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♠ A green micro-algal growth media modified for use as a stringent minimal media for bacteria.

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1 Works for me dx.doi.org/10.17504/protocols.io.bgzujx6w

ABSTRACT

Introduction:

Chlamydomonas reinhardtii, a green micro-alga can be grown at the lab heterotrophically or photo-heterotrophically at room temperature using the <u>Tris-Phosphate-Acetate</u> (TAP) medium which contains 0.1 % acetate (acetic acid) as the sole carbon source. Standard TAP medium recipe can be found at the website of Chlamydomonas Resource Center: https://www.chlamycollection.org/methods/media-recipes/tap-and-tris-minimal/. Hutner's trace element solution is an ingredient in the TAP medium. Hutner's trace element recipe can be found at https://www.chlamycollection.org/methods/media-recipes/hutners-trace-elements/. M9 medium is the standard minimal medium for growing bacteria (https://www.thelabrat.com/protocols/m9minimal.shtml). When grown in TAP medium, https://www.thelabrat.com/protocols/m9minimal.shtml). When grown in TAP medium, <a href="https://w

In our lab we use a slightly modified TAP medium recipe which has a final concentration of phosphate, nitrogen, magnesium and calcium approximately 10-fold, 2-fold, 2-fold and 2-fold higher than that in the TAP recipe described on the Chlamydomonas Resource Center website, respectively. Final concentrations of acetate and Hutner's trace elements are same in both TAP recipes. Attached Table 1 compares the chemical ingredients in our lab's TAP recipe (full recipe can be found on protol.io) with that present in the standard M9 medium described at http://www.thelabrat.com/protocols/m9minimal.shtml. M9 medium has a final concentration of phosphate, nitrogen, magnesium and carbon, approximately 70-fold, 2.5-fold, 4-fold and 5-fold higher than that present in our lab's TAP medium, respectively. Additionally, M9 contains 0.05% salt (8.56 mM) (Table 1). TAP has additional trace elements like iron, zinc, copper, manganese, cobalt, boron and molybdenum which are components in the Hutner's trace element solution (Table1). In summary, TAP medium is chemical composition wise, a more stringent minimal medium than the M9 medium used by microbiologists.

Applications:

There are many bacteria that can also utilize acetate as a carbon source. Some of these bacteria also have glyoxylate cycle like *Chlamydomonas* and green plants. Hence, TAP medium can be used to isolate acetate-requiring bacteria. If acetate is removed from the TAP medium, it becomes the TP medium which lacks a carbon source and will not allow Chlamydomonas or other acetate-requiring bacteria to grow. The 0.1 % acetate in the TAP medium can be substituted with alternative carbon sources (e.g. glucose, sucrose, lactose, hydrocarbons, aromatic compounds and polyhydroxyalkanoates etc.) to test physiological abilities of candidate bacteria to use the tested chemicals as the sole alternative carbon and energy source.

EXTERNAL LINK

https://f1000research.com/articles/9-656/v1

THIS PROTOCOL ACCOMPANIES THE FOLLOWING PUBLICATION

Mitra M, Nguyen K, Box T et al. Isolation and characterization of a novel bacterial strain from a Tris-Acetate-Phosphate agar medium plate of the green micro-alga Chlamydomonas reinhardtii that can utilize common environmental pollutants as a carbon source [version 1; peer review: awaiting peer review]. F1000Research 2020, 9:656

ATTACHMENTS



06/12/2020

Citation: Mautusi Mitra (06/12/2020). A green micro-algal growth media modified for use as a stringent minimal media for bacteria.. https://dx.doi.org/10.17504/protocols.io.bgzujx6w

Table 1.pdf

DOI

dx.doi.org/10.17504/protocols.io.bgzujx6w

PROTOCOL CITATION

Mautusi Mitra 2020. A green micro-algal growth media modified for use as a stringent minimal media for bacteria.. **protocols.io**

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MANUSCRIPT CITATION please remember to cite the following publication along with this protocol

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KEYWORDS

TAP medium, M9 medium, Chlamydomonas reinhardtii, alternative carbon source, bacteria grown in TAP medium, TP medium, acetate-consuming bacteria, alternative carbon souces, bacterial minimal medium

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GUIDELINES

Do not use NaOH for pHing Hutner's trace element. Use KOH pellets and solution for pHing Hutner's trace element.

MATERIALS

NAME	CATALOG #	VENDOR
Cobaltous chloride hexahydrate		
Zinc sulfate heptahydrate	204986	Sigma Aldrich
Boric acid	BP1681	Fisher Scientific
Manganese chloride	7773-01-5	Fisher Scientific
Ammonium molybdate (VI) tetrahydrate	12054-85-2	Fisher Scientific
Calcium chloride, dihydrate	CD0050.SIZE.500g	Bio Basic Inc.
Copper (II) sulfate pentahydrate	CDB0063.SIZE.500g	Bio Basic Inc.
Iron (II) sulfate, heptahydrate	FB0461.SIZE.500g	Bio Basic Inc.
Potassium hydroxide	PB0441.SIZE.500g	Bio Basic Inc.
Magnesium Sulfate Heptahydrate, ACS Grade	M-020	Gold Biotechnology



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NAME	CATALOG #	VENDOR
Acetic acid Glacial	A38-212	Fisher Scientific
Tris Base	604204	Fisher Scientific
Potassium phosphate Dibasic		Sigma Aldrich
Potassium Phosphate, Monobasic, Molecular Biology Grade, Calbiochem™, , 1kg	5295681KG	Thermo Fisher
Ammonium Chloride	A661-500	Fisher Scientific

BEFORE STARTING

You can purchase all chemicals needed to make the TAP medium from any vendor that sells them and do not have to use the vendors that I have selected under "Materials".

Preparation of TAP stock nutrients

Make TAP salts (TAP stock nutrients/FBS solution [Filner's Beijerinck solution X40]) by dissolving Ammonium chloride (16g), Magnesium sulfate (4g) and Calcium chloride (2.65g). This will give you a final concentration of: Ammonium chloride (7.48 mM), Magnesium sulfate (16.2 mM) and Calcium chloride (18.10 mM).

Preparation of Phosphate solution

2 Make Phosphate solution (1M Potassium phosphate solution, pH7) by dissolving 28.8 g of K₂HPO₄ and 14.4g of KH₂PO₄. Note: You don't have to adjust pH for this solution as it automatically gives a pH of 7-7.2 when mixed in the above stated ratio. Phosphate solution has the final concnetration of K₂HPO₄ (1.65M) and KH₂PO₄ (1.058M).

Preparation of Hutner's trace element

Preparation of Hutner's trace element (X 1000) (Note: If you do not want to make it, you can purchase this solution from Chlamydomonas Resource center at at https://www.chlamycollection.org/methods/media-recipes/hutners-trace-elements/). In order to make this solution, dissolve the following salts in order in 800 mL of E-pure water—dissolve each fully before adding the next.

Salt	Molecular	Final	Amount
	Weight	concentration in 1L	to be added in 1L
FeSO4.7H2O	278.01	18 mM	4.99 g
ZnS04.7 H20	287.56	76.5mM	22 g
H3B03	61.83	184 mM	11.4 g
MnCl2.4H2O	197.91	25.6 mM	5.06 g
CuSO4.5H2O	249.68	6.3mM	1.57 g
(NH4)6Mo7024.	1235.86	0.89mM	1.10 g
4 H2O			
CoCl2.6H2O	237.93	6.8mM	1.61 g

Table of chemical ingredients needed to make Hutner's trace element.

- 4 Bring the salt mixture to a slow boil. Add 50 grams of disodium salt of EDTA to the boiling mixture, acid form (Na₂ EDTA.2H₂O; molecular weight is 372.24; final concentration in the 1L solution is 134 mM).
- Add KOH pellets (and not NaOH) to the boiling mixture to adjust the pH to 6.5. Make up the volume to 1L with pure water after adjusting the pH to 6.5. The solution should be clear green at this point.

 Pour the green solution in a 1 L bottle. Close the cap not too tightly. Shake it occasionally every week. The color will slowly change to dark magenta/purplish color over time. If you see any brown precipitate, filter the solution through two layers of Whatman#1 filter paper, repeating the filtration if necessary until the solution is clear.

Preparation of TAP medium

- 7 To make the final TAP medium, mix the following in 950 L of water: 2.42 g Tris base/Trizma, 25 ml solution #1 (salts), 0.375 ml solution #2 (phosphate), 1.0 ml solution #3 (trace elements) and 1.0 ml glacial acetic acid (0.1% in the final 1 L of TAP medium).
- 8 Check the pH of the solution before making up the volume to 1L. pH of TAP medium should be approximately between 7-7.2 once you make it but if it is not, adjust pH with acid/base. The final concentration sof all chemical ingredients that you have added to make TAP medium is given in the attached Table 1 pdf.

Preparation of TAP-agar plates

9 For preparing solid TAP-agar media plates to maintain *Chlamydomonas* in lab, add 15 grams of agar per liter of the media after checking the pH, shake it and then autoclave. **Note:** Make sure to have a magnetic stir bar inside the agar media bottle before you autoclave the bottle. Stirring using a magnetic stirrer is required during the cooling of the hot media after autoclaving to mix agar uniformly in the media solution before pouring media plates.

Preparation of TP medium

TP medium is prepared exactly the same way as the TAP medium, except glacial acetic acid is not added. Hence TP medium has no carbon souce. Nothing will grow on it unless you add a carbon source to the TP medium. You will pH the TP medium like you would for making TAP medium. You can add your preferred alternative carbon source (e.g. glucose, sucrose, lactose, hydrocarbons, aromatic compounds and polyhydroxyalkanoates etc.) for testing biochemical abilites of bacteria to use exogenous carbon sources for energy production and growth.