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Protocol status: Working
We use this protocol and it's working

Created: Mar 03, 2022

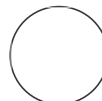
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Seahorse protocol for islets using Xfe24 Analyzer

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ABSTRACT

Using Agilent's seahorse Xfe Analyzer with the islet capture microplates to assess whole islet bioenergetics in vitro. Agilent Seahorse XFe24 Analyzers measure the oxygen consumption rate (OCR) and extracellular acidification rate (ECAR) of live cells in a 24-well plate format.

PROTOCOL integer ID:
59039

MATERIALS

Seahorse XF24 Islet Capture Microplates cat #[101122-100](#), Agilent Technologies

Seahorse XF Calibrant Solution 500 mL cat#[100840-000](#), Agilent Technologies

Seahorse XF DMEM medium, pH 7.4, 500 mL cat#[103575-100](#), Agilent Technologies

D-glucose 1KG, cat#G8270, Sigma-Aldrich

Fetal bovine serum (FBS) 500ml cat#12483-020, Gibco

Sodium Pyruvate (100mM), 100ml, cat#11360-070, Gibco

L-glutamine (200mM), 100ml, cat#25030-081, Gibco

MilliporeSigma™ Stericup™ Quick Release-GP Sterile Vacuum Bottle Top Filtration Systems cat#S2GPU05RE, Fisher Scientific

Oligomycin from *Streptomyces diastatochromogenes*, cat#04876-5MG, Sigma-Aldrich

FCCP, Carbonyl cyanide 4-(trifluoromethoxy)phenylhydrazone, cat#C2920-10MG, Sigma-Aldrich

Rotenone, cat#R8875-5G, Sigma-Aldrich

Antimycin A from *Streptomyces* spp., cat#A8674-25MG, Sigma-Aldrich

DMSO, Dimethyl Sulfoxide, cat#D128-1, Fisher Scientific

Solution Prep - Calibrant Solution

- 1 Calibrant Solution (Agilent 100840-000, 500ml) is provided by Agilent and stored at room temperature.

Solution Prep - MA media

- 2 Use the DMEM (Agilent 103575-100, 500ml) provided by Agilent; stored at 4 °C. Needs to be supplement with 1% Fetal Bovine Serum (FBS, Gibco 12483-020). Add 5ml of heat inactivated FBS to the 500ml

DMEM. Filter sterilize and store at 4°C.

- 2.1** On the day of experiment, supplement with 2mM or 2.8mM D-glucose (Sigma G8270), 2mM sodium pyruvate (Gibco 11360-070) and 2mM L-glutamine (Gibco 25030-081). Add 0.018g or 0.025g of glucose to 50ml MA media. Pipette 1ml of (100mM) sodium pyruvate and 500ul of (200mM) L-glutamine to the 50ml MA media. Mix until glucose has dissolved.

Solution Prep - Stocks

- 3** Glucose 2M stock in Dimethyl sulfoxide (DMSO, Fisher D128-1) - A

Weigh 18.2g of glucose into 50ml DMSO. Aliquot and store at -20°C.

- 4** Oligomycin (Sigma 04876-5MG) 5mM stock in DMSO - B

Pipette 1.27ml DMSO to the 5mg Oligomycin. Aliquot and store at -20°C.

- 5** FCCP (Carbonyl cyanide 4-(trifluoromethoxy)phenylhydrazone, Sigma C2920-10MG) 1mM stock in DMSO - C

Prepare 50mM stock of FCCP by adding 787ul of DMSO to the vial containing 10mg of FCCP. To make 1mM stock, use 20ul of the 50mM FCCP into 980ul of DMSO. Aliquot the 1mM stock and store both stocks at -20°C.

- 6** Rotenone (Sigma R8875-5G) 5mM stock in DMSO - D

Weigh 1.97g of Rotenone into 1ml of DMSO. Aliquot and store at -20°C.

- 7** Antimycin A (Sigma A8674-25MG) 5mM stock in DMSO - D

Weigh 2.7mg of Antimycin A into 1ml DMSO. Aliquot and store -20°C.

Solution Prep - working stocks

- 8** Glucose (167mM) - A

Pipette 167 μ l of 2M glucose stock to 1833 μ l MA media. Working stock concentration of 167mM with an injected final concentration of 16.7mM.

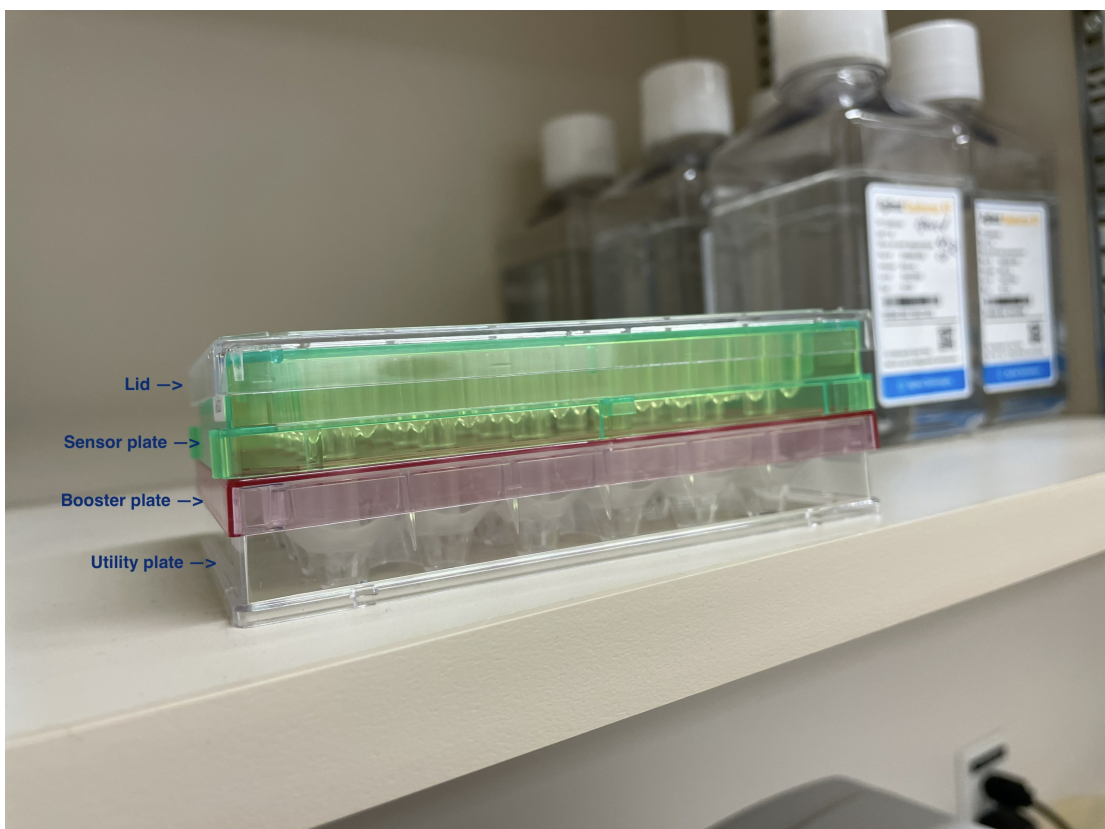
- 8.1** Glucose (200mM) - A
Pipette 200 μ l of 2M glucose stock to 1800 μ l MA media. Working stock concentration of 200mM with an injected final concentration of 20mM. **
- 9** Oligomycin (55uM) - B
Pipette 22 μ l of 5mM oligomycin stock to 1978 μ l MA media. Working stock concentration of 55uM with an injected final concentration of 5 μ M.
- 10** FCCP - C
Mouse (55.5 μ M)
Pipette 111 μ l of 1mM FCCP stock to 1889 μ l MA media. Working stock concentration of 55.5 μ M with an injected final concentration of 5 μ M.
Human (33.3 μ M)
Pipette 66.6 μ l of 1mM FCCP stock to 1933.4 μ l MA media. Working stock concentration of 33.3 μ M with an injected final concentration of 3 μ M.
FCCP needs to be titrated for every lot
- 10.1** FCCP cat#C2920-10MG, Sigma, lot#0000120405
optimal concentration on mouse islets = 5 μ M
optimal concentration on human islets - 3 μ M
- 11** Rotenone (55.2 μ M)/Antimycin A (56.2 μ M) - D
Pipette 22.5 μ l of 5mM rotenone stock, 22.5 μ l of 5mM antimycin A stock to 1955 μ l MA media. Working stock concentration of 55.2 μ M with an injected final concentration of 5 μ M

Day one Protocol

- 12** The day before the assay, add 1 ml of calibrant solution to each of the wells of the utility plate of the extracellular flux assay kit.



Incubate the entire set of plates at 37°C with NO CO₂ overnight.



Day two Protocol

- 13 Warm MA media in waterbath to 37°C.

14 Place screens into a dish with MA media.

15 Place islets (min 70 islets per well), into tube with 500 μ l of MA media. Let islets gravity settle to the bottom of the tube. Pipette islets (100 μ l) and carefully dispense into center of the well of islet capture microplate. Place screen, flat side down, to well using forceps ensuring that the screens click into the well. Ensure there are no bubbles trapped under the screens. Add the remaining MA media to a total volume of 500 μ l in each well (by pipetting onto the side of the well). Incubate at 37°C with NO CO₂ for 1hr.

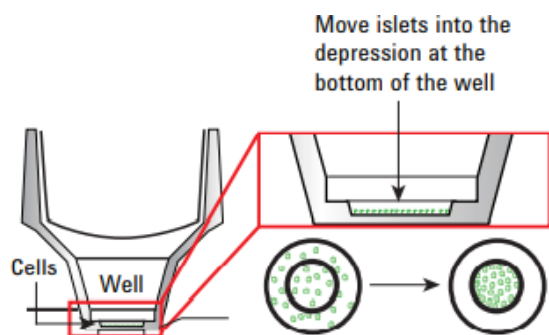


Figure 2. Islets should be in the depression at the bottom of the well.

- 16** Ensure to not place islets on the wells that are designated as blanks. See template below.

A	B	C	D	E	F
Blank	sample 1	sample 2	sample 3	sample 4	sample 5
sample 6	sample 7	sample 8	Blank	sample 9	sample 10
sample 11	sample 12	Blank	sample 13	sample 14	sample 15
sample 16	sample 17	sample 18	sample 19	sample 20	Blank

- 17** While islets are incubating, turn on seahorse Xfe Analyzer, turn on computer and open Wave. Open protocol and set up plate details.

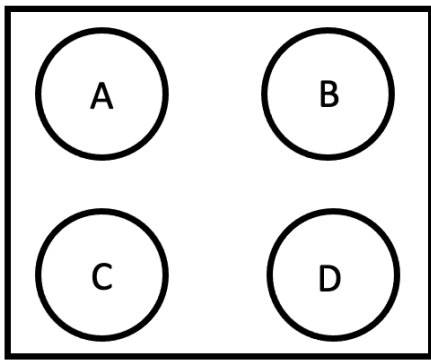
Protocol parameters:

A	B	C	D	E
	Cycles	Mix	Wait	Measure
Basal (2 or 2.8mM Glucose)	4	3:00	2:00	3:00
Glucose (16.7 or 20mM)	6	3:00	2:00	3:00
Oligomycin (5µM)	8	3:00	2:00	3:00
FCCP (3 or 5µM)	6	3:00	2:00	3:00
Rotenone/Antimycin A (5µM)	8	3:00	2:00	3:00

- 18** Make working stocks solutions.

- 19** Load solutions into the ports of the extracellular flux assay kit plate.

A - Glucose (50µl)
B- Oligomycin (55µl)
C- FCCP (60µl)
D- Rotenone/Antimycin A (65µl)



- 20 Press run on the seahorse program. Place extracellular flux assay plate (with booster plate and lid removed). Once initialization and 1hr incubation is complete remove the utility plate and replace with islet plate.
- 21 Once complete, remove plate, and collect for DNA or protein.
 - 21.1 Collect islets from well, including scraping off islets from the capture screens into a 1.5ml tube. Use extra MA media to wash wells
 - 21.2 Centrifuge tubes at 1000rpm for 4 minutes for protein or 1500rpm for 6 minutes for DNA.
 - 21.3 Aspirate supernatant. Store dry pellet at -20°C until ready to run protein or DNA assay.