03_04_PCR-for-ssDNA-samples

MITTWOCH, 2.6.2021

Goal setting

• Performing a PCR reaction using the Phusion™ High-Fidelity DNA Polymerase

Terms / abbreviationsTerms / abbreviations

- DMSO = Dimethyl sulfoxide
- PCR = Polymerase chain reaction

Risk areas

• If spilled, always wipe surface with alcohol



Required materials/information

- Chemicals:
 - o 5x Phusion HF buffer, ThermoFisher
 - o 5x Phusion GC buffer, ThermoFisher
 - o 10 mM dNTP, ThermoFisher
 - o 10 μM Forward primer, Ella Biotech
 - $\circ~$ 10 μM Reverse primer, Ella Biotech
 - o 50 mM MgCl₂ solution, ThermoFisher
 - o Autoclaved MilliQ water, Sartorius arium pro VF
 - o DMSO, ThermoFisher
 - Phusion DNA Polymerase (2 U/μL), ThermoFisher
 - Template DNA
- Material:
 - o lce or cooling rack
 - o PCR tubes, Sarstedt
 - o Trash bags, Th. Geyer GmbH & Co. KG

Templates, devices, software

- Nanodrop spectrophotometer, ThermoFisher
- Thermocycler, Eppendorf
- Pipettes, Eppendorf

Preliminary work

- Any reaction creating DNA that should be amplified
- 04_01_Spectralphotometer-NanoDrop to detected desired amount of template DNA

Operation

- 1. Put the PCR tube on ice or the cooling rack
- 2. Add the components in the order listed in the following table

Composition of Master Mix									
	А	В	С	D	Е				
1	Component	20 [μL]	50 [μL]	Final concentration	Unit				
2	MilliQ water	add to 20 μL	add to 50 μL						
3	5X Phusion HF Buffer	4	10	1	x				
4	10 mM dNTPs	0.4	1	200	μΜ				
5	Forward primer 10 μM	1	2.5	0.5	μΜ				
6	Reverse primer 10 μM	1	2.5	0.5	μΜ				
7	Template DNA *	×	×	0.2					
8	Phusion High-Fidelity DNA Polymerase	0.2	0.5	0.02	U/µl				

 $^{^{*}}$ use 1 μ L from standard TdT reaction as template

3. Run the PCR program in the thermocycler according to the following table

PCR Program for Thermocycler									
	А	В	С	D					
1	Temperature [°C]	Time	Unit	Repeats					
2	98	30	S	1x					
3	98	5	S						
4	63	10	S	30x					
5	72	30	S						
6	72	5	min	1x					

Increase elongation time for fragments > 1 kb

Disposal

• Autoclave trash bags, discard in S1 waste

Troubleshooting

- Wear gloves to reduce the risk of DNase and RNase contamination
- The Phusion DNA Polymerase should be pipetted carefully and gently as the high glycerol content (50%) in the storage buffer may otherwise lead to pipetting errors

Sources

MAN0012393_Phusion_HighFidelity_DNAPolymerase_UG.pdf

Follow-up work

- Purify fragments with a cleanup-kit
- If materials are empty, care about new order