

Oct 23, 2019

PCR V.2

Claudia Troncone Clemente¹

¹Universidad Complutense de Madrid

1 Works for me dx.doi.org/10.17504/protocols.io.8nyhvfw

AEGIS - Madrid iGEM 2019



ABSTRACT

Our aim with this protocol is to amplify DNA. This protocol has been optimized has a general amplification

As the quantity of DNA is exponentially increased during the performance of the selection, further modification in the numbers of cycle will be needed to be implemented.

GUIDELINES

We sure to have all the surfaces and materials clean before the start.

All the procedures must be done in an sterile environment to avoid contamination.

MATERIALS

NAME ~	CATALOG #	VENDOR ~
Speedy Supreme Green Master Mix	MB39102	NZYtech
Agarose (LM-ultrapure grade)	MB123	NYZtech

MATERIALS TEXT

- Aptamer library (order to IDT)
 - 5'- G TTG CTC GTA TTT AGG GAA TG $N_{\tiny{10}}$ ACA CCA GTC TTC ATC CGC TTT $_{\tiny{6}}$ 3
- Forward primer (order to IDT):
 G TAG GCG AAA₆ Cy3 5`
- Reserve primer (order to IDT):
 - 5`- BiodTG TTG CTC GTA TTT AGG GAA TG
- Thermocycler
- TAE buffer

1 Prepare the PCR reaction mixture following the specifications below:

Component	Positive control (V; ul)	Negative control (V; ul)
Template	5	0
Fwd primer	1.25	1.25
Rev primer	1.25	1.25
dH2O	15	20
DMSO	2.5	2.5
Master Mix	25	25

2 Perform the amplification in a general thermocycler in the following conditions. Adjust the annealing temperature according to the primers used, and the hotstart to the specifications of your polymerase:

Hot start	95 º	5 min		
Amplification cycles				
Denaturing	95 º	30s		
Annealing	52 °	30s	X 15 cycles	
Extension	72 º	30s		
Final extension	72 º	3 min		
Hold	40			

- 3 Prepare a 3% agarose gel. Load the samples and perform the electrophoresis at 90V for 50 min.
- 4 Remove the gel and observe the bands under UV light.

This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited