

# Q-Extract DNA Extraction Hot Start PCR Kit

With TEMPase Hot Start 2x Master Mix A BLUE 1.5 mM MgCl<sub>2</sub> final concentration

Cat. No.: A574499 20 Reactions

MADE IN **DENMARK** 

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-	Q-Extract DNA Extraction Solution	TEMPase Hot Start 2x Master Mix A BLUE, 1.5 mM MgCl₂
ID No.	4900100	5200600
Cap colour	Clear	Red
Content	1 x 2 ml	1 x 0.25 ml

## **Product description**

Q-Extract DNA Extraction Hot Start PCR Kit consists of Q-Extract DNA extraction solution and TEMPase 2x Master Mix A BLUE, which is required for the subsequent PCR.

The Q-Extract DNA Extraction solution is designed for rapid and efficient extraction of PCR-ready DNA from various sample types; mammalian tissues (such as mouse tail and ear snips), plant leaves, saliva and bacteria. The non-toxic Q-Extract DNA Extraction Solution enables the extraction of DNA from tissues in just 8 minutes. The extraction protocol is divided into two simple heating steps, which is directly followed by PCR using TEMPase Hot Start 2x Master Mix A BLUE. This method is ideal for PCR analysis such as screening and genotyping.

The one-reagent DNA extraction set-up is easily scaled and can be conducted by robotic automation platforms. Depending on the sample size, the DNA extraction can be performed in PCR tubes or 1.5 ml tubes, using either a thermocycler or heating block.

TEMPase 2x Master Mix A BLUE is a ready-to-use 2x reaction mix. Each PCR reaction requires 12.5  $\mu$ l of the master mix. Simply add primers, DNA extract and water to a total reaction volume of 25  $\mu$ l to successfully carry out PCR.

TEMPase Hot Start DNA Polymerase is a modified form of Ampliqon Taq DNA polymerase, which is activated by heat treatment. A chemical moiety is attached to the enzyme at the active site, which renders the enzyme inactive at room temperature. Thus, during setup and the first ramp of thermal cycling, the enzyme is not active and misprimed primers are not extended. The result is higher specificity and greater yields when compared to standard DNA polymerases.

There is no need to use separate loading dyes. Simply load a portion of the reaction product onto an agarose gel for electrophoresis and subsequent visualization. The blue dye front runs at 400 - 500 bp on a 0.5 - 1.5% agarose gel.

Q-Extract DNA Extraction Hot Start PCR Kit allows for DNA extraction and amplification hereof in less than 2 hours, as compared to  $\geq 1$  day with conventional protocols.

## **Composition of Q-Extract DNA Extraction Solution**

Optimized DNA extraction solution

Composition of TEMPase Hot Start DNA Polymerase 2x Master Mix A BLUE, 1.5 mM MgCl<sub>2</sub>

- Tris-HCl pH 8.5, (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 3.0 mM MgCl<sub>2</sub>, 0.2% Tween<sup>®</sup> 20
- 0.4 mM of each dNTP
- TEMPase Hot Start DNA Polymerase
- Inert blue dye and stabilizer

# **Recommended Storage and Stability of Kit Components**

Q-Extract DNA Extraction Solution: Long term storage at -20 °C. Product expiry at -20 °C is stated on the label. Can be stored short term at +4 °C for up to 3 months. Q-Extract DNA Extraction Solution tolerates up to 20 freeze-thaw cycles. It is recommended to aliquot the Q-Extract into smaller volumes.

TEMPase 2x Master Mix A BLUE: Long term storage at -20 °C. Product expiry at -20 °C is stated on the label. Can be stored at +4 °C for up to 6 months.

## **Quality Control**

Each batch of Q-Extract DNA Extraction Solution is functionally tested.

TEMPase Hot Start DNA Polymerase is functionally tested for contaminating activities, with no traces of endonuclease activity, nicking activity or exonuclease activity.

## **Extraction Protocol**

Preparation of DNA extraction should be performed in a separate area from that used for setting up the PCR.

- Thaw Q-Extract DNA Extraction Solution. For the first time use, aliquot the Q-Extract DNA Extraction solution into smaller volumes. (Q-Extract DNA Extraction Solution has a cloudy appearance).
- Add your sample to a tube containing 100 µl Q-Extract DNA Extraction Solution. Recommended sample sizes are shown in Table 1.
- 3. Vortex the tube containing the sample and the DNA extraction solution for 15 sec.
- 4. Transfer the tube to a heat block or a thermal cycler and incubate for
  - 1. 65 °C for 6 min
  - 2. 98 °C for 2 min
  - 3. 4 °C (or cool down on ice)

The DNA extract is now ready for PCR. See PCR protocol and table 2.

DNA extracts are stable at -20  $^{\circ}\text{C}$  for one week or long term at -80  $^{\circ}\text{C}.$ 

Table 1. Sample sizes

Sample	Q-Extract DNA Extraction Solution		
	100 μΙ	500 μl	
Tissue*	0.5 – 10 mg	10 – 50 mg	
Plant**	2 – 10 mg	10 – 50 mg	
E. coli	1 colony (Φ 0.5 - 2 mm)	1 colony (Φ 0.5 - 5 mm)	
Saliva	10 – 20 μΙ	50 - 100 μΙ	

<sup>\*</sup> Examples of tested tissues include mouse tail snip, mouse organs and chicken breast.

<sup>\*\*</sup>Examples of tested plant materials include leaves from stinging nettle and ivy.

### **PCR Protocol**

This protocol serves as a guideline to ensure optimal PCR results when using TEMPase 2x Master Mix A BLUE. Optimal reaction conditions such as incubation times, temperatures, and amount of template DNA may vary and must be determined individually.

Thaw TEMPase 2x Master Mix A BLUE and primers.
 It is important to thaw the solutions completely and mix thoroughly before use to avoid localized concentrations of salts.

#### Keep all components on ice.

2. Prepare a reaction mix. Table 2 shows the reaction set up for a final volume of 25  $\mu$ L. If desired, the reaction size may be scaled up or down.

Table 2. Reaction components (reaction mix and template DNA)

Comments (reaction mix and template E				
Component Vol./reaction*		Final concentration*		
2x Master Mix	12.5 μΙ	1x		
25 mM MgCl <sub>2</sub>	Optional	1.5 mM (1.5 – 4.5 mM)		
Primer A (10 μM)	0.5 μl (0.25 – 2.5 μl)	0.2 μΜ (0.1 – 1.0 μΜ)		
Primer B (10 μM)	0.5 μl (0.25 – 2.5 μl)	0.2 μΜ (0.1 – 1.0 μΜ)		
PCR-grade H <sub>2</sub> O	ΧμΙ	-		
DNA Extract**	2 - 5 μΙ	Variable		
TOTAL volume	25 μΙ	-		

<sup>\*</sup> Suggested starting conditions; theoretically used conditions in brackets

- 3. Mix gently.
- Add extracted DNA to the individual tubes containing the reaction mix.
- 5. Program the thermal cycler according to the manufacturer's instructions. See table 3 for an example.
  - For maximum yield and specificity, temperatures and cycling times should be optimized for each new template target or primer pair.
- 6. Place the tubes in the thermal cycler and start the reaction.
- 7. At the end of the run, simply load a portion of the reaction product (e.g.  $10~\mu l$ ) onto an agarose gel for analysis.

Table 3. Three-step PCR program

Cycles	Duration of cycle	Temperature
1	15 minutes	95 ℃
25 - 35	20 – 30 seconds	95 ℃
	20 – 40 seconds	50 – 65 °C
	30 seconds	72 °C
1	5 minutes	72 °C

## Notes:

- For genotyping of fish fins and other applications please visit our website.
- The final MgCl<sub>2</sub> concentration of TEMPase 2x Master Mix A BLUE is 1.5 mM. In some PCR applications, more than 1.5 mM MgCl<sub>2</sub> is required for best results. Use 25 mM to adjust the Mg<sup>2+</sup> concentration according to table 4.

Table 4. Additional volume (μl) of MgCl<sub>2</sub> per 25 μl reaction:

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Final MgCl <sub>2</sub> conc. in reaction (mM)	1.5	2.0	2.5	3.0	3.5	4.0	4.5
Volume of 25 mM MgCl <sub>2</sub>	0	0.5	1	1.5	2	2.5	3

#### **Related Products**

**Q-Extract DNA Extraction PCR Kit \*** 

100 DNA extractions + PCR reactions	A570001
500 DNA extractions + PCR reactions	A570004
* Q-Extract DNA Extraction Solution + Taq DNA Pol. 2x Master Mix RED, 1.5 mM	
Q-Extract DNA Extraction Hot Start PCR Kit *	Cat. No.
100 DNA extractions + PCR reactions	A574401

Cat. No.

\* Q-Extract DNA Extraction Solution + TEMPase Hot Start DNA Pol. 2x Master Mix A BLUE, 1.5 mM MgCl<sub>2</sub>

Q-Extract DNA Extraction Solution	Cat. No.
100 DNA extractions	A560001
500 DNA extractions	A560004

Taq Master Mixes (500 x 50 μl reactions) *	Cat. No.
2x Master Mix, 1.5 mM MgCl <sub>2</sub> final concentration	A140303
2x Taq OptiMix CLEAR, 1.5 mM MgCl <sub>2</sub> final concentration	A370503
2x Master Mix RED, 1.5 mM MgCl <sub>2</sub> final concentration	A180303
TEMPase Hot Start Master Mixes (500 x 50 μl reactions)*	Cat. No.
2x Master Mix A**, 1.5 mM MgCl <sub>2</sub> final concentration	A230303
2x Master Mix A**BLUE, 1.5 mM MgCl <sub>2</sub> final concentration	A290403

\*Master mixes available also in 1.1x variants as well as 2 mM MgCl $_2$  variants, \*\*Mix A is Ammonium Buffer based, also available as Mix C based on Combination Buffer.

Special TEMPase Master Mixes (500 x 50 μl reactions)	Cat. No.
Multiplex 2x Master Mix, 3 mM MgCl <sub>2</sub> final concentratio	n <b>A260303</b>
GC TEMPase 2x Master Mix I – for GC-rich templates	A331703
GC TEMPase 2x Master Mix II – for GC-rich templates	A332703

Taq DNA Polymerase (500 units) *	Cat. No.
Taq DNA Polymerase 5 U/μl	A110003
with 10x Ammonium Buffer	A111103

\*Available in kits including one or two buffers (Ammonium Buffer, Standard Buffer or Combination Buffer). All kits include extra 25 mM MgCl<sub>2</sub>

Hot Start DNA Polymerase (500 units) *	Cat. No.
TEMPase Hot Start DNA Polymerase, 5 U/μl	A220003
with 10x Ammonium Buffer	A221103

\*Available in kits including one or two buffers (Ammonium Buffer, Standard Buffer or Combination Buffer). All kits include extra 25 mM MgCl<sub>2</sub>

Buffers for DNA polymerases *	Cat. No.
10x Ammonium Buffer, 3 x 1.5 ml	A301103
10x Standard Buffer, 3 x 1.5 ml	A302103
10x Combination Buffer, 3 x 1.5 ml	A303103
PCR Grade Water, 6 x 5 ml	A360056

\*Ammonium Buffer, Standard Buffer and Combination Buffer are also available as Mg<sup>2+</sup> free buffers, detergent free buffers and Mg<sup>2+</sup> and detergent free buffers.
\*\*For direct gel loading and visualisation.

For Research Use Only. Not for use in diagnostics procedures.

Other product sizes, combinations and customized solutions are available. Please look at www.ampliqon.com or ask for our complete product list for PCR Enzymes. For customized solutions please contact us.

Made in Denmark

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<sup>\*\*</sup> If the PCR yields are poor or one experience no bands, it might help to dilute the DNA extract 1:10. DNA extracts from plant leaves should be diluted 1:10 or 1:100, especially when analysing chloroplast DNA.