# iGEM 2013 – SDU

Title: ON culture of <i>E. coli</i>	Date issued: 2012.10.25
SOP number: SOP0001_v01	<b>Review date:</b> 2013.12.01
Version number: 01	Written by: Michelle Madelung

# 1. Purpose

To prepare ON culture of *E.coli* for use in experiments

### 2. Area of application

This procedure is valid for all *E. coli* ON cultures

# 3. Apparatus and equipment

Apparatus/equip ment	Location (Room number)	Check points	Criteria for approval/rejection
Incubator	Laboratory (class 1) - V16-407-2 Laboratory (class 2) – V15-501b-2	<ul> <li>Preheated</li> </ul>	37°C
Spectrometer	Laboratory (class 1) - V16-407-2 Laboratory (class 2) – V15-501a-2	<ul> <li>Set to wavelength 600</li> </ul>	
Vortex	Laboratory (class 1) Laboratory (class 2) – V15-501a-2	•	
Pipette boy		<ul> <li>Remember to recharge</li> </ul>	
Racks		•	
Sterile glass culture tube	Laboratory (class 1) – opposite elevator Laboratory (class 2) – V16-501a-2 filing cabinet	•	
Refrigerator		•	
Pipettes (p1000,200)		•	

# 4. Materials and reagents – their shelf life and risk labelling

Name	Component s	Supplier / Cat. #	Room (hallway storage)	Safety considerations
Appropriate medium ex. LB	1% Tryptone 1% NaCl 0.5% Yeast extract	Oxoid Sigma-Aldrich Merck	Media lab or V18-405-0	
Appropriate antibiotic if needed				
5 ml graduated pipettes		Fisher Scientific / CCI 4487	Micro storage	
Cuvettes		Contact lab-manager	BMB storage	
Blue pipette tips		Contact lab-manager	Micro storage	
Green pipette tips		Contact lab-manager	Micro storage	
Plate Bag		Contact lab-manager	BMB storage	

### 5. QC – Quality Control

When measuring  $OD_{600}$ , measurements can't go above 0.300, if this is the case dilute solution with medium 10 times.

6. List of other SOPs relevant to this SOP

### 7. Environmental conditions required

#### 8. Procedure

- 8.1 Take an agar plate with appropriate medium and antibiotic
- 8.2 Scrap surface of frozen bacterial stock
- 8.3 Streak this bacterial stock onto agar plate (primary streak)
- 8.4 Take a new pipette tip streak again (secondary steak)
- 8.5 Take a new pipette tip streak again (tertiary streak)
- 8.6 Place the plate in a plate bag
- 8.7 Leave 16 hours in incubator set to 37 °C
- 8.8 Move plate to refrigerator (4 °C)
- 8.9 Fill 5 ml medium in culture tube
- 8.10 Add antibiotic to appropriate concentration
- 8.11 Take single colony from agar plate
- 8.12 Vortex medium with colony
- 8.13 Place culture tube in incubator set to 37 °C with aeration (155 rpm)
- 8.14 Leave 16 hours
- 8.15 Add 1 ml medium to a cuvette and calibrate spectrometer
- 8.16 Add 0.9 ml medium to a cuvette and 0.1 ml ON culture and mix
- 8.17 Measure OD

### 9. Waste handling

Chemical name	Concentration	Type of waste (C, Z)	Remarks
ON Culture		Liquid bacterial waste	
Once use plastic		GMO yellow waste	

#### **10.** Time consumption



- Total-time 30 hours and 20 min.
- Hands-on-time 30 min.

### 11. Scheme of development

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

# 12. Appensdixes