# Media preparation

Version 1.4

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#### General notes on autoclaving

- Place lid on bottle, screw until it starts gripping and then turn it back a full rotation. Then tape the lid in this position with a piece of autoclaving tape. This is to ensure that expanding gases can escape the bottle and it doesn't explode
- We usually run a short 15/20 minute liquid cycle (number 006), but longer liquid cycles are permissible

## Minimal Marine Sea Water (MSW) + D-Gluconic Acid 0.1% (1 liter)

- 1. Autoclave one empty 1 L bottle and 1 L of MilliQ in a separate bottle
- 2. After autoclaving, add MilliQ and magnetic stir bar to beaker. Then, add the following chemicals (use high-precision scale in antibiotic station for the last two ingredients):
  - 1.0g D-Gluconic acid sodium salt
  - 1.0g KH<sub>2</sub>PO<sub>4</sub>
  - 1.0g  $H_4 \bar{N}_2 O_3$
  - 10.0g NaCl
  - $0.2g \text{ MgSO}_4 \cdot 7H_2O$
  - $10 \text{mg FeSO}_4 \cdot 7 \text{H}_2 \text{O}$
  - 10mg CaCl<sub>2</sub>·2H<sub>2</sub>O

## 3. Adjust pH to 7

- 1. Turn on pH meter, take electrode out of storage buffer, rinse with MilliQ water and carefully dry with lab wipes
- 2. While stirring, lower the electrode into the liquid medium and press the measure button
- 3. Add NaOH slowly until the pH reaches 7 (6.95 7.05 are permissible). 1000 mL medium require  $\sim 1000$  uL of NaOH
- 4. When done, rinse and dry the electrode before putting in back into the storage solution and turninig off the pH meter

#### 4. Sterile filtration

- 1. Inside a sterile hood, screw the sterile filter onto the autoclaved empty bottle
- 2. Back in J18, turn on the vacuum pump and stick the hose onto the sterile filter nozzle
- 3. Add the liquid medium (and any remaining autoclaved water) to the sterile filter
- 4. First take the hose off the nozzle, then turn off the vacuum pump
- 5. Finally, screw off the sterile filter in a hood and close the bottle with its cap

#### 5. Clean up and storage

- 1. Wipe bench and scales
- 2. Turn off stirring plate, scales and pH meter (if not off already)
- 3. Put away dry beakers, spatulas and weighing boats before washing your own ones
- 4. Label liquid medium bottle with "MSW + D-Gluc.", your name and the date
- 5. Store in the fridge at 4°C

## Minimal Marine Sea Water Agar 1% (MSWA) + D-Gluconic Acid 0.1% (1 liter)

- 1. Dissolve 15g of BactoAgar in 800 mL of MilliQ, then add to 1 L bottle
- 2. Autoclave agar and 200 mL of MilliQ in a separate (e.g. 250 mL) bottle
- 3. After autoclaving, add 200 mL MilliQ and magnetic stir bar to beaker. Then, add the following chemicals (use high-precision scale in antibiotic station for the last two ingredients)
  - 1.0g D-Gluconic acid sodium salt
  - 1.0g KH<sub>2</sub>PO<sub>4</sub>
  - $1.0g H_4 N_2 O_3$
  - 10.0g NaCl
  - $0.2g MgSO_4 \cdot 7H_2O$
  - 10mg FeSO<sub>4</sub>·7H<sub>2</sub>O
  - $10 \text{mg CaCl}_2 \cdot 2 \text{H}_2 \text{O}$

#### 4. Adjust pH to 7

- 1. Turn on pH meter, take electrode out of storage buffer, rinse with MilliQ water and carefully dry with lab wipes
- 2. While stirring, lower the electrode into the liquid medium and press the measure button
- 3. Add NaOH slowly until the pH reaches 7 (6.95 7.05 are permissible). 1000 mL medium require  $\sim 1000$  uL of NaOH
- 4. When done, rinse and dry the electrode before putting in back into the storage solution and turninig off the pH meter

#### 5. Sterile filtration

- 1. Inside a sterile, screw the sterile filter onto the autoclaved bottle containing  $800\,$  mL agar
- 2. Back in [18, turn on the vacuum pump and stick the hose onto the sterile filter nozzle
- 3. Add the liquid medium (and any remaining autoclaved water) to the sterile filter
- 4. First take the hose of the nozzle, then turn off the vacuum pump
- 5. Finally, screw off the sterile filter in a hood and close the bottle with its cap. Mix well by swirling and inverting

#### 6. Clean up and storage

- 1. Wipe bench and scales
- 2. Turn off stirring plate, scales and pH meter (if not off already)
- 3. Put away dry beakers, spatulas and weighing boats before washing your own ones
- 4. Label medium bottle with "MSWA + D-Gluc.", your name and the date. Preferably pour agar plates immediately (otherwise heat-sensitive components might decay). Otherwise store in media incubator at 50°C

## Marine broth (MB) (1 liter)

- 0. First, check the Marine broth 2216 (Sigma Aldrich) bottle for instructions. If they exist, follow those instead of this protocol.
- 1. Add 37.4 g of Marine broth 2216 powder to roughly 900 mL of MilliQ. Once dissolved, fill volume up to 1000 mL, then add to 1L bottle
- 2. Autoclave and store in the fridge at 4°C

## Marine agar (MA) (1 liter)

- 0. First, check the Marine broth 2216 (Sigma Aldrich) bottle for instructions. If they exist, follow those instead of this protocol
- 1. Add 15 g of BactoAgar to roughly 900mL of MilliQ
- 2. Add 37.4 g of Marine broth 2216 powder the beaker. Once dissolved (excluding the agar), add to 1 L bottle
- 3. After autoclaving, store bottle in 50°C incubator to prevent solidifying

## Casitone-Tris (CTT) liquid (1 liter)

- 1. Dissolve the following components in 900 mL MilliQ:
  - 10 mL 1.0 M Trish-HCl (pH 8.0)
  - 10 mL 0.8 M MgSO<sub>4</sub>
  - 10g Bacto Casitone
- 2. Fill up volume to 1 L with MilliQ and adjust pH to 7.6
- 3. Autoclave using cycle 003 (liquid media, 50 min at 121°C)
- 4. After cooling <50°C add 1 mL 1 M KH<sub>2</sub>PO<sub>4</sub>-K<sub>2</sub>HPO<sub>4</sub>

## Phosphate-Buffered Saline (PBS; 1 liter of 10x solution)

Instructions according to Sigma

- 1. Dissolve the following components in 900 mL MilliQ:
  - 17.8 g of Na<sub>2</sub>HPO<sub>4</sub>·2H<sub>2</sub>O
  - 2.4 g of KH<sub>2</sub>PO<sub>4</sub>
  - 80 g of NaCl
  - 2 g of KCl
- 2. Adjust final volume to 1 L with MilliQ
- 3. Measure pH using the pH meter. The pH of the 10X stock will be approximately 6.8, but when diluted to 1x PBS the pH should change to 7.4
- 4. Autoclave or filter-sterilize into a sterile bottle
- 5. Store in the fridge at  $4^{\circ}C$