PCR

Monday, December 05, 2011
1:10 PM

Sean Weise

http://signal.salk.edu/tdnaprimers.2.html

PCR

Materials:
Sterile dH₂O
5 U/µl Taq (Invitrogen 18038-042)
10X PCR Buffer (comes with Invitrogen Taq)
50 mM MgCl₂ (comes with Invitrogen Taq)
2.5 mM each dNTP solution (Invitrogen 18038-042)
10 µM Forward and Reverse primers
Agarose
TAE solution (1X and 10X)
Gel casting mold, combs, and electrophoresis apparatus
EZ-vision Loading buffer / DNA dye (Amresco, N472-Q-0.5ml)
1 Kb ladder (Amresco, K181-500µl)

Method:
Do the following methods on ice
1. Prepare a “Master Mix” in a 0.5 ml microfuge tube for your PCR reactions according to the following table

<table>
<thead>
<tr>
<th>Reagent (conc in PCR tube)</th>
<th>1X (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sterile dH₂O</td>
<td>12.5</td>
</tr>
<tr>
<td>10X PCR Buffer (1X)</td>
<td>2</td>
</tr>
<tr>
<td>50 mM MgCl₂ (2.5 mM)</td>
<td>1</td>
</tr>
<tr>
<td>10 mM dNTPs (150 µM)</td>
<td>0.3</td>
</tr>
<tr>
<td>5 U/µl Taq (1 U)</td>
<td>0.2*</td>
</tr>
<tr>
<td><strong>Sub Total</strong></td>
<td><strong>16</strong></td>
</tr>
<tr>
<td>10 µM Forward Primer (1 µM)</td>
<td>1</td>
</tr>
<tr>
<td>10 µM Reverse Primer (1 µM)</td>
<td>1</td>
</tr>
<tr>
<td>DNA</td>
<td>2</td>
</tr>
<tr>
<td><strong>Total Volume</strong></td>
<td><strong>20</strong></td>
</tr>
</tbody>
</table>

* Could go down to 0.04 - 0.1 µl if needed according to invitrogen protocol

2. Add 0.5 - 4 µl of DNA sample to each PCR tube
   0.5 µl for purified plasmid to 4 µl for colony PCR from liquid culture
3. Add Master Mix to each PCR tube
4. If you did not add your primers to the master mix add them now
5. If using the old Perkin Elmer thermocycler add 20 µl of mineral oil to top
6. Place your PCR tubes in the thermocycler and run your program
7. While you are waiting for the PCR you can make a gel to check your results
8. In a 250 - 500 ml bottle or flask make up ≈ 1 % agarose solution
   For small gels add 0.5 g agarose to 50 ml 1X TAE
   For large gels add 2 g agarose to 200 ml 1X TAE
9. Microwave agarose and TAE buffer for 1 minute or until boiling
10. Set up gel mold in empty try by placing combs in mold and clamp mold tight
11. Slowly pour agarose TAE solution into gel mold, allow to cool and solidify for 30 min
12. When the agarose gel has solidified carefully and slowly remove comb and place in electrophoresis apparatus, don’t worry if some of the wells are stuck closed they will open when you add TAE buffer in the next step. You can also store your gel in the refrigerator overnight. If your using a mold or comb that is in “high demand” remove the gel from the mold too. Place the gel in a plastic bag so that it does not dry out.
13. Place your gel in the electrophoresis system and fill with 1X TAE to the fill line, you can use the TAE solution that is already
in the system, no need to drain and refill.

14. When your PCR is finished, add 5 µl of loading buffer to 20 µl DNA in separate PCR tube
15. Mix 10 µl of 1 Kb ladder with 2.5 µl EZ-vision loading dye
16. Place 5 – 20 µl of PCR product with loading buffer into each well.
17. After loading place cover on electrophoresis system and run gel at approximately 85 V for 30 – 45 minutes
18. Turn power supply off and remove cover. Carefully place gel in an empty tray to carry to the imager.

1 The program you will run in the thermocycler will depend on the size of your primers and the size of the expected product, and the Taq polymerase you are using, below is an example program

**Typical Program**
1. 95°C – 2 min *1
2. 95°C – 30sec
3. 55°C*2 – 30sec
4. 72°C – 1.5 min*3
5. Cycle to step 2 35 more times
6. 72°C – 4.5 min (3 * the extension time length)
7. Incubate at 4°C forever

*1 The initial melting temperature may be as long as 7.5 minutes for colony PCR to break open cells

*2 The annealing temperature is the most critical, a rule of thumb is to subtract 5°C from the Tm given to you by the primer 3 web program or you can check this temp by entering your primer sequence in promega’s website program getting the “base stacking Tm” subtracting 2 - 3°C and using that as your annealing temperature

http://fokker.wi.mit.edu/primer3/input.htm
http://www.promega.com/biomath/calc11.htm

*3 The extension time should be ≈ 1 kb per minute

2 The % of agarose in your gel will be determined by the size of the PCR product you are trying to isolate. The smaller the product size (in base pairs) the higher the % of agarose you will need to use. In general 0.5% = 20 kbp – 2 kpb, 1% = 10kpb – 500bp, 1.5% = 5kbp – 250bp, 2% 2.5kbp – 100bp

**PCR Reagents**

**10 X TAE Buffer 1L**
+ 48.4 g Tris
+ 7.44g Na2EDTA * 2H2O
+ 11.42 ml glacial acetic acid
+ 100 µl 0.5 M EDTA
Add H2O up to 1 L
pH to 8.5
Note – Do not make up more then 1 L it will go bad before you can use it all

**1 X TAE Buffer 1 L**
In 900 ml dH2O
+ 100 ml 10 X TAE buffer (40 mM Tris)

**100 µM Primers ≈ 1 ml**
Multiply 10 * the nmol amount of primer printed on the side of the tube
This is the amount of sterile dH2O you should add to the tube
Mix by up and down pipetting try not to vortex
store in -20°C

**10 µM Primers 500 µl**
In 450 µl sterile dH2O
+ 50 µl of 100 µM primer
store in -20°C