





| Cat.# Size |                   |  |
|------------|-------------------|--|
| BG10110S   | 40 R(50 μL vol)   |  |
| BG10110M   | 200 R(50 μL vol)  |  |
| BG10110L   | 1000 R(50 μL vol) |  |

Source: E.coli strain with a cloned pol gene from Thermus aquaticus YT1.

Storage temperature: -20°C Exp.: 12 months(-15°C ~ -25°C)

2. PCR product for TA cloning

3. Colony PCR.

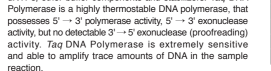


website: syngen.tech

Component 2 × Tag Colorless Master Mix contains Tag DNA Polymerase. dNTPs, other buffer components and stabilizers. Tag DNA 10 uM forward primer 10 uM reverse primer Template DNA Nuclease-Free water

| Cat.#    | Size              | Quantity  | Nuclease-Free Water | cond |
|----------|-------------------|-----------|---------------------|------|
| BG10110S | 40 R(50 μL vol)   | 1 mL      | 1 mL                |      |
| BG10110M | 200 R(50 μL vol)  | 5 × 1 mL  | 5 × 1 mL            | Init |
| BG10110L | 1000 R(50 μL vol) | 25 × 1 mL | 25 × 1 mL           |      |

1. Routine PCR amplification of DNA fragments up to 5 kb



## Reagent supplied:

|    | Size              | Quantity  | Nuclease-Free Water | conditions:          |
|----|-------------------|-----------|---------------------|----------------------|
| 0S | 40 R(50 μL vol)   | 1 mL      | 1 mL                | Step                 |
| OM | 200 R(50 μL vol)  | 5×1 mL    | 5×1 mL              | Initial Denaturation |
| 0L | 1000 R(50 μL vol) | 25 × 1 mL | 25 x 1 mL           | Denaturation         |

## Protocol

o prepare several parallel reactions with minimized Extension pipetting errors, prepare a PCR master mix by mixing Nuclease-Free water, primers and 2 x Tag Master Mix. Final Extension Prepare sufficient master mix for the number of reactions General guidelines: plus one extra. Aliquot the master mix into individual PCR tubes and then add template DNA.

- 1. Gently vortex and briefly centrifuge all solutions after
- 2. Place a thin-walled PCR tube on ice and add the DNA following component for each 25 µL or 50 µL reaction:

calculated Tm

5 min at 68°C is recommended.

25 μL 50 μL reaction co 1 uL variable 0.1-100 ng × Tag Colorless Master Mix | 12.5 μL | 25 μL Denaturation to 25 μL to 50 μL An initial denaturation of 2 min at 94°C is sufficient for

- 3. Gently vortex the samples and spin down.
- 4.Perform PCR using recommended thermal cycling

| Step            | Temperature | Time       | Number of cycles | colony          |
|-----------------|-------------|------------|------------------|-----------------|
| al Denaturation | 94°C        | 2-5 min    | 1                | recomn          |
| Denaturation    | 94°C        | 15-30 s    |                  | 4. Anne         |
| Annealing       | Tm-5°C      | 10-30 s    | 25-40            | The and is base |
| Extension       | 68°C        | 10-60 s/kb |                  | 68°C. A         |
| nal Extension   | 68°C        | 5-15 min   | 1                | a temp          |

The recommended extension temperature is 68°C. Extension time are generally 20-30 s/kb for complex, genomic samples, success of PCR reaction. Recommended amounts of DNA but can be reduced to 10 s/kb for simple templates (plasmid. template for a 50 uL reaction are as follows: virus). To determine specific extension time, we recommend

| DNA                  | Amount    |
|----------------------|-----------|
| Genomic DNA          | 1 ng-1 μg |
| Plasmid or Viral DNA | 1 pg-1 μg |

## Cycle number:

Oligonucleotide primers are generally 20-40 nucleotides in Generally. 25-35 cycles yields sufficient product. Up to 45 length and ideally have a GC content of 40-60%. Software such as Oligo 7 can be used to design or analyze primers. 7. PCR product: The final concentration of each primer in a PCR reaction may be 0.05-1 μM, typically 0.1-0.5 μM.

most amplicons from pure DNA templates. For difficult Chien A. Edgar D B. Trela J M. Deoxyribonucleic-acid templates such as GC-rich sequences or crude samples a longer denaturation of 2-8 min at 94°C is recommended Journal of Bacteriology, 1976, 127. prior to PCR cycling to fully denature the template. With 2. Kaledin A S. Sliusarenko A G. Gorodetski S I. Isolation and PCR, an initial 2-5 min denaturation at 94°C is

1 min/kb and reduce the extension time until 10 s/kb. 2 min/

kb is recommended for crude samples. A final extension of

nnealing step is typically 15-60 s. Annealing temperature Research only, not for the rapeutic or diagnostic purposes. sed on the Tm of the primer pair and is typically 45 Annealing temperatures can be optimized by doing a temperature gradient PCR starting at 5°C below the

644-651.

## 2× Tag Colorless Master Mix 使用说明书

【产品来源】重组表达柄热水生菌 DNA 聚合酶基因的大肠杆

2× Tag Colorless Master Mix 是一种加入了 Tag DNA 聚合

合生基因的 Tag DNA 聚合酶对微量的 DNA 样品具有很好的扩

增能力。 Tag DNA 聚合酶具有热稳定性,  $5' \rightarrow 3'$  端 DNA 聚合

酶活性和  $5' \rightarrow 3'$  端核酸外切酶活性,无  $3' \rightarrow 5'$  端核酸外切酶

BG10110L 1000 R(50 μL vol) 25 × 1 mL 25 × 1 mL

数量

无核酸酶水

1 mL

5 × 1 mL

【随酶提供试剂】

规格

BG10110M | 200 R(50 uL vol) | 5 × 1 mL

BG10110S | 40 R(50 µL vol)

货号

【产品规格】 cycles may be required to detect low-copy-number templates 货号

40 R(50 μL vol) The PCR products generated by Taq DNA Polymerase BG10110M 200 R(50 μL vol) contain dA overhangs at the 3' end, therefore can be BG10110L 1000 R(50 μL vol) ligated to dT/dU-overhang vectors.

- 【储存条件】-20℃ 【保质期】12 个月(-15℃~-25℃) polymerase from extreme thermophile Thermus-aquaticus[J]. 【应用范围】
- 1. 常规 PCR 扩增 5 kb 以内的 DNA 片段 2. 生成用于 TA 克降的 PCR 产物 properties of DNA polymerase from extreme thermophylic 3. 菌落 PCR bacteria Thermus aquaticus YT-1[J]. Biokhimiia. 1980. 45(4): 【产品说明】

酶, dNTPs, 其它一些缓冲液组分和 PCR 稳定剂的预混液。 V1.0.3 © 13-14 Beijing SyngenTech Co., Ltd. All rights reserved

## 【操作指南】

如果要同时讲行几个平行反应,混合无核酸酶水,反应缓

冲液, 引物和 2× Tag Colorless Master Mix 制备 PCR 混合液 为减少加样误差的影响,制备的反应混合液要在需要的反应数 量上添加上一个额外反应, 分装母液到单个的 PCR 管中, 然 后加入模板 DNA。

- 1. 温和震荡后, 快速离心解冻后的所有溶液
- 2. 加入 25 uL 或者 50 uL 的组份(见下表)到冰上的薄壁

| 风分                           | 25 µL 反应 | 50 µL 反应 | 终浓度        |
|------------------------------|----------|----------|------------|
| 10 µM 正向引物                   | 0.5 μL   | 1 μL     | 200 nM     |
| 10 μM 反向引物                   | 0.5 μL   | 1 μL     | 200 nM     |
| 模板 DNA                       | 可变量      | 可变量      | 0.1-100 ng |
| 2 × Taq Colorless Master Mix | 12.5 μL  | 25 μL    | 1×         |
| 无核酸酶水                        | 至 25 μL  | 至 50 μL  | -          |

- 3. 温和震荡并短暂离心。
- . 使用推荐的热循环条件讲行 PCR.

| J-3K | /III./X | L 1 [m]    | 122  |
|------|---------|------------|------|
| 预变性  | 94℃     | 2-5 min    | 1    |
| 变性   | 94℃     | 15-30 s    |      |
| 退火   | Tm-5℃   | 10-30 s    | 25-4 |
| 延伸   | 68℃     | 10-60 s/kb |      |
| 最后延伸 | 68℃     | 5-15 min   | 1    |

## 【使用说明】

模板。高质量、纯化的 DNA 模板会大幅提升 PCR 反应的

功率。对于一个 50 uL 反应体系, 推荐量的 DNA 模板如7

| DNA   | 用量        |  |
|-------|-----------|--|
| 基因组   | 1 ng-1 μg |  |
| 质粒或病毒 | 1 pg-1 ng |  |

- 2. 引物, 寡核苷酸引物—般长度在 20-40 nt. GC 含量为 40-60%。如 Oligo 7 软件可被用来设计或分析引物。在 PCR 反应 中每个引物的终浓度范围是 0.05-1 μM, 通常为 0.1-0.5 μM。
- 3. 变性: 94℃ 2 min 预变性即可满足干大部分纯化的 DNA 模 05 1 5 50 1 5 4 4 4 4 6 板的扩增。对于高 GC 含量序列或者粗样品、推荐使用 2-8 min 的变性时间来完全变性模板。对于菌落 PCR, 推荐使用 2-5 min 的预变性时间。
  - 4. 退火; 退火时间通常为 15-60 s, 退火温度基于引物的 Tm 值, 一般在 45℃到 68℃之间。退火温度可以通过低于 Tm-5℃的温 度梯度 PCR 进行优化。
  - 5. 延伸,推荐的延伸温度为68℃。复杂或基因组样品的延伸 时间一般为 20-30 s/kb。简单模板(质粒,病毒)可以为 10 s/ kb。为确定具体的延伸时间,我们推荐从 1 min/kb 开始降低弧 伸时间直到 10 s/kb。粗样品推荐使用 2 min/kb。68℃最终延 伸时间推荐使用 5 min。
  - 6. 循环数: 一般来说, 25 到 35 个循环即可以产生足够的 PCR 产物。最高到 45 个循环可用干检测低拷贝模板。
  - 7. PCR 产物、使用 Tag DNA 聚合酶产生的 PCR 产物在 3'端 会含有 dA 突出,因此,PCR 产物可以连接到 dT/dU 粘性末端 载体上。

# 【参考文献】

1. Chien A. Edgar D B. Trela J M. Deoxyribonucleic-acid polymerase from extreme thermophile Thermus-aquaticus[J]

Journal of Bacteriology, 1976, 127.

2. Kaledin A S. Sliusarenko A G. Gorodetski S I. Isolation and properties of DNA polymerase from extreme thermophylic bacteria Thermus aquaticus YT-1[J]. Biokhimiia, 1980, 45(4):

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