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🌐 Cortical neuron differentiation using forced NGN2 expression

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ABSTRACT

This is a modified protocol based on (Zhang et al.2013 and Meijer et al., 2019) to generate cryopreserved NGN2 neurons, using hPSCs that carry a doxycycline-inducible NGN2 cassette.

The voluming are calculated for 10cm TC dishes and should be adapted accordingly



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

Created: Jan 29, 2024

Last Modified: Jan 29, 2024

MATERIALS

PROTOCOL integer ID: 94363

Media:

Keywords: ASAPCRN

KSR medium

	500ml	250ml	50ml	Reagent
	415ml	207.5ml	41.5ml	KO DEMEM
	75ml	37.5ml	7.5ml	KO SRM
	5ml	2.5ml	0.5ml	GlutaMax, NEAA, P/S
	0.5ml	0.25ml	0.05ml	β -mercapto ethanol

N2B medium

	500ml	250ml	50ml	Reagent
	480ml	240ml	48ml	DMEM F12 + HEPES
	5ml	2.5ml	0.5ml	GlutaMax, N2, P/S
	7.5ml	3.75ml	0.75ml	20% Dextrose

D2 N2B w/o B27 supplement

D3 N2B 1:100 B27 supplement

NBM medium

	500ml	250ml	50ml	Reagent
	485ml	242.5ml	48.5ml	Neurobasal
	5ml	2.5ml	0.5ml	GlutaMax, P/S
	7.5ml	3.75ml	0.75ml	20% Dextrose
	10ml	5ml	1ml	B27 supplement
	2.5ml	1.25ml	0.25ml	NEAA

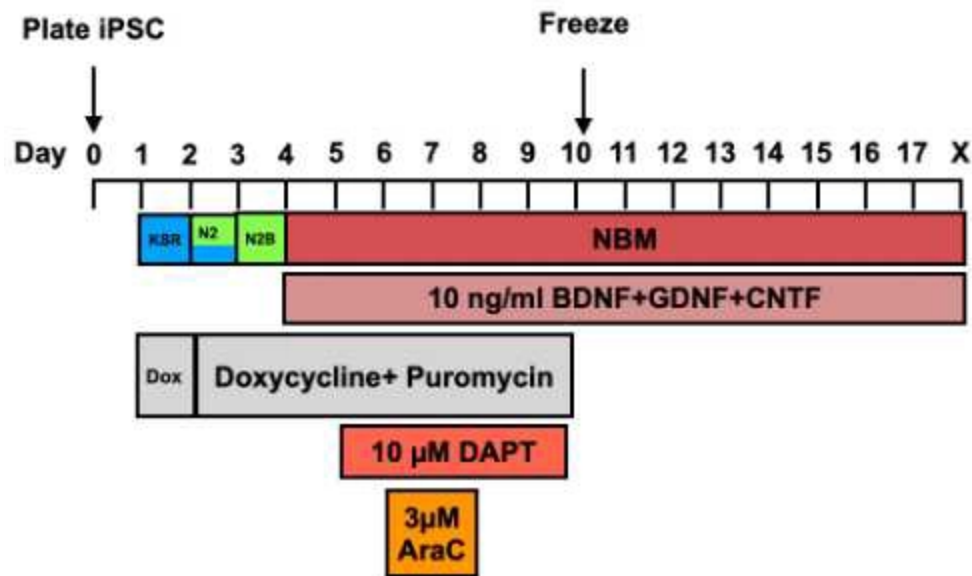
Consumables: For preparation of Growth factors check Cell culture recipes sheet

A	B	C	D	E	F
Reagent	Vendor	Cat no.	Stock con.	Working conc.	Dilution
Neurobasal medium	Thermofisher Scientific	#21103049	n.a.	n.a.	n.a.
Knock DMEM	Gibco	#10829-018	n.a.	n.a.	n.a.
KnockOut Serum Replacement	Invitrogen	10828-028	n.a.	n.a.	n.a.

A	B	C	D	E	F
MEM non-essential amino acids	Invitrogen	11140-050	n.a.	n.a.	n.a.
Beta-mercaptoethanol	Invitrogen	21985-023	n.a.	n.a.	n.a.
GlutaMax	Invitrogen	35050061	n.a.	n.a.	n.a.
DMEM F12 HEPES	Gibco	11330057	n.a.	n.a.	n.a.
N2 supplement	Invitrogen	17502048	n.a.	n.a.	n.a.
20% Dextrose			20%	n.a.	n.a.
Penicillin/Strep tomycin	Invitrogen	15140122	100x	1x	1:100
Trehalose	Sigma	#T9531-25G	n.a.	n.a.	n.a.
DMSO	Sigma	#D8418-250M	n.a.	n.a.	n.a.
mTeSR plus	Stem cell Technologies	# 100-0276	n.a.	n.a.	n.a.
Matrigel (Differentiation)	Corning	354234	n.a.	n.a.	n.a.
Dnase	NEB	#M030	n.a.	n.a.	n.a.
BDNF	Peprotech	#450-02	10µg/ml	10ng/ml	
GDNF	Peprotech	#450-10	10µg/ml	10ng/ml	
CNTF	Peprotech	# 450-13	10µg/ml	10ng/ml	
DAPT	Fisher Scientific	#26-341-0	10mM	10µM	
Doxycyclin Hyclate	Sigma	D9891-5g	2mg/ml	2µg/ml	
Puromycin	Life technology	A11138-03	10mg/ml		
Accutase	Gibco	#A11105			
Y27632 ROCK Inh.	Stem cell technologies	#72304	10mM	10µM	
Cell strainer (40µm)	Falcon/Corning	#352340	n.a.	n.a.	n.a.
B27 Supplement	Invitrogen	17504044	n.a.	n.a.	n.a.

Overview

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Differentiation overview

D0

- Seed 4×10^6 single cells onto 1x10 cm dish (coating MG-GFR or with GF 2mg) in iPSC medium supplemented with 10 μ M RI in 7ml medium

D1

- Add 8ml KSR with 2 μ g/ml DOX

D2

- 9ml 1:1 KSR/N2B w/o B27 2 μ g/ml DOX and 5 μ g/ml Puro



Ensure 24h window between this media change and the previous one

D3

- 5 10ml N2B containing 2µg/ml DOX) and 5µg/ml Puro

D4

- 6 10 ml NBM containing 2µg/ml DOX, 5µg/ml Puro, BDNF 10µg/ml , GDNF 10µg/ml , CNTF 10µg/ml

D5

- 7 10 ml NBM containing 2µg/ml DOX, 5µg/ml Puro, BDNF 10µg/ml , GDNF 10µg/ml , CNTF 10µg/ml and 10µM DAPT

D6 AraC treatment

- 8 10 ml NBM 2µg/ml DOX, 5µg/ml Puro, BDNF 10µg/ml, GDNF 10µg/ml, CNTF 10µg/ml, 10 µM DAPT
add 3 µM AraC varies between cell lines

! Discard AraC trash according to environmental health guidelines!

D7 AraC treatment

- 9 10 ml NBM 2µg/ml DOX, 5µg/ml Puro, BDNF 10µg/ml, GDNF 10µg/ml, CNTF 10µg/ml, 10 µM DAPT
add 3 µM AraC varies between cell lines

! Discard AraC trash and used medium according to environmental health guidelines!

D8

- 10 2x wash with 5ml NBM w/o GF to remove AraC completely
add 8-10 ml NBM 2µg/ml DOX, 5µg/ml Puro, BDNF 10µg/ml, GDNF 10µg/ml, CNTF 10µg/ml and 10 µM DAPT


! Discard AraC trash and used medium according to environmental health guidelines!



D9


- 11 **add 8-10 ml** NBM 2µg/ml DOX, 5µg/ml Puro, BDNF 10µg/ml, GDNF 10µg/ml, CNTF 10µg/ml and 10 µM DAP


D10 Cryopreservation

- 12 remove medium

- 12.1 1x wash with  4 mL | PBS

- 12.2 add  4 mL Accutase supplemented with 10µM RI for 90-120min at  37 °C

- 12.3 After incubation time; If needed add  200 µL DNase I

- 12.4 Add  3 mL NBM and dissociate using 10 ml serological pipette

- 12.5 Pool 3x10cm dishes and collect cells in 1x50 ml falcon after pipetting through a 40µm cell strainer on Ice

!collect all cells in a T175 TC fask on ice!

- 12.6 Rinse dishes with 3 ml NBM
- 12.7 Take a sample from the flask to determine cell number
!Always store cells suspension on ice from now on!
- 12.8 Aliquot pooled cells into 15 ml falcons and centrifuge at 300g for 5min at 4°C
- 12.9 Freeze cells in 1.5M or 3M aliquots and store at -80°C overnight and **transfer the next day** into the liquid nitrogen tank
- Freezing medium:
70% KOSRM
20% 1M Trehalose
10% DMSO