IR750

For general laboratory use.

Shipping: shipped on gel packs

Storage Conditions: store at -20 °C

Additional Storage Conditions: avoid freeze/thaw cycles, store dark

Shelf Life: 12 months

Spectroscopic Properties: λ_{exc} 753 nm, λ_{em} 775 nm, ϵ 255.0 L mmol⁻¹ cm⁻¹ (Tris-HCl pH 7.5)

Description:

HighFidelity IR750 PCR Labeling Kit is designed to produce randomly IR750-modified DNA probes by PCR. Such probes are ideally suited for Fluorescence *in situ* hybridization (FISH) and Northern Blot experiments. PCR-based labeling is superior to random-primed labeling with Klenow fragment if template amounts are limited or amplification of a specific DNA fragments is required. Amplification of probes up to 4kbp is feasible.

dUTP-XX-IR750 is efficiently incorporated into DNA as substitute for its natural counterpart dTTP using an optimized reaction buffer and a High Fidelity Polymerase blend consisting of *Taq* polymerase and a proofreading enzyme. 25 % dUTP-XX-IR750 substitution typically results in an optimal balance between reaction and labeling efficiency. Individual optimization of dUTP-XX-IR750/dTTP ratio however, can easily be achieved with the single nucleotide format.

The kit contains sufficient reagents for 10 labeling reactions (S-Pack) or 50 labeling reactions (L-Pack) of 20 μl each (25% dUTP-XX-IR750 substitution, 100 μM dATP/dGTP/dCTP, 75 μM dTTP, 25 μM dUTP-XX-IR750).

Content: High Fidelity Polymerase

in storage buffer with 50% glycerol (v/v) #APP-101-IR750-S: 1x 40 µl (100 units, 2.5 units/µl) #APP-101-IR750-L: 2x 40 µl (2x 100 units, 2.5 units/µl)

High Fidelity Labeling Buffer 1x 500 µl (10x)

dATP - Solution 1x 20 μl (100 mM)

dGTP - Solution 1x 20 μl (100 mM)

dCTP - Solution

1x 20 µl (100 mM)

dTTP - Solution

1x 20 µl (100 mM)

dUTP-XX-IR750

#APP-101-IR750-S: 1x 10 μl (1 mM) #APP-101-IR750-L: 5x 10 μl (1 mM)

Lambda DNA

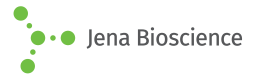
1x 20 µl (100 ng/µl)

500 bp forward primer 1x 20 μl (10 $\mu M)$



Jena Bioscience GmbH Löbstedter Str. 71 | 07749 Jena, Germany | Tel.:+49-3641-6285 000 https://www.jenabioscience.com

DATA SHEET





HighFidelity IR750 PCR Labeling Kit

Preparation of IR750-labeled DNA probes by PCR

500 bp reverse primer 1x 20 μl (10 μM)

PCR-grade water 1x 1.2 ml

To be provided by user DNA template

Primer DNA purification tools (optional)

1. Preparation of working solutions

1.1 Preparation of 1 mM dATP/dCTP/dGTP working solution

- Thaw 100 mM dATP, 100 mM dCTP and 100 mM dGTP solutions on ice, voretex and spin-down briefly.
- Prepare a 1:100 dilution with PCR-grade water to achieve a final concentration of 1 mM (e.g. 2 µl 100 mM dATP + 2 µl 100 mM dCTP + 2 µl 100 mM dGTP + 194 µl PCR-grade water).
- 1 mM ATP/CTP/GTP working solution can be stored at -20°C. Prepare aliquots to avoid freeze/thaw cycles.

1.2 Preparation of 1 mM dTTP working solution

- Thaw 100 mM dTTP solution on ice, voretex and spin-down briefly.
- Prepare a 1:100 dilution with PCR-grade water to achieve a final concentration of 1 mM (e.g. 2 μl 100 mM dTTP + 198 μl PCR-grade water).
- 1 mM dTTP working solution can be stored at -20 °C. Prepare aliquots to avoid freeze/thaw cycles.

3. Standard PCR Labeling protocol

The standard protocol is set-up for labeling of a 500 bp DNA fragment. An optimal balance between reaction and labeling efficiency is typically achieved with 25% dUTP-XX-IR750 substitution following the standard protocol below however, individual optimization might improve results for individual applications.

- Assemble the PCR on ice in the order stated below (DNAse-free reaction tube).
- Voretex and spin-down briefly.
- Perform assay set-up and reaction under low-light conditions.

Component	Volume	Final concenctra- tion		
PCR-grade water	Xμl			
High Fidelity La- beling Buffer (10x)	2 µl	1x		
1 mM dATP/dCTP/ dGTP working so- lution (s. 1.1)	2 µl	100 µM		
1 mM dTTP working solution (s. 1.2)	1.5 µl	75 μΜ		
1 mM dUTP-XX- IR750	0.5 μl	25 μΜ		
forward primer (10 μM)	Χ μί	0.1 - 1 μM (e.g. 0.3 μM 500 bp forward primer)		
reverse primer (10 µM)	ΧμΙ	0.1 - 1 μM (e.g. 0.3 μM 500 bp reverse primer)		
template DNA	ΧμΙ	1 - 10 ng genomic DNA (e.g. 1 ng Lambda DNA)		
High Fidelity Polymerase (2.5 units/µl)	1 μl	2.5 units		
Total volume	20 µl			

Recommended cycling conditions

Cycle step	Temperature	Time	Cycles
Initial denaturation	95°C	2 min	1x
Denaturation Annealing ¹⁾ Elongation ²⁾	95°C 58°C 68°C	20 sec 30 sec 60 sec	30x
Final Elongation	68°C	2 min	1x

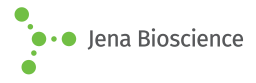
¹⁾The annealing temperature depends on the melting temperature of primers used.

²⁾The elongation time depends on the length of fragments to be amplified. A time of 2 min/kbp is recommended. Elongation at 72°C works as well.

For optimal amplification results and high incorporation rates an individual optimization of the recommended PCR assay and cycling conditions may be necessary for each new primer-template pair.









HighFidelity IR750 PCR Labeling Kit

Preparation of IR750-labeled DNA probes by PCR

4. Probe purification:

Probe purification is not required for most hybridization experiments. If a downstream application requires purification (e.g. concentration determination by absorbance measurement) we recommend silica-membrane or gel filtration-based purification.

Related Products:

Aminoallyl-dUTP-XX-IR750, #NU-803-XX-IR750

