

JAN 29, 2024

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DOI:

dx.doi.org/10.17504/protocols.io. n2bvj3wwwlk5/v1

**Protocol Citation:** Beatrice Weykopf 2024. Cortical neuron differentiation using forced NGN2 expression. **protocols.io** https://dx.doi.org/10.17504/protoc ols.io.n2bvj3wwwlk5/v1

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

Created: Jan 29, 2024

## 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 doxycline-inducible NGN2 cassette.

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

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	ß-mercapto ethanol

#### N2B medium

500ml	250ml	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

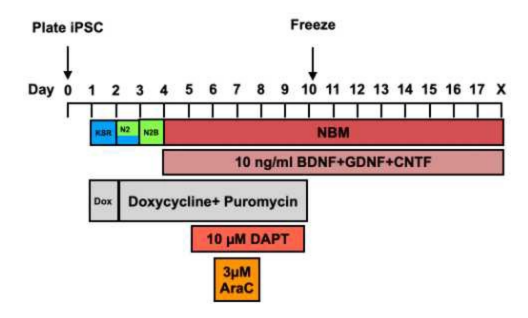
## Consumables: For preparation of Growth factors check Cell culture recipes sheet

A	В	С	D	E	F
Reagent	Vendor	Cat no.	Stock con.	Working conc.	Dilution
Neurobasal medium	Thermofisher Scientific	#2110304 9	n.a.	n.a.	n.a.
Knock DMEM	Gibco	#10829- 018	n.a.	n.a.	n.a.
KnockOut Seru m Replacement	Invitrogen	10828- 028	n.a.	n.a.	n.a.

A	В	С	D	E	F
MEM non- essential amino acids	Invitrogen	11140- 050	n.a.	n.a.	n.a.
Beta- mercaptoethan ol	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/m I		
Accutase	Gibco	#A11105			
Y27632 ROCK Inh.	Stem cell technologies	#72304	10mM	10μΜ	
Cell strainer (40µm)	Falcon/Cornin g	#352340	n.a.	n.a.	n.a.
B27 Supplement	Invitrogen	17504044	n.a.	n.a.	n.a.

# Overview

1



Differentiation overview

D0

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

**D1** 

3 Add 8ml KSR with 2μg/ml DOX

**D2** 

4 9ml 1:1 KSR/N2B w/o B27 2μg/ml DOX and 5μg/ml Puro

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

<b>&amp;</b>	protocols.io					
		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				
١		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				
	a	10 ml NBM 2ug/ml DOX 5ug/ml Puro RDNF 10ug/ml GDNF 10ug/ml CNTF 10ug/ml 10 uM DAPT				

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 I PBS
  - 12.2 add Δ 4 mL Accutase supplemented with 10μM RI for 90-120min at \$\mathbb{8}\$ 37 °C
  - 12.3 After incubation time; If needed add Δ 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 falsk on ice!

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:

Freezing medium: 70% KOSRM 20% 1M Trehalose 10% DMSO