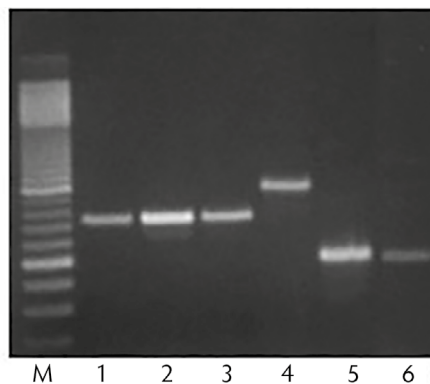


QuickExtract DNA Extraction Solution

Simple, rapid extraction of PCR-ready DNA

- The QuickExtract™ DNA Extraction Solution extracts PCR-ready genomic DNA from almost any sample in just 3-8 minutes.
- Many publications support the use of QuickExtract DNA Extraction Solution with samples such as hair follicles, quill-end cells of feathers, tissue-culture cells, buccal cells, zebrafish organs and scales, mouse tail snips, and more. The simple, single-tube procedure can accommodate one to hundreds of samples, and it is easily adapted to multiwell plates with robotic automation systems.
- The extracted DNA is suitable for PCR-based analysis, such as: genomic, transgenic, or viral DNA screening in animals; genetic or environmental research and screening in humans and other organisms; and CRISPR/Cas9 library screening. QuickExtract has also been used to isolate viral RNA for subsequent SARS-CoV-2 detection by RT-PCR or RT-LAMP.
- The convenient, scalable protocol involves gentle lysis and extraction that provides high yields of intact nucleic acid – all without the use of toxic chemicals or spin columns.
- **Rapid procedure:** eight-minute protocol for most sample types
- **Simple method:** single-tube protocol with no spin columns
- **Automation-friendly:** process one or hundreds of samples
- **Safe workflow:** no phenol, chloroform, or guanidinium salts
- **Many applications:** suitable for genotyping, human identity testing, viral/microbial screening, and more

Figure 1. FailSafe™ PCR amplifications of genomic DNA isolated using the QuickExtract procedure. All samples were treated with QuickExtract DNA Extraction Solution. PCR was performed using primers to amplify the regions indicated: Lanes 1-3, human β -globin (from human buccal cells, HeLa cells, and human hair follicle, respectively); lane 4, transgenic mouse GAPDH (from mouse tail snip); lane 5, 16S ribosomal RNA gene (from *E. coli*); lane 6, transgenic SV40 T antigen (from mouse tail snip).

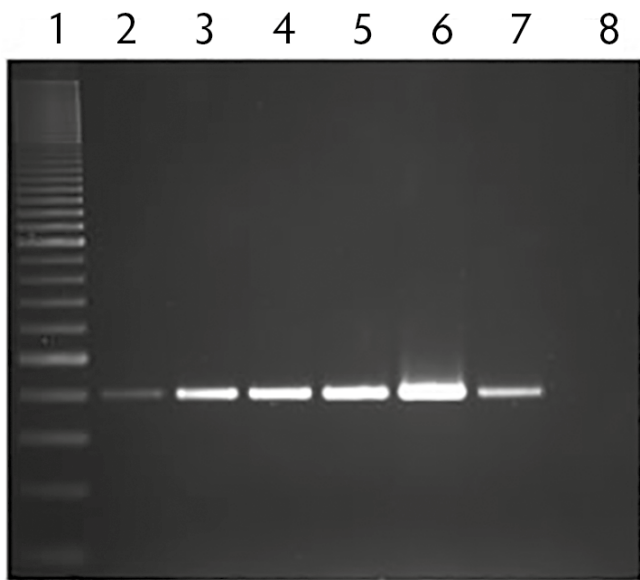


PCR-ready DNA from a variety of samples

Ordering information

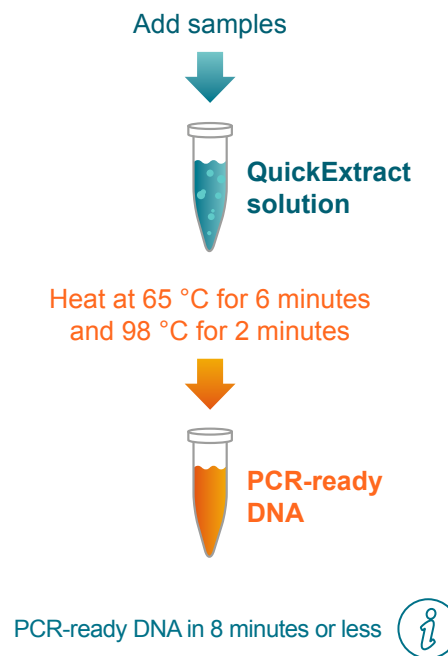
Cat no.	Size	Description
QE0905T	5 mL (10 extractions)	QuickExtract DNA Extraction Solution
QE09050	50 mL (100 extractions)	
QE0901L	1000 mL (2000 extractions)	

Figure 2. PCR amplification of DNA extracted from multiple zebrafish (*Danio rerio*) organs using QuickExtract DNA Extraction Solution. DNA was extracted from the following organs using 100 μ L of QuickExtract DNA Extraction Solution, and 1 μ L of each extracted sample was used to amplify a single-copy crystallin-like gene. Lane 1, 100-bp ladder; lanes 2-3, fins; lanes 4-5, eyes; lanes 6-7, scales; lane 8, no-DNA control.



Sensitive PCR detection from extracted DNA

Figure 3. The QuickExtract DNA Extraction Solution workflow.



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