Modulation of GABA_A activity: Investigations in hiPSC-derived neuronal co-cultures and human ion channel assays



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INTRODUCTION

A balance between inhibitory neurotransmission and neuronal excitation is critical for normal brain function. γ -aminobutyric acid (GABA) is the most abundant inhibitory neurotransmitter, which acts on GABA_A receptors. Perturbation of GABA_A signalling by drug-induced inhibition and potentiation are common mechanisms producing seizure and sedation, respectively. The introduction of commercially available human induced pluripotent stem cell (hiPSC-) derived neurons facilitates the *in vitro* study of neuronal function and, in our work, the detection of seizure liability during drug discovery. It is known that GABA_A antagonists such as picrotoxin increase neuronal firing and induce a seizure-like phenotype in hiPSC-derived neurons, however further characterisation of GABA_A modulation within these cell models is lacking. This study aimed to address this by assessing the effects of a selection of GABA modulators on the electrical activity of hiPSC-derived neuronal co-cultures, and the ion flux of $\alpha_1\beta_2\gamma_2$ -GABA_A.

METHODS

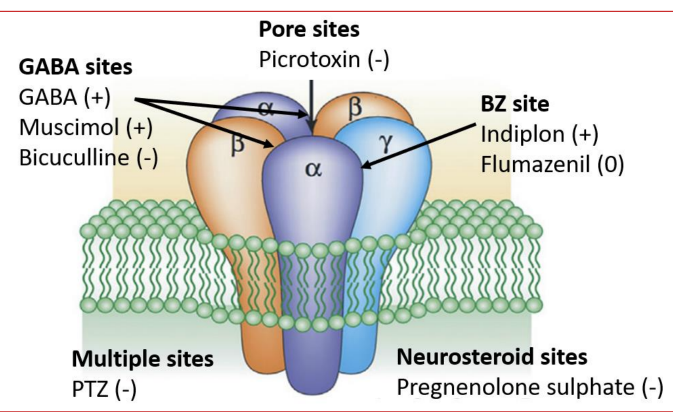
hiPSC-DERIVED NEURONAL CO-CULTURES

- iCell Glutaneurons (80% glutamatergic/20% GABAergic neurons) were plated with astrocytes (85%:15%) and monitored using a microelectrode array (MEA) system (Maestro Edge, Axion).
- On DIV22 and DIV23, spontaneous electrical activity was recorded at baseline and 1 hour after exposure to GABA_A modulators and solvent controls.
- Cells exposed to agonists were subsequently challenged with antagonists and spontaneous electrical activity was measured 15 minutes after application.

HUMAN $\alpha_1\beta_2\gamma_2$ -GABAA ION CHANNEL ASSAYS

- The activity of GABA modulators was assessed by automated patch-clamp (QPatch II, Sophion) using a CHO $\alpha_1\beta_2\gamma_2$ -GABA_A cell line.
- All modulators except for ligands were coapplied with 30μM GABA.
- 6-point dose-response curves were generated for all modulators.
- For agonists, a 5-point dose-response curve plus subsequent antagonist challenge was generated.

COMPOUND SELECTION



RESULTS

MEA PARAMETERS

Firing rate - Weighted mean firing rate based on electrodes with activity greater than minimum spike rate, set by the neural statistics calculator.

Burst duration - Average time between the first and last spike in a burst.

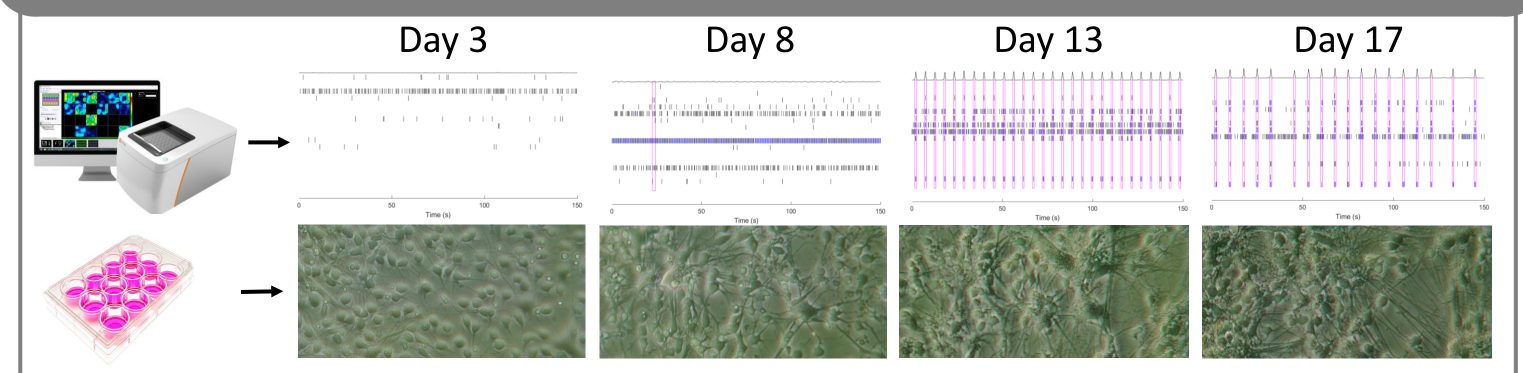
Network burst freq. - Total number of electrode bursts divided by recording time. **Network burst duration** - Average time between the first and last spike in a network burst.

No. spikes per network burst - Average number of spikes in a network burst.

↑↑↑≥100%
个个50 to 99%
↑20 to 50%
\leftrightarrow within +/-20%
↓-20 to -50%
$\downarrow \downarrow$ -50 to -99%
↓ ↓ ↓ ≥-100 %
***p<0.001
**p<0.01

* p<0.05

DEVELOPMENT OF SPONTANEOUS ELECTRICAL ACTIVITY



AGONISTS INCREASE GABA_A RESPONSE, DECREASE POPULATION ACTIVITY

	JI J INCKLASE GAI	DA _A RESPONSE, DE	CALASE POPULATA	JNACIIVIII		
	GABA Agonist, GABA site	MUSCIMOL Agonist, GABA site	GABA & MUSCIMOL	INDIPLON Positive allosteric modulator, BZ site		
OVERVIEW OF ACTIVITY						
Mean firing rate	$\downarrow \downarrow \downarrow$	$\downarrow\downarrow\downarrow^*$	$\downarrow \downarrow$	<u> </u>		
Burst duration	\leftrightarrow	$\downarrow \downarrow *$	$\downarrow \downarrow *$	\downarrow		
Network burst freq.	$\downarrow \downarrow$	$\downarrow\downarrow\downarrow\downarrow^*$	$\downarrow\downarrow\downarrow\downarrow$	\downarrow		
Network burst duration	$\downarrow \downarrow$	$\downarrow\downarrow\downarrow\downarrow^*$	$\downarrow\downarrow\downarrow\downarrow$	\leftrightarrow		
No. spikes per network burst	$\downarrow \downarrow$	$\downarrow\downarrow\downarrow\downarrow^*$	$\downarrow\downarrow\downarrow\downarrow$	↓		
RASTER PLOTS						
Baseline 1 hour after addition	1ΟμΜ 1 το μπο	3 μΜ 0 50 100 150 Time (s)	1μΜ each 1 μ	3 n M Time (s) Time (s) Time (s) Time (s)		
$HUMAN$ $α_1β_2γ_2$ - $GABA_A$ ION CHANNEL ASSAYS						
6-point concentration- dose response	Weshouse (%) 100 100 GABA conc. (pM)	Sesponse (%) 1 10 100 Muscimol conc. (pM)	100 Sesponse 50 1 10 100 GABA and Muscimol conc. (MM)	Sesponse (%) 100-100 100 Indiplon conc. (nM)		

 $EC_{50} = 2.8 \mu M$

 $EC_{50} = 1.1 \mu M$

 $EC_{50} = 4.3 \text{nM}$

 $EC_{50} = 11.1 \mu M$

ANTAGONISTS DECREASE GABA RESPONSE, MIXED POPULATION ACTIVITY **PREGNENOLONE BICUCULLINE** PTZ **PICROTOXIN SULFATE** Competitive Non-competitive Non-competitive Negative allosteric antagonist, multiple antagonist, GABA antagonist, pore modulator, modes of action sites site neurosteroid sites **OVERVIEW OF ACTIVITY** Mean firing rate \leftrightarrow **Burst duration** \leftrightarrow Network burst freq. Network burst 个个 \leftrightarrow \leftrightarrow duration No. spikes per $\uparrow \uparrow \uparrow \uparrow$ \leftrightarrow network burst RASTER PLOTS 10μM $3\mu M$ 300μΜ 1μ M Baseline 1 hour after addition HUMAN $\alpha_1\beta_2\gamma_2$ -GABA_A ION CHANNEL ASSAYS 6-point concentrationdose response PTZ conc. (mM) Pregnenolone Sulfate conc. (M)

3 REVERSAL OF AGONIST-INDUCED SEDATION BY ANTAGONISTS

 $IC_{50} = 1.4 \mu M$

 $IC_{50} = 3.1 \text{mM}$

 $IC_{50} = 28.8 \mu M$

 $IC_{50} = 5.5 \mu M$

	INDIPLON	GABA	MUSCIMOL	MUSCIMOL			
	FLUMAZENIL silent antagonist, BZ site	BICUCULLINE	BICUCULLINE	PICROTOXIN			
OVERVIEW OF ACTIVITY							
Mean firing rate	$\uparrow \uparrow$	^^^*	\leftrightarrow	^^^*			
Burst duration	\leftrightarrow	$\uparrow \uparrow$	\uparrow	^^^*			
Network burst freq.	\leftrightarrow	个个个	个 (from 0)	个** (from 0)			
Network burst duration	\downarrow	个个个	个 (from 0)	个** (from 0)			
No. spikes per network burst	\leftrightarrow	个个	个个个	^^^*			
RASTER PLOTS							
	3nM	10μΜ	3μΜ	3μΜ			
1 hour after agonist addition							
	3nM	3μM	3μM	10μM			
15 minutes after			σμινι				
antagonist addition							
$HIINANN \sim R \times _CARA IONICHANNIELACCAVC$							
HUMAN α ₁ β ₂ γ ₂ -GABA _A ION CHANNEL ASSAYS							
Antagonist		%) esuoc 50-	100 3μM) sundsea	Picrotoxin (%) esuodsey			
challenge	10 30 100 300 300 Indiplon (nM)	1 3 10 30 30 GABA concentration (pM)	0.3 1 3 10 30 30 Muscimol concentration (M)	0.3 1 3 10 30 30 Muscimol concentration (M)			

DISCUSSION AND CONCLUSIONS

- Agonists GABA and muscimol induced sedation in hiPSC-derived neuronal co-cultures and increased $\alpha_1\beta_2\gamma_2$ -GABA_A current (fig.1) while antagonists bicuculline and picrotoxin induced seizure in hiPSC-neuronal co-cultures and reduced $\alpha_1\beta_2\gamma_2$ -GABA_A current (fig.2).
- PTZ is used *in vivo* to induce seizure. This often involves chronic repeat-dose application, suggesting PTZ may not translate well to single-dose *in vitro* studies (fig.2).
- Pregnenolone sulfate (PS) did not induce seizure in hiPSC-derived neuronal co-cultures, yet inhibited $\alpha_1\beta_2\gamma_2$ -GABA_A current. This suggests the expression of other subtypes in neuronal cells, possibly GABA_C which is considerably less sensitive to PS than GABA_A.
- In ion channel assays, bicuculline blocked GABA- and muscimol-induced current (fig.3).
 In hiPSC-derived neuronal co-cultures however, muscimol-induced sedation was not reversed by bicuculline. It is known that bicuculline cannot compete with muscimol at GABA_C, further suggesting its expression in hiPSC-derived neuronal co-cultures.
- Indiplon, a marketed sleeping aid, induced sedation in hiPSC-derived neuronal cocultures and increased $\alpha_1\beta_2\gamma_2$ -GABA_A current (fig.3). It was competitively antagonized by flumazenil, a clinical antidote to indiplon overdose. As a silent antagonist, Flumazenil was inactive alone in both assays.
- These studies have further characterised modulation of GABA_A activity within hiPSC-derived neuronal co-cultures by recapitulating expected clinical outcomes. This further validates the model as a translationally relevant screen for seizure detection which also shows promise for sedation.