illustra™ puReTaq Ready-To-Go PCR Beads

Product Booklet

Codes:
- 27-9557-01 (0.2 ml tubes/plate of 96)
- 27-9557-02 (0.2 ml tubes/5 plates of 96)
- 27-9558-01 (0.5 ml tubes, 100 reactions)
- 27-9559-01 (0.2 ml tubes, 96 reactions)
Page finder

1. Legal 3
2. Handling 4
   2.1. Safety warning and precautions 4
   2.2. Storage 4
   2.3. Quality control 4
3. Components of the kit 5
4. Materials not supplied 6
5. Introduction 7
6. Protocols 8
   6.1. Avoiding nucleic acid contamination 8
   6.2. Preliminary preparations and general handling instructions 8
   6.3. PCR with Ready-To-Go PCR Beads 9
   6.4. Thermal Cycling 10
7. Appendix 11
   7.1. Primers and templates – general considerations 11
   7.2. Primer annealing temperatures 11
   7.3. Magnesium chloride concentration 12
   7.4. Thermal cycling parameters 12
8. Troubleshooting 13
9. References 18
10. Companion products 19
1. Legal

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This product is sold under licensing arrangements with Roche Molecular Systems, F Hoffmann-La Roche Ltd and the Perkin-Elmer Corporation. Purchase of this product is accompanied by a limited license to use it in the Polymerase Chain Reaction (PCR) process for research in conjunction with a thermal cycler whose use in the automated performance of the PCR process is covered by the upfront license fee, either by payment to Perkin-Elmer or as purchased, i.e. an authorized thermal cycler.

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http://www.gehealthcare.com/lifesciences

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Buckinghamshire, HP7 9NA UK
2. Handling

2.1. Safety warnings and precautions

Warning: For research use only. Not recommended or intended for diagnosis of disease in humans or animals. Do not use internally or externally in humans or animals.

All chemicals should be considered as potentially hazardous. We therefore recommend that this product is handled only by those persons who have been trained in laboratory techniques and that it is used in accordance with the principles of good laboratory practice. Wear suitable protective clothing such as laboratory overalls, safety glasses and gloves. Care should be taken to avoid contact with skin or eyes. In the case of contact with skin or eyes wash immediately with water. See material safety data sheet(s) and/or safety statement(s) for specific advice.

2.2. Storage

Store at ambient room temperature in the airtight foil pouch with the desiccant. Once opened, completely reseal the pouch, fold the sealed edge over several times and seal with a clip. Store unopened and resealed pouches in a dessicator to maximize product lifetime.

2.3. Quality control

Each batch of puReTaq Ready-To-Go PCR beads are thoroughly tested for quality performance. Specifications for release are based upon performance, purity, and reproducibility of each individually dispensed reaction.
3. Components of the kit

The newly developed puReTaq™ Ready-To-Go™ Polymerase Chain Reaction (PCR) Beads are premixed and predispensed complete reactions for performing PCR amplifications. The beads contain recombinant puReTaq DNA polymerase, an industry leader for ambient stability and purity. These PCR beads have been optimized for endpoint- and intercalating dye-based real-time PCR amplifications. With the exception of primer and template, the convenient, ambient temperature-stable beads provide all the necessary reagents to perform 25-μl polymerase chain reactions. The beads are available predispensed into either 0.2-ml thin-walled micro-well plates (sufficient for 96 reactions), 0.2-ml thin-walled tubes or 0.5-ml PCR tubes.

**The following components are included in this product:**
puReTaq Ready-To-Go PCR Beads: Room temperature-stable beads containing stabilizers, BSA, dATP, dCTP, dGTP, dTTP, ~2.5 units of puReTaq DNA polymerase and reaction buffer. When a bead is reconstituted to a 25 μl final volume, the concentration of each dNTP is 200 μM in 10 mM Tris-HCl (pH 9.0 at room temperature), 50 mM KCl and 1.5 mM MgCl₂.

**World Wide Web address:** www.gehealthcare.com/lifesciences
Visit the GE Healthcare home page for regularly updated product information.
4. Materials not supplied

Reagents
- Water - Use only deionized or distilled water that is sterile and free of contaminating nucleic acid.
- DNA template - Best results are obtained with high quality DNA, as discussed on page 11.
- Primers (template-specific) - Guidelines for primer design and use are described on page 11.
- Mineral oil - If required for the thermal cycler being used.

Equipment
- Supplies for liquid handling - Gloves, vials, and pipette tips should be sterile; pipettor and microcentrifuge. Perform all reactions in the plastic microcentrifuge tubes provided in the kit; these tubes are suitably sterile for thermal cycling.
- Ice bucket or cold block - For maintaining puReTaq PCR beads at 4°C during rehydration and prior to thermal cycling.
- Thermal cycler - For cycling according to the specified conditions.
5. Introduction

PCR, or Polymerase Chain Reaction, is a method for the *in vitro* amplification of a specific sequence of DNA. A typical PCR contains template DNA containing the target sequence to be amplified, two primers that are complementary to the target DNA sequence, nucleotides, and a thermal-stable DNA polymerase. The reaction mixture is repeatedly cycled through alternating periods of thermal denaturation, annealing, and extension, resulting in exponential amplification of the target DNA sequence (1,2,3). puReTaq Ready-To-Go PCR beads are designed and manufactured using a proprietary technology licensed to GE Healthcare. Optimized specifically for PCR, the Ready-To-Go reaction beads are formulated with high quality buffer, nucleotides (dNTPs), and recombinant puReTaq DNA polymerase - only template DNA and template-specific primers need to be added. The Ready-To-Go PCR bead format significantly reduces the number of pipetting steps, thereby decreasing handling errors and increasing reproducibility.
6. Protocols

6.1. Avoiding nucleic acid contamination
puReTaq PCR beads have passed rigorous quality tests to ensure the lowest possible levels of contaminating prokaryotic and eukaryotic nucleic acids. Adherence to some simple precautions will prevent the reintroduction of contamination. Use sterile pipette tips with filters for dispensing, select molecular biology grade water (or better) to formulate all buffers, and wherever possible, autoclave all buffers prior to use. In addition, work in a laminar air flow hood or PCR bubble that has been illuminated with UV light. It is important to analyze or pipette PCR products in an area separate from that where the reactions will be set up.

To minimize environmental contamination with amplified products, routinely treat all work surfaces with a 10% (v/v) bleach solution. Ideally, allow the bleach to contact the surface for at least 10 minutes prior to wiping away with sterile water.

6.2. Preliminary preparations and general handling instructions
Please note that the beads contain buffers, dNTPs, enzyme, stabilizers, and BSA, all which have been pretreated to minimize contamination. Discard any beads that were accidentally dislodged from their respective containers.

Prepare the PCR beads as follows:
1. Remove the desired quantity of tubes from the foil pouch. Remove individual tubes from a strip of eight by cutting the plastic link between tubes with scissors.
2. Examine these tubes to verify that a bead is visible at the bottom of each tube. The beads are carefully screened by weight and appearance. Please discard any beads that appear substantially...
smaller or misshapen - an indication of moisture contamination.
Please refer to the recommended storage conditions.

3. If necessary, gently tap the tube against a hard surface to force each bead to the bottom of the tube.

4. Place the tubes into a container that allows easy access during your experiment. The beads are now ready for use.

6.3. PCR with Ready-To-Go PCR Beads
For general information concerning primer design and cycling parameters, refer to Appendixes 1–4.

When performing PCR amplifications, exercise extreme care to prevent DNA contamination as described above. Each PCR bead is designed for use in a 25 μl reaction volume (one PCR bead/tube). When resuspended in a final volume of 25 μl, each reaction will contain 1.5 mM MgCl₂. Please refer to Appendix 3 if a higher concentration of MgCl₂ is desired.

1. For each reaction, add the following to a tube containing a PCR bead:

   Note: Do not mix the tube contents until all the components (below) have been added to the tube containing the bead.

   5’ (forward) primer (5–25 pmol) X μl
   3’ (reverse) primer (5–25 pmol) Y μl
   Template DNA* Z μl
   Sterile high-quality water to a final volume of 25 μl

   *Start with 50 pg for a simple template such as plasmid DNA, or 50 ng for a complex template such as genomic DNA. Avoid template amounts > 1 μg.

2. Snap the caps (provided) onto the tubes, pushing down firmly to ensure a tight fit. Mix the tube contents by gently flicking the tube
with a finger. Vortex gently and then centrifuge the tube for a few seconds to bring the components to the bottom of the tube. The reaction is fully dissolved and mixed when it appears clear.

3. Place the reaction mixtures on ice or in a cold block until ready for cycling. Minimize the time on ice prior to cycling to prevent formation of background reaction products.

6.4. Thermal Cycling

The optimal cycling profile for a given PCR system and thermal cycler will vary and must be determined empirically. Cycle number can range from 20 to 40 depending on the desired yield of product. Thermal cycling results and product yield can vary with cycle conditions and thermal cycler used. Read the instructions provided with your thermal cycler and optimize reaction conditions accordingly.
7. Appendix

7.1. Primers and templates - general considerations:
In general, PCR primers should be 15–30 bp in length with a GC content of ~ 50%. Complementarity between primer pairs and within each primer must be avoided to minimize the production of primer-dimers. There are a number of excellent Web sites for designing primers - some of our favorites are: www.oligo.net, www.abrf.org, and www.basic.nwu.edu/biotools/oligocalc.html.

Template DNA purified by a variety of methods is a suitable substrate for PCR, but high quality DNA produces the most reproducible results. A typical PCR contains < 1 μg of template DNA and primers at a concentration of 0.2–1.0 μM. The optimal quantity of template and primers must be determined empirically for each new combination of template and primer. The reaction conditions described in this protocol are general recommendations only.

7.2. Primer annealing temperatures
The optimal annealing temperature depends on the sequence of the primers and their homology to the template DNA. With the Ready-To-Go PCR bead format, annealing temperatures slightly higher than those used in standard reactions should be used if nonspecific products are present following amplification. The annealing temperatures of the chosen primers often vary from their estimated melting temperatures ($T_m$). Although amplification is more specific at higher annealing temperatures, there might be some loss of longer target sequences during amplification and an overall reduction in yield. In contrast, when lower annealing temperatures are used, nonspecific amplification often increases along with an increase in specific target amplification. On occasion, it is necessary
to reoptimize annealing temperatures to achieve maximum performance. We recommend altering the temperature in increments of 2–5°C, starting at 10°C below the original annealing temperature.

### 7.3. Magnesium chloride concentration
When each PCR bead is rehydrated in a reaction volume of 25 μl, the mixture will contain 1.5 mM MgCl₂. If a higher concentration of Mg²⁺ is desired, the following table can be used to determine the volume of a sterile 10 mM MgCl₂ solution that should be added to increase the Mg²⁺ concentration of the reaction. If MgCl₂ is added to the reaction, decrease the amount of water added to the reaction to maintain a final reaction volume of 25 μl.

<table>
<thead>
<tr>
<th>Final [MgCl₂]</th>
<th>Volume of 10 mM MgCl₂ to add</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0 mM</td>
<td>1.25 μl</td>
</tr>
<tr>
<td>2.5 mM</td>
<td>2.50 μl</td>
</tr>
<tr>
<td>3.0 mM</td>
<td>3.75 μl</td>
</tr>
<tr>
<td>3.5 mM</td>
<td>5.00 μl</td>
</tr>
<tr>
<td>4.0 mM</td>
<td>6.25 μl</td>
</tr>
<tr>
<td>4.5 mM</td>
<td>7.50 μl</td>
</tr>
<tr>
<td>5.0 mM</td>
<td>8.75 μl</td>
</tr>
</tbody>
</table>

### 7.4. Thermal cycling parameters
A standard three-step endpoint PCR cycling protocol consists of multiple cycles of denaturation (95°C), annealing (40–60°C) and extension (72°C). An initial denaturation step (95°C for 5 minutes) is recommended to ensure complete denaturation of the template DNA.

In some cases, it may be possible to amplify a target sequence using a two-step PCR where the denaturation step (95°C) is followed by a combined annealing/elongation step (50–65°C). For most standard, three-step PCRs, 35 cycles produces a 105–109-fold amplification of the target sequence. PCR product yield can be improved by increasing the number of cycles to 40.
8. Troubleshooting

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible causes/solutions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. No amplification</td>
<td>1. The thermal cycler did not function properly. Improper cycling conditions can result in poor amplification. Consider testing PCR beads in a control reaction with template DNA and primers previously shown to work successfully.</td>
</tr>
<tr>
<td></td>
<td>2. Primer was omitted from the reaction. PCR beads contain no primer; they must be added to the reaction by the researcher.</td>
</tr>
<tr>
<td></td>
<td>3. The reaction volume was incorrect. Each PCR bead should be resuspended to a final reaction volume of 25 μl. Further dilution of the beads will severely compromise performance. Volumes less than 25 μl will alter salt concentrations and alter primer annealing stringency.</td>
</tr>
<tr>
<td></td>
<td>4. Insufficient DNA was used in the PCR. The amount of DNA required to generate good signal can vary between</td>
</tr>
<tr>
<td>Problem</td>
<td>Possible causes/solutions</td>
</tr>
<tr>
<td>---------</td>
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</tr>
<tr>
<td>4. Continued different PCR systems. Titrate the amount of template in the reaction, starting with 50 pg for a simple template such as plasmid DNA, or 50 ng for a complex template such as genomic DNA.</td>
<td></td>
</tr>
<tr>
<td>5. The quality of the DNA template was poor. Impure DNA might fail to amplify. To inactivate contaminating enzyme activities, heat the template DNA to 95°C for 5 minutes before use in PCR. Alternatively, isolate template DNA by another method.</td>
<td></td>
</tr>
<tr>
<td>6. Examine beads for size and uniformity. If the pouch was previously opened and the beads subsequently stored without properly resealing the pouch, atmospheric moisture will rehydrate the beads. The result will be beads that are misshapen or markedly smaller. Reactions containing such beads should be discarded. To achieve optimum performance and maximum</td>
<td></td>
</tr>
<tr>
<td>Problem</td>
<td>Possible causes/solutions</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>2. Excessive background amplification</td>
<td>6. Continued storage lifetime, store the beads in a dessicator after resealing the pouch by folding over the opened edge several times and clipping shut with a paper clip or the equivalent.</td>
</tr>
<tr>
<td></td>
<td>1. Too much template DNA was added to the reaction. Reduce the amount of template DNA in the reaction until the smearing is eliminated.</td>
</tr>
<tr>
<td></td>
<td>2. The reaction utilized more than 35 cycles. Although the yield of PCR product can be increased by increasing the number of cycles to 40, this can produce spurious bands and increased background. Reduce the number of cycles until the smearing is eliminated.</td>
</tr>
<tr>
<td></td>
<td>3. Cycling conditions vary depending on the thermal cycler used. Optimize cycling conditions based on the manufacturer’s recommendations.</td>
</tr>
<tr>
<td>Problem</td>
<td>Possible causes/solutions</td>
</tr>
<tr>
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</tr>
<tr>
<td>4. The annealing temperature was too low. The optimal annealing temperature depends on the sequence of the primers and their homology to the template DNA. With the Ready-To-Go PCR bead format, annealing temperatures might vary slightly from those used in standard master mix PCR. Reoptimization of annealing temperature might be required. Increase the annealing temperature by 2–5°C increments.</td>
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</tr>
<tr>
<td>5. The quality of the DNA template was poor. Impure DNA can fail to amplify properly. Use freshly prepared DNA or isolate the template by another method.</td>
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</tr>
<tr>
<td>6. Too much primer was added to the reaction, resulting in the formation of primer-dimer bands. Excessive primer:template ratios can cause an abundance of low molecular weight bands and smearing. Titrate the amount</td>
<td></td>
</tr>
<tr>
<td>Problem</td>
<td>Possible causes/solutions</td>
</tr>
<tr>
<td>---------</td>
<td>---------------------------</td>
</tr>
<tr>
<td>6. Continued of primers in the reaction until the primer-dimer band is eliminated.</td>
<td></td>
</tr>
<tr>
<td>7. Primers were not properly designed. Complementarity between primer pairs and within each primer should be avoided.</td>
<td></td>
</tr>
<tr>
<td>3. Nonspecific amplification</td>
<td></td>
</tr>
<tr>
<td>1. The primers hybridized to a secondary site on the template. Design new primers that are less specific for the secondary site. Increase the annealing temperature by 2–5°C increments until the desired specificity is achieved.</td>
<td></td>
</tr>
<tr>
<td>2. Contamination in primers, template, or buffers. Prepare fresh materials.</td>
<td></td>
</tr>
</tbody>
</table>
9. References


10. Companion products

<table>
<thead>
<tr>
<th>Product</th>
<th>Pack size</th>
<th>Product number</th>
</tr>
</thead>
<tbody>
<tr>
<td>GenomicPrep™ Blood DNA Isolation Kit</td>
<td>100 purifications</td>
<td>27-5236-01</td>
</tr>
<tr>
<td>GenomicPrep Cells and Tissue DNA Isolation Kit</td>
<td>55 purifications</td>
<td>27-5237-01</td>
</tr>
<tr>
<td>GFX™ MicroPlasmid Prep Kit</td>
<td>250 purifications</td>
<td>27-9601-02</td>
</tr>
<tr>
<td>GFX PCR DNA and Gel Band Purification Kit</td>
<td>100 purifications</td>
<td>27-9602-01</td>
</tr>
<tr>
<td>GFX 96 PCR Purification Kit</td>
<td>2 x 96 plates</td>
<td>25-6902-01</td>
</tr>
<tr>
<td>100 Base-Pair Ladder</td>
<td>100 μg</td>
<td>27-4007-01</td>
</tr>
</tbody>
</table>