

## WARRANTY

Products are guaranteed to conform to the quality and content indicated on each vial and external labels during their shelf life. BIOTOOLS obligation and purchaser's rights under this warranty are limited to the replacement by BIOTOOLS of any product that is shown defective in fabrication, and that must be returned to BIOTOOLS, freight prepaid, or at BIOTOOLS' option, replacement of the purchasing price. Any complaint on damaged goods during transport must be directed to the handling or transport agent.

Product for Research Use Only. This product must be used by qualified professionals only. It is the responsibility of the user to ascertain that a given product is adequate for a given application. Any product not fulfilling the specifications included in the product sheet will be replaced. This warranty limits our responsibility to the replacement of the product. No other warranties, of any kind, express or implied, including, without limitation, implicit warranties of commercialisation ability or adequacy for a given purpose, are provided by BIOTOOLS. BIOTOOLS will not be held responsible for any direct, indirect, consequential or incidental damage resulting of the use, misuses, results of the use or inability to use any product.

### Produced by:

BIOTOOLS, Biotechnological & Medical Laboratories, S.A. have been evaluated and certified to accomplish ISO 9001:2000 requirements for the following activities: Research and development of biotechnology products and manufacture of biotechnology and in vitro products. Valle de Tobalina - 52 - Nave 39, 28021 Madrid - Spain

© 2008 BIOTOOLS, Biotechnological & Medical Laboratories, S.A. All rights reserved



BIOTOOLS B&M Labs, S.A.  
Valle de Tobalina - 52 - Nave 39  
28021 Madrid  
Spain

Tel. (34) 91 710 00 74

Fax (34) 91 505 31 18

E-mail: [info@biotools.eu](mailto:info@biotools.eu)

[www.biotools.eu](http://www.biotools.eu)



**BIOTOOLS**  
BIOTOOLS B & M LABS. S.A.

## QUANTIMIX HOTSPLIT EASY KIT

**Kit for Real Time DNA Amplification and Quantification to use with Intercalating Fluorophores.**

*(Includes Biotools HotSplit DNA Polymerase)*

Ref.	FORMAT	CONTENT
10.691	100 rxn	Quantimix HotSplit Easy Kit
10.692	200 rxn	Quantimix HotSplit Easy Kit
10.693	500 rxn	Quantimix HotSplit Easy Kit

**Store at -20°C**

**Research Use Only. Not for use in diagnosis procedures**

**Notice to users:** Some of the applications which may be performed with this product are covered by applicable patents in certain countries. The purchase of this product does not include or provide a license to perform patented applications. Users may be required to obtain a license depending on the country and/or application.

Ed. 02 -February 2011

## 1. DESCRIPTION

QUANTIMIX HOTSPLIT EASY KIT has been optimised to deliver maximum efficiency, precision, and sensitivity during nucleic acid amplification in real time using intercalating fluorophores. The kit includes a chemically modified version of Biotools DNA Polymerase, Biotools HotSplit DNA Polymerase, which presents little or no activity at low temperatures and provides greater specificity and yield in DNA amplifications.

Real time detection of amplified products is performed by monitoring the increase of fluorescence after each amplification cycle. In QUANTIMIX HOTSPLIT EASY Kit the fluorescent signal is generated by the incorporation of DNA-binding dyes to the amplified products. In the exponential phase of the amplification there is a correlation between the amount of product and the initial template DNA. Thus, the amount of fluorescence is proportional to the amplified DNA in each cycle.

The Biotools HotSplit DNA Polymerase ensures high specificity and sensitivity. It overcomes the problems due to nonspecific priming, primer-dimer formation or unwanted reactions occurring at low temperatures during the PCR setting, because of the presence of thermolabile blocking groups acting over the amino-acid residues involved in enzyme polymerization. The polymerase activity is restored during the Initial Denaturation Step. On the other hand, Biotools HotSplit DNA Polymerase has a low base misincorporation rate in comparison with similar commercial enzymes.

QUANTIMIX HOTSPLIT EASY KIT is a ready-to-use 2X Master Mix (QUANTISPLIT), which includes all reaction components except intercalating fluorophore, template and primers.

## 2. REAGENTS INCLUDED IN THE KIT

- QUANTISPLIT:** It is a 2X ready-to-use solution that contains all necessary components for real time amplification assays: Biotools HotSplit DNA Polymerase, all four dNTPs, Reaction Buffer and MgCl<sub>2</sub> is also included at the appropriate concentration (4mM).
- 50 mM MgCl<sub>2</sub> Solution:** Used only for specific real time assays which require an additional optimisation.

**The intercalating fluorophore is not provided with the kit**

## 3. STORAGE AND HANDLING INSTRUCTIONS

Store all components of the QUANTIMIX HOTSPLIT EASY Kit at **-20°C** in a constant temperature freezer (frost-free freezers are not recommended). All reagents must be thawed and handled on ice.

**QUANTISPLIT:** Minimise the number of freeze-thaw cycles by storing in working aliquots. Mix well prior to use.

**MgCl<sub>2</sub> Solution:** Mix thoroughly before use.

If stored under the recommended conditions, the product will maintain performance through the indicated date on the label.

## 4. GENERAL CONSIDERATIONS

**Template:** DNA condition is a key point to obtain optimal results. DNA prepared using standard isolation techniques is a suitable substrate for amplification. Nevertheless, numerous compounds inhibit amplification including ionic detergents, some gel loading dyes, phenol and hemin.

The employed methods for DNA purification can be either phenol-based or resin-based, provided they yield enough amount of pure DNA and guarantee the absence of amplification inhibitors. When using purification methods based on a silica matrix, it is important to check the complete absence of silica particles in the sample since it inhibits amplification and fluorescence reading. We recommend the use of our Speedtools line for extraction of genomic DNA from blood (Speedtools DNA Extraction); from tissue (Speedtools Tissue DNA Extraction Kit) from food (Speedtools Food DNA Extraction Kit) and from plant material (Speedtools Plant DNA Extraction Kit).

Samples should be transported and stored frozen. In samples that have been stored without refrigeration, DNA can be degraded. In case of working with clinical samples, handle them as if they are capable of transmitting infectious agents.

The quantity of template DNA to be added in each reaction depends on the source and quality of the template. We recommend determining the concentration by A260/280 measurement. If you do not know the concentration of template DNA, add a fixed volume of the extraction mixture to the problem samples. The purpose of this recommendation is to obtain comparable quantitative results.

**MgCl<sub>2</sub> Concentration:** Magnesium ion concentration affects primer annealing and template denaturation, as well as enzyme activity and fidelity. Generally, excess Mg<sup>2+</sup> results in accumulation of nonspecific amplification products, whereas insufficient Mg<sup>2+</sup> results in reduced yield of the desired PCR product. The recommended range of concentration is 4-6 mM. A 4mM total MgCl<sub>2</sub> concentration is present in the final 1X dilution of the QUANTISPLIT (appropriate concentration). However, the kit is provided with an additional vial with 50 mM MgCl<sub>2</sub> Solution for additional optimisation.

**Primer Design:** Primers typically are 15-30 bases in length and contain approximately 40-60% G+C residues: the annealing temperatures of primer pairs should be nearly identical. Care must be taken to design primers that do not form hairpin loop structures or are self-complementary. The 5' end of a primer may contain mismatches between the primer and template, whereas this is not recommended at the 3' end.

The recommended primer concentration is 0.1-0.5 µM. A final concentration of 0.2 µM for each primer is effective for most reactions.

Unlike regular PCR, amplicons for real time PCR tend to be less than 500 bp. The optimal real-time PCR results can be achieved with amplicon sizes of 100-300 bp; longer products do not amplify as efficiently.

**Cycling Parameters:** Many parameters influence both the specificity and efficiency of amplification including the temperature and duration of denaturation, annealing and elongation; ramp speed; and total cycle number. Due to the shorter length of real time PCR amplicons, cycling programs for real time PCR are typically shorter than regular PCR. Variations to the program may be required depending on the amplicon size.

QUANTIMIX HOTSPLIT EASY Kit contains a chemically modified polymerase which presents little or no activity at low temperature. The full enzyme activity is recovered during the Initial Denaturation Step; 10 min at 96°C should be enough to completely activate the polymerase.

Biotoools HotSplit DNA Polymerase also has a low base misincorporation rate (1-10 x 10-6 bp) and requires more time for extension at 72°C as compared to other polymerases.

The annealing temperature should be 5°C lower than the theoretical melting temperature of the primers. If the annealing temperature is set too low, there is an added risk of primer-dimer extension or nonspecific products.

A **melting curve** is essential when running assays using intercalating fluorophores in order to check the melting profile of PCR products.

## 5. STANDARD PROTOCOL

**Materials to be supplied by user:**

- Intercalating Fluorophore
- Downstream oligonucleotide primer
- Upstream oligonucleotide primer
- Nuclease-free water

**Laboratory workflow must be unidirectional, from pre-amplification to amplification areas. Specific equipment for each working area must be used, in order to avoid cross contaminations.**

For optimal results, it is essential to KEEP THE REACTION VIALS REFRIGERATED until their introduction in the thermal cyclers. When working with standard conical amplification vials, be sure to keep them on ice or in coolers, avoiding wetting the optical cap.

**Proceed to the Reagent Preparation Area in a laminar flow cabinet. Wear disposable gloves and use sterile and nuclease free plastic material in order to avoid contaminations and false negative results. We also recommend the use of filter tips.**

- 1.-Thaw and thoroughly mix all the reagents before dispensing (avoid vortex).
- 2.-**Dilute the fluorophore:** Before using, dilute the stock of your intercalating fluorophore to a working solution (follow the instructions and advice provided by the manufacturer). Protect the dye from prolonged exposure to light.
- 3.-Determine the number of samples to be analysed. If quantification is performed include samples of known concentrations that will be used for the standard curve. The inclusion of positive and negative controls is highly recommended in each experiment.
- 4.-Prepare the reactions according to Table 1. PROTECT MASTER MIX FROM PROLONGED EXPOSURE TO LIGHT.

**TABLE 1. Master Mix preparation**

COMPONENT	Final Concentration	20 µl rxn
<b>2X QUANTISPLIT</b>	1 X	10 µl
50 mM MgCl <sub>2</sub> Solution*	4-6 mM	x µl
Primers	0.1-0.5 µM	x µl
Diluted Intercalating fluorophore <sup>†</sup>	-	x µl
Sterile bidistilled water	-	Up to 20 µl
DNA template	variable	x µl

\*Only necessary for concentrations of MgCl<sub>2</sub> >4mM

<sup>†</sup> Final concentration: 0.1-0.5 X for SYBR<sup>®</sup> Green I or 0.5-1.5 X for Chromofly<sup>™</sup> Dye

**Proceed to DNA Purification Area separate from other sources of DNA.**

Never introduce DNA in the laminar flow cabinet from the reagent preparation area. Amplification must start in the next 10 minutes after adding purified DNA and primers to the amplification mix. Keep all reagents on ice until their introduction in the thermal cyclers.

- 5.-Add template DNA to each reaction tube. Close the tubes and mix gently.
- 6.-Centrifuge the amplification vials briefly.

**Proceed to Amplification Area**

- 7.-Place the tubes in the thermal cyclers and begin cycling.
- 8.-Program the thermal cyclers following the recommendations provided by the manufacturer (see TABLE 2).

**TABLE 2. PCR cycling parameters for QUANTIMIX HOTSPLIT EASY Kit**

Cycle Step	Nº Cycles	Temperature	Time
Initial Denaturation and Enzyme Activation	1	96°C	5-10 min
Denaturation Annealing Extension*	30-50	95-98°C	5-20 sec
		2-5°C<T <sub>m</sub> of the primers	5-10 sec
		70-72°C	30-40 sec
Fluorescence Acquisition (See Note 1)	1	T <sub>m</sub> of nonspecific products <X<T <sub>m</sub> specific products	10-15 sec <sup>+</sup>
Melting*	1	60-95°C	Start with a 0.5 °C/sec ramp

\* Fluorescence Acquisition during Extension and Melting Steps.

<sup>+</sup> The shortest time needed for reading fluorescence (different for each thermal cyclers).

**Note 1: When primer-dimer or nonspecific products appear, include an additional Fluorescence Acquisition Step after the Extension Step**

*This protocol has been optimised for the following real time quantification equipments: iCycler (Biorad), SmartCycler I and II (Cepheid), Rotor-Gene 3000 and 6000 (Corbett Research) and ABI PRISM 7500 series (Applied Biosystems). For other thermal cyclers, optimisation of the reaction parameters may be required. Please contact our Technical Department (info@biotoools.eu).*

The interpretation of the results is performed with the help of specific software. Therefore, follow the instructions and advice provided by the manufacturer.

## 6. TROUBLESHOOTING

**Little or no amplification detected**

1. **Check template quality and quantity.** Check the quality of template DNA by agarose gel electrophoresis or fluorimetry. Organic extraction followed by ethanol precipitation may remove some amplification inhibitors. Use of excess template can reduce PCR product yield.
2. **Redesign primers.** Design primers that have higher annealing temperatures and do not form hairpin loops or primer-dimers.
3. **Optimise primer concentration.** Although lower primer concentration can prevent primer-dimer formation, sufficient primers are needed for successful real-time PCR. Increase primer concentration in increments of 0.1 µM.
4. **Increase initial template denaturation time.** Increase the length of initial template denaturation up to 10 minutes in order to ensure that HotSplit DNA Polymerase has been completely activated.
5. **Lower annealing temperature.** Lower the annealing temperature in 2°C decrements.
6. **Increase number of cycles.** Perform additional cycles in increments of 5 cycles.
7. **Increase extension time.** Increase the extension time by increments of 30 sec. Generally 20 seconds/100 bp of PCR product should be enough.
8. **Activate detection step.** Check to make sure that the fluorescence detection step was activated when programming the cycles.
9. **Check detection step accuracy.** Ensure the fluorescence detection step takes place during the correct step of the PCR cycling program.
10. **Choose a Dye-compatible filter.** For a real-time instrument that is equipped with a multiple dye detection system, ensure that a filter compatible with your DNA-binding dye is activated.

**Multiple products or a smear detected**

1. **Decrease concentration of reaction components.** Check the concentration of template DNA by agarose gel electrophoresis or fluorimetry. Decrease the amount of DNA and/or primer added to the reaction.
2. **Check the primers for degradation.** Check by electrophoresis in a denaturing acrylamide gel.
3. **Redesign primers.** Design primers that have higher annealing temperatures and do not form hairpin loops or primer dimers.
4. **Increase annealing temperature.** Increase the annealing temperature in 2°C increments.
5. **Decrease number of cycles.** Decrease number of cycles in decrements of 5 cycles.
6. **Decrease PCR product size.** For optimal real time PCR, design primers that amplify PCR products between 100-300 bp in length.
7. **Include an additional Fluorescence Acquisition Step.** See Note 1.

**No linearity in amplification curve**

1. **Check template quantity.** An excess of template can affect the linearity.
2. **Include an additional Fluorescence Acquisition Step.** Non specific amplification products can affect the linearity (See Note 1).

**Fluorescence in no template control (NTC)**

1. **Repeat the assay with new reagents and take appropriate safety precautions.** If the T<sub>m</sub> of the NTC is similar to the T<sub>m</sub> of the target (contamination).
2. **Include an additional Fluorescence Acquisition Step.** If T<sub>m</sub> of NTC is lower than the T<sub>m</sub> of the target (primer-dimers) (See Note 1).

## 7. ORDERING INFORMATION

DESCRIPTION	Size	Reference
<b>QuantiSplit</b>	1100 µl	10.691
	2 x 1100 µl	10.692
	5 x 1100 µl	10.693
<b>50mM MgCl<sub>2</sub> Solution</b>	1.8 ml	10.691
	1.8 ml	10.692
	1.8 ml	10.693