Hot Start PCR

**JumpStart™ Taq DNA Polymerase**

*Increase Specificity and Yields*

Sigma's JumpStart Taq DNA Polymerase is an antibody inactivated hot start enzyme designed to minimize non-specific amplification while increasing target yield. Once the reaction temperature reaches 70 °C, Taq DNA Polymerase activity is restored and the resulting PCR exhibits a higher specificity and yield. This antibody-enzyme complex allows for easy and convenient set-up with less contamination risk than manual hot start techniques. Since the enzyme can be included in the master mix preparation, more consistent results are obtained from one reaction to the next.

**Features and Benefits**

- Minimize non-specific amplification while increasing target yield and specificity
- Superior amplification regardless of template concentration
- Reduce set-up time and eliminate concerns associated with manual or wax hot start methods

JumpStart Taq DNA Polymerase is provided with a 10× Reaction Buffer available with and without MgCl₂. The 10× Buffer without MgCl₂ includes a separate tube of 25 mM MgCl₂ for optimization.

**Components:** JumpStart Taq DNA Polymerase
10× PCR Buffer or 10× PCR Buffer without MgCl₂ and a separate tube of 25 mM MgCl₂

**Unit definition:** One unit incorporates 10 nmol of total dNTPs into acid-precipitable DNA in 30 min at 74 °C

**Concentration:** 2.5 units per µl

**Storage:** –20 °C
Shipped in wet ice

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Inactivated by a Taq-Directed Antibody, JumpStart Taq Provides Increased Sensitivity, Specificity and Yield

Inactivated by a Taq-directed antibody, JumpStart Taq provides increased sensitivity, specificity and yield. Ten nanograms (even lanes) or 100 nanograms (odd lanes) total human genomic DNA was amplified with primers targeted to a 5 kb section of β-globin. Each PCR was performed according to supplier’s recommendations.

**Ordering Information**

<table>
<thead>
<tr>
<th>Cat. No.</th>
<th>Product Description</th>
<th>Quantity</th>
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<tbody>
<tr>
<td>D9307</td>
<td>JumpStart Taq DNA Polymerase</td>
<td>50 units</td>
</tr>
<tr>
<td></td>
<td></td>
<td>250 units</td>
</tr>
<tr>
<td></td>
<td></td>
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<tr>
<td>D4184</td>
<td>JumpStart Taq DNA Polymerase without MgCl₂</td>
<td>50 units</td>
</tr>
<tr>
<td></td>
<td></td>
<td>250 units</td>
</tr>
<tr>
<td></td>
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<td>1,500 units</td>
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JumpStart™ REDTaq®
DNA Polymerase

JumpStart REDTaq DNA Polymerase is a specialized blend of REDTaq Genomic DNA Polymerase and JumpStart Taq Polymerase. JumpStart Taq Polymerase is an antibody inactivated hot start enzyme designed to minimize non-specific amplification while increasing target yield. Unlike other hot start methods (i.e., chemical inactivation), JumpStart Taq Polymerase does not require a pre-incubation step prior to cycling because polymerase activity is fully restored during the first denaturation cycle of the PCR reaction.

JumpStart REDTaq DNA Polymerase has the benefits of REDTaq Genomic DNA Polymerase including enhanced amplification of genomic templates, easy visualization of enzyme addition, complete reaction mixing and direct loading of samples following amplification.

The inert red dye does not affect automated sequencing, restriction enzyme digestion, ligation or other downstream applications. The PCR product can be easily separated from the dye by standard purification methods.

Features and Benefits

- Minimize non-specific amplification while increasing target yield and specificity
- Superior amplification regardless of template concentration
- Reduce set-up time and eliminate concerns associated with manual or wax hot start methods. JumpStart Taq minimizes non-specific amplification while increasing target yield. The red dye also provides quick recognition and confirmation of appropriate mixing. Aliquots can be loaded directly on an agarose gel without the need for loading buffers or tracking dyes.

Components:
JumpStart Taq DNA Polymerase
10× PCR Buffer

Unit definition: One unit incorporates 10 nmol of total dNTPs into acid-precipitable DNA in 30 min at 74 °C

Concentration: 1 unit per µl

Storage: −20 °C
Shipped in wet ice

Ordering Information

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<th>Product Description</th>
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<tr>
<td>D8187</td>
<td>JumpStart REDTaq DNA Polymerase</td>
<td>50 units, 250 units, 2,500 (10 × 250) units</td>
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Hot Start PCR

JumpStart™ Taq ReadyMix™ PCR Reaction Mixes

JumpStart Taq ReadyMix PCR Reaction Mix is a prepared solution combining the performance benefits of hot start PCR with the convenience of a ReadyMix. The mix includes Sigma’s high quality JumpStart Taq DNA Polymerase, 99% pure deoxynucleotides and buffer in a 2× optimized reaction concentrate. Add 25 µl of ReadyMix to template, primers and water to a final reaction volume of 50 µl. JumpStart Taq Polymerase is an antibody inactivated hot start enzyme designed to minimize non-specific amplification while increasing target yield. Unlike other hot start methods (i.e., chemical inactivation), JumpStart Taq polymerase does not require a pre-incubation step prior to cycling because polymerase activity is fully restored during the first denaturation cycle of the PCR reaction.

Using JumpStart Taq ReadyMix reduces pipetting steps and risk of contamination. The hot start mechanism allows for room temperature set-up, making this the product of choice for high throughput applications.

Features and Benefits

- Minimize non-specific amplification while increasing target yield and specificity
- Superior amplification regardless of template concentration
- Reduce set-up time and eliminate concerns associated with manual or wax hot start methods. JumpStart Taq minimizes non-specific amplification while increasing target yield

Available in Direct Load

JumpStart REDTaq ReadyMix PCR Reaction Mix combines the advantages of JumpStart Taq ReadyMix with the added convenience of an inert red dye. This dye provides quick visual confirmation that the enzyme has been added and properly mixed. After PCR, an aliquot can be loaded directly onto an agarose gel without the need for loading buffers or tracking dyes.

Available in a High Throughput, Genomic Formulation

JumpStart REDTaq ReadyMix PCR Reaction Mix for High Throughput PCR is formulated with REDTaq Genomic DNA Polymerase for amplification of more complex or genomic templates. This mix contains optimized enzyme and dye concentrations to provide increased length and yield on more difficult templates.

Exceptional Performance with JumpStart REDTaq ReadyMix Hot Start PCR in the Convenience of a ReadyMix

Exceptional performance with JumpStart REDTaq ReadyMix Hot Start PCR in the convenience of a ReadyMix. 200 ng Lambda phage DNA was amplified with Sigma’s JumpStart REDTaq ReadyMix (odd numbered lanes) and Competitor I’s Direct Load ReadyMix (even numbered lanes). Taq was activated per the supplier’s recommendations.

Unit definition: One unit incorporates 10 nmol of total dNTPs into acid-precipitable DNA in 30 min at 74 °C

Storage: –20 °C
Shipped in wet ice

Ordering Information

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<tr>
<td>P2893</td>
<td>JumpStart Taq ReadyMix 1.5 units Taq/reaction (50 µl reaction volume)</td>
<td>100 reactions</td>
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<td>P0982</td>
<td>JumpStart REDTaq ReadyMix PCR Reaction Mix 1.5 units Taq/reaction (50 µl reaction volume)</td>
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<td></td>
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<td>100 reactions</td>
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<td>P1107</td>
<td>JumpStart REDTaq ReadyMix PCR Reaction Mix for High Throughput PCR 0.75 units Taq/reaction (50 µl reaction volume)</td>
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<td>400 reactions</td>
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JumpStart™ AccuTaq™ LA DNA Polymerase

**Increase Yields, Fidelity and Specificity**

JumpStart AccuTaq LA DNA Polymerase is a combination of AccuTaq LA DNA Polymerase plus a Taq-directed antibody. JumpStart Taq Polymerase is an antibody inactivated hot start enzyme designed to minimize non-specific amplification while increasing target yield. Unlike other hot start methods (i.e., chemical inactivation), JumpStart Taq Polymerase does not require a pre-incubation step prior to cycling because polymerase activity is fully restored during the first denaturation cycle of the PCR reaction. JumpStart AccuTaq LA DNA Polymerase can generate long products with higher fidelity (up to 6.5× of Taq DNA Polymerase).

**Features and Benefits**

- Minimize non-specific amplification while increasing target yield and specificity
- Fidelity up to 6.5× that of Taq DNA Polymerase making it the enzyme of choice for multiplex PCR
- Produce amplicons up to 22 kb on genomic templates and up to 40 kb on less complex templates such as lambda or bacterial genomic DNA
- Reduce set-up time and eliminate concerns associated with manual or wax hot start methods. JumpStart Taq minimizes non-specific amplification while increasing target yield

Supplied with 10× Reaction Buffer.

**Available in Direct Load**

JumpStart REDAccuTaq LA DNA Polymerase combines all the advantages of JumpStart AccuTaq LA with the added convenience of an inert red dye. This dye provides quick visual confirmation that the enzyme has been added and properly mixed. After PCR, an aliquot can be loaded directly onto an agarose gel without the need for loading buffers or tracking dyes.

**Greater Specificity and Increased Yield with JumpStart AccuTaq and JumpStart REDAccuTaq**

Greater specificity and increased yield with JumpStart AccuTaq and JumpStart REDAccuTaq. Long and accurate hot start enzymes were used to amplify a 5 kb fragment starting with 25 ng of total human genomic DNA. All reactions were performed according to the manufacturer’s specifications.

Lane M: Wide Range DNA marker (D7058)
Lane 1: JumpStart AccuTaq LA
Lane 2: JumpStart REDAccuTaq LA
Lane 3: Supplier I, enzyme P
Lane 4: Supplier I, enzyme P
Lane 5: Supplier I, enzyme HF

**Components:** JumpStart AccuTaq LA DNA Polymerase or JumpStart REDAccuTaq LA DNA Polymerase

10× AccuTaq LA Buffer

**Unit definition:** One unit incorporates 10 nmol of total dNTPs into acid-precipitable DNA in 30 min. at 74 °C

**Storage:** −20 °C

Shipped in wet ice

**Ordering Information**

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<thead>
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<th>Cat. No.</th>
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<tr>
<td>D5809</td>
<td>JumpStart AccuTaq LA DNA Polymerase 2.5 units per µl</td>
<td>125 units 500 units</td>
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<td>D1313</td>
<td>JumpStart REDAccuTaq LA DNA Polymerase 1 unit per µl</td>
<td>50 units 250 units</td>
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Hot Start PCR

JumpStart™ Taq Antibody

The primary purpose of all hot start PCR methods is to prevent Taq DNA Polymerase activity prior to thermal cycling. Even if set-up is conducted on ice, Taq DNA Polymerase remains active and may elongate unwanted products such as misprimed or other nonspecific events. Primer-dimer interactions may also occur, which will reduce overall yield and efficiency.

One method commonly used to prevent unwanted amplification products is to add JumpStart Taq Antibody to the reaction. This efficient, yet simple, procedure takes only 10 minutes and effectively inactivates the Taq DNA polymerase until the first denaturation cycle. Unlike other hot start methods (i.e., chemical inactivation), JumpStart Taq Polymerase does not require a pre-incubation step prior to cycling because polymerase activity is fully restored during the first denaturation cycle of the PCR reaction. Upon heating to 70 °C, the antibody dissociates and full activity is restored to the Taq DNA Polymerase for the remainder of the PCR. JumpStart Taq antibody works effectively on a variety of Taq based DNA Polymerases that are commercially available.

Two units of Taq DNA Polymerase are inactivated by 1 test of JumpStart Taq Antibody.

Features and Benefits

- Minimize amplification while increasing target yield
- Reduce set-up time and eliminate concerns associated with manual or wax hot start methods

Components: JumpStart Taq Antibody

Dilution Buffer for JumpStart Taq Antibody

Storage: –20 °C
Shipped in wet ice

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<thead>
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<th>Quantity</th>
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<td>JumpStart Taq Antibody</td>
<td>200 tests</td>
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<td>500 tests</td>
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