iGEM 2013 – SDU	
Title: Colony PCR with MyTaq	Date issued: 2012.11.26
SOP number: SOP0021_v01	<b>Review date:</b> 2013.12.03
Version number: 01	Written by: Tina Kronborg

### 1. Purpose

Colony PCR with My Taq

### 2. Area of application

This procedure is valid for all E. coli strains

### 3. Apparatus and equipment

Apparatus/equipm ent	Location (Room number)	Check points	Criteria for approval/rejection
Heating block	Laboratory (class 1) – V18-404-2 Laboratory (class 2) – V18-501b-2	<ul> <li>Preheat to 96</li> <li>°C</li> </ul>	
PCR machine	Laboratory (class 1) – V18-403b-2	<ul> <li>Check program</li> </ul>	Appropriate PCR program
Pipettes (p100,20,10)			
Container for ice			
PCR tube rotator	Laboratory (class 1) – V18-403b-2		

### 4. Materials and reagents - their shelf life anod risk labelling

Name	Components (Concentrations)	Manufac turer/Cat . #	Ro om	Safety consid eration s
Water	Demineralised milli-Q autoclaved water	Milli-Q water purificati on system (Millipor e)	RT	
MyTaq™HS Red Mix	http://www.bioline.com/documents/product_inser ts/MyTaq%E2%84%A2%20HS%20Red%20Mix.pdf#z oom=130	Bioline	V18 -40 5a- 2	
Reverse primer	Made specific to the template	Sigma-Al drich		
Forward primer	Made specific to the template	Sigma-Al drich		
PCR tubes		Eppendo rf	Mi cro sto rag e	
1.5 ml tubes		Contact lab-mana ger	BM B sto rag e	
Ice			V1 8-4 03 a-2	
Green pipette tips		Contact lab-mana ger	Mic ro sto rag e	
Purple pipette tips		Contact lab-mana ger	Mi cro sto rag e	

# 5. QC – Quality Control

For more than 2 samples a premix of primers, MyTaq, primers and water is mixed and aliquoted before adding template

#### Designing primers:

Generally, the annealing temperature is about 5°C below the lowest melting temperature  $(T_m)$  of the pair of primers used, and should be around 55°C so  $T_m$  should be around 60°C.

 $T_m = 4(G + C) C + 2(A + T) C$ 

Each primer should be about 20-30 nucleotides.

#### 6. List of other SOPs relevant to this SOP

JMJ\_SOP0006\_v01\_MM\_Agarose\_gel\_DNA

#### 7. Environmental conditions required

When the bacteria samples have been boiled they can be removed from class II and transferred to class I

#### 8. Procedure

- 1. Add 50 μl sterile water to a 1.5 ml tube one for each sample
- 2. Dip a green tips in the colony and transfer it to the eppendorf tube
- 3. Boil at 96 °C for 5 min in a heating block
- 4. Or a colony sample can be transferred into a PCR-tube and with open lid be microwaved for 2 min at max effect
- 5. Place the samples on ice
- 6. Mix primers, water and MyTaq as described under PCR set up; paragraph 12 with the template
- 7. Be sure that all the samples are in the bottom of the PCR tubes by spinning on PCR tube rotator
- 8. Place in PCR machine
- 9. Start the appropriate PCR program for MyTaq or design one yourself, see paragraph 12
- 10. Keep at 4-5 °C until use, if more than 2 days waiting time place in -20°C

#### 9. Waste handling

Chemical name	Concentration	Type of waste (C, Z)	Remarks
Boiled bacteria waste		GMO yellow waste	
Once used plastic		GMO yellow waste	

### 10. Time consumption

- Total-time 2 hours
- Hands-on-time 1 hour

## 11. Scheme of development

Date / Initials	Version No.	Description of changes
12.11.26 / TK	01	The SOP has been written
13.05.22 / TK & MM	01	The SOP has been approved

## 12. Appendixes

### PCR set up, 1 sample:

#### Notes: Smaller volume than recommended by Bioline

Template:	200 ng	Template: 0.5 $\mu$ l of 50 $\mu$ l sterile water boiled with some of the bacteria colony or mix directly into the PCR tube with the microwaved colony
Primers (20 pmol):	0.2 μl each	
MyTaq HS Red Mix, 2x:	5 μΙ	
Water (sterile):	4.1 μl	
Total:	10 μl	

# PCR cycling conditions:

Step 1: Initial denaturation:	95 °C	2 min
Step 2: Denaturation:	95 °C	15 sec
Step 3: Annealing:	55 °C	15 sec (depending on the primer sequences,
		2-5 °C below the lowest Tm of the primers)
Step 4: Extension/Elongation:	72 °C	10-30 sec (30 sec. pr. kilo bp)
Step 5: Repeat step 2-4:		30 times
Step 6: Extra elongation:	72 °C	2.5 min
Step 7: Keep the samples cold	4 °C	until the samples is removed