Inhibitory Synapse Formation in a Co-culture Model Incorporating GABAergic Medium Spiny Neurons and HEK293 Cells Stably Expressing GABA$_A$ Receptors

Laura E. Brown$^1$, Celine Fuchs$^1$, Martin W. Nicholson$^1$, F. Anne Stephenson$^1$, Alex M. Thomson$^1$, Jasmina N. Jovanovic$^1$

$^1$UCL School of Pharmacy, University College London

Correspondence to: Jasmina N. Jovanovic at j.jovanovic@ucl.ac.uk

URL: http://www.jove.com/video/52115
DOI: doi:10.3791/52115

Keywords: Neuroscience, Issue 93, Developmental neuroscience, synaptogenesis, synaptic inhibition, co-culture, stable cell lines, GABAergic, medium spiny neurons, HEK 293 cell line

Date Published: 11/14/2014

Citation: Brown, L.E., Fuchs, C., Nicholson, M.W., Stephenson, F.A., Thomson, A.M., Jovanovic, J.N. Inhibitory Synapse Formation in a Co-culture Model Incorporating GABAergic Medium Spiny Neurons and HEK293 Cells Stably Expressing GABA$_A$ Receptors. J. Vis. Exp. (93), e52115, doi:10.3791/52115 (2014).

Abstract

Inhibitory neurons act in the central nervous system to regulate the dynamics and spatio-temporal co-ordination of neuronal networks. GABA (γ-aminobutyric acid) is the predominant inhibitory neurotransmitter in the brain. It is released from the presynaptic terminals of inhibitory neurons within highly specialized intercellular junctions known as synapses, where it binds to GABA$_A$ receptors (GABA$_A$Rs) present at the plasma membrane of the synapse-receiving, postsynaptic neurons. Activation of these GABA-gated ion channels leads to influx of chloride resulting in postsynaptic potential changes that decrease the probability that these neurons will generate action potentials.

During development, diverse types of inhibitory neurons with distinct morphological, electrophysiological and neurochemical characteristics have the ability to recognize their target neurons and form synapses which incorporate specific GABA$_A$Rs subtypes. This principle of selective innervation of neuronal targets raises the question as to how the appropriate synaptic partners identify each other.

To elucidate the underlying molecular mechanisms, a novel in vitro co-culture model system was established, in which medium spiny GABAergic neurons, a highly homogenous population of neurons isolated from the embryonic striatum, were cultured with stably transfected HEK293 cell lines that express different GABA$_A$R subtypes. Synapses form rapidly, efficiently and selectively in this system, and are easily accessible for quantification. Our results indicate that various GABA$_A$R subtypes differ in their ability to promote synapse formation, suggesting that this reduced in vitro model system can be used to reproduce, at least in part, the in vivo conditions required for the recognition of the appropriate synaptic partners and formation of specific synapses. Here the protocols for culturing the medium spiny neurons and generating HEK293 cells lines expressing GABA$_A$Rs are first described, followed by detailed instructions on how to combine these two cell types in co-culture and analyze the formation of synaptic contacts.

Video Link

The video component of this article can be found at http://www.jove.com/video/52115/

Introduction

GABA is one of the earliest neurotransmitters found in the embryonic brain, preceding the most abundant excitatory neurotransmitter glutamate. During development, GABA depolarizes and excites immature neurons, playing a key role in regulating cell proliferation, migration and formation of neuronal networks without inducing excitotoxicity. In the adult brain, the reversal potential for GABA$_A$ receptor channels is shifted to more negative potentials due to a decrease in the intracellular concentration of chloride. This shift is caused by up-regulation of the potassium-chloride co-transporter (KCC2), which transports chloride out of the cell, and, in parallel, down-regulation of the sodium-potassium-chloride transporter (NKCC1), which has the opposite effect.

In the brain, GABA primarily binds to either GABA$_A$ or GABA$_B$ receptors to mediate fast or slow synaptic inhibition, respectively. GABA$_A$Rs are a class of receptors also known as heteropentameric ionotropic or ligand-gated Cys-loop ion channels. Two molecules of GABA are required for activation of the receptor, which is permeable to chloride ions and to a lesser degree, bicarbonate ions. The increase in chloride conductance decreases the effectiveness of depolarizing, excitatory events in activating the postsynaptic neuron.

Structural diversity of GABA$_A$Rs has long been recognized as a key factor in determining their wide range of functional and pharmacological properties. Native GABA$_A$Rs are hetero-pentamers composed of subunits with multiple isoforms classified as: α(1-6), β(1-3), γ(1-3), δ, ε, π and θ, with a common transmembrane topology comprising a large N-terminal extracellular domain, four transmembrane domains (TMs), and a major intracellular domain between TMs 3 and 4. The β3 and γ2 subunits are essential for synaptic inhibition and organism survival, because mice bearing genetic deletion of these subunits die after birth. In contrast, individual isoforms of α subunit are important for the function of specific synaptic connections in the brain associated with different behaviors such as anxiety, sedation, arousal, and others, but are...
not, individually, essential for life.\textsuperscript{7,9} GABA\textsubscript{A}Rs are the main sites of action for a variety of drugs with potent sedative, hypnotic, anxiolytic and anticonvulsant effects, such as benzodiazepines, barbiturates, neurosteroids and anesthetics.\textsuperscript{7,10,11}

Synaptic GABA\textsubscript{A}Rs typically contain a γ2 subunit, two β subunits (most commonly β2 or β3) and two α subunits (α1, α2, α3 or α5).\textsuperscript{12,13} The predominant class of extra-synaptic receptors contains the δ subunit in combination with two α subunits (α4 or α6), and two β subunits (β2 or β3).\textsuperscript{14} Subcellular localization of GABA\textsubscript{A}Rs to axons, dendrites or soma, and insertion into the plasma membrane are dependent on the presence of β-subunits.\textsuperscript{15,16} However, selective incorporation of different GABA\textsubscript{A}R subtypes into distinct types of synapses correlates well with the presence of specific α subunits (α1, α2, α3 or α5).\textsuperscript{17,18} Importantly, deletion of α1 or α2 subunit in mice causes ultrastructural changes at inhibitory synapses.\textsuperscript{19} This suggests that GABA\textsubscript{A}Rs themselves may play a direct role in regulating synapse formation.

Evidence indicates that GABAergic synapse development is a precisely co-ordinated sequence of events, in which both the neuronal targets contacted by different types of inhibitory axons and the receptors that are clustered at each class of inhibitory synapse are selective and functionally attuned.\textsuperscript{17,20,22} This fundamental principle of specificity at GABAergic synapses raises the question as to how the pre- and postsynaptic partners recognize each other during the initiation of synaptic contacts.

In vitro co-culture assays have been applied successfully to study some of the mechanisms of synapse formation and to test the role of individual synaptic cleft-spanning proteins in this process. One of the common trans-synaptic interacting protein combinations that function bi-directionally to mediate synapse formation and maturation, are the Neurexins (Nrxns) and Neuroligins (NLs). Nrxns are presynaptic proteins that exhibit alternative splicing within their laminin-neurexin-sex hormone-binding protein domains, giving rise to many different isoforms.\textsuperscript{23} While the Nrxns also interact with other proteins, NLs are thought to be their ubiquitous postsynaptic partners.\textsuperscript{24} Together these proteins contribute to holding the presynaptic and postsynaptic membranes in close and rigid apposition.\textsuperscript{25} The two most abundant isoforms are NL-1 and NL-2 which are present at excitatory and inhibitory synapses, respectively.\textsuperscript{26} One of the earliest culture model systems, designed to investigate trans-synaptic protein interactions, employed different types of non-neuronal cells, most commonly immortal cell lines such as Human Embryonic Kidney (HEK) 293 cells, to over-express NL-2. When these cells were cultured with pontine neurons, an accumulation of presynaptic proteins in close proximity to the surface of the HEK cells was observed, indicating formation of synapse-like contacts. Addition of soluble β-neurexin to these co-cultures inhibited the formation of contacts, suggesting that trans-synaptic interactions between Nrxns and NLs are necessary for synaptic contact formation.\textsuperscript{27} Moreover, transient expression of β-neurexin in COS (CV-1 simian in Origin, and carrying the SV40 genetic material) cells co-cultured with dissociated hippocampal glutamatergic and GABAergic neurons induced expression of the postsynaptic protein gephyrin and of GABA\textsubscript{A}R subunits γ2 and δ2 at points of contact between these two cell types.\textsuperscript{28} Another example of a co-culture model used to study synapse formation involved HEK293 cells, transiently transfected with GABA\textsubscript{A}R subunits α2/β3/γ2 and NL-2, and a mixed population of hypothalamic neurons.\textsuperscript{29} This study concluded that the expression of NL-2 is an absolute requirement for formation of inhibitory synapses.

However, in the recent co-culture study, stably transfected α1/β2/γ2 GABA\textsubscript{A}Rs in HEK293 cells were found to be sufficient to induce functional synapses when co-cultured with GABAergic medium spiny neurons, without the need for additional trans-synaptic or postsynaptic adhesion proteins. However, a prominent increase in synapse formation and strength was observed when NL-2 was co-expressed with GABA\textsubscript{A}Rs.\textsuperscript{30} This indicates that this co-culture model system has advantages over previously described model systems, most evidently an increased sensitivity and reliability of synaptic contact detection. Two important factors contributing to the overall improvement in detection of synaptic contacts are: i) The use of stably transfected HEK293 cell lines with high and consistent expression of GABA\textsubscript{A}R subunits at the surface of individual cells. This consistency facilitates quantitative comparisons between different co-culture conditions. ii) The use of a pure population of GABAergic medium spiny neurons cultured from the embryonic striatum removes complications and ambiguities resulting from the use of mixed neuronal populations and allows, for example, selection of the most appropriate postsynaptic GABA\textsubscript{A}R types that can be compared with each other during synapse formation.

Formation of synapses is thought to involve many trans-synaptic signals within pre- and postsynaptic cell adhesion complexes. Due to the bi-directional nature of synaptic signaling and the sheer numbers of cell adhesion molecules, it is difficult to identify key components involved in synapse formation. Thus, transfecting a single cell adhesion protein into a non-neuronal cell (in this case, the two most prevalent postsynaptic targets for GABAergic medium spiny neurons in vivo, α1/β2/γ2 or α1/β3/γ2 GABA\textsubscript{A}Rs) greatly reduces the complexity of trans-synaptic signals available at the postsynaptic surface and allows precise quantitative analysis of the efficacy of this protein in promoting synapse formation.

**Protocol**

Sprague-Dawley rats or BAB/c inbred mice (Harlan, UK; the number of pregnant females used was 30) were housed and sacrificed according to UK Home Office [and European Communities Council directive of 24 November 1986 (86/609/EEC)] guidelines. The project was formally approved by the UCL School of Pharmacy Ethics Committee.

1. **Preparation of Instruments, Culture Medium, and Dishes**

   1. Turn on and clean the laminar flow hood with 70% ethanol in order to work under sterile conditions at all times.
   2. Prepare HEK293 cell culture medium, containing Dulbecco’s Modified Eagle Medium pH 7.4 (DMEM, 500 ml), L-glutamine (2 mM), penicillin (50 Units/ml), streptomycin (50 μg/ml), and fetal bovine serum (10%).
      
      **NOTE:** penicillin and streptomycin are irritants.
   3. Prepare the serum-free neuronal culture medium, containing Neurobasal medium pH 7.4 (500 ml), B27 supplement (25 ml), L-glutamine (2 mM), penicillin (50 Units/ml), streptomycin (50 μg/ml), and glucose (6 mM).
   4. Prepare 500 ml of HEPES-buffered saline solution (HBSS), containing HBSS 10x stock (50 ml), HEPES (1 M) (5 ml) and water (445 ml), pH 7.4.
   5. Autoclave phosphate buffered saline solution (PBS, pH 7.4; 1 L), water (1 L), glass coverslips (13 mm in diameter) and glass Pasteur pipettes to sterilize them.
2. Preparation of HEK293 Stable Cell Line Expressing α1/β3/γ2-GABA<sub>R</sub>

1. Plate 2x10<sup>5</sup> HEK293 cells into a 10 cm sterile tissue culture plate and incubate at 37 °C in a humidified 5% carbon dioxide (CO<sub>2</sub>) atmosphere (CO<sub>2</sub> incubator) in order to reach 70-90% confluency overnight.

2. The following day, transfet Hek293 cells with GABA<sub>R</sub> α1 subunit cDNA in the pcDNA 3.1(+) expression vector incorporating the G418 disulfate (Table 1) resistance gene, and the GABA<sub>R</sub> β3 subunit cDNA in the expression vector incorporating the phleomycin D1 (Table 1) resistance gene (both under the regulation of a human cytomegalovirus immediate-early (CMV) promoter), using a cationic liposome formulation, which complexes with negatively charged nucleic acid molecules (Table 1) according to manufacturer’s protocol.

3. Briefly, add 500 μl of reduced serum medium (pH 7.4; Table 1) and 7.5 μg of each cDNA construct to a sterile 15 ml centrifuge tube, followed by 15 μl of liposomal transfection buffering reagent, and gently mix before leaving at room temperature for 5 min.

4. To this mixture, add 8.75 μl of liposomal transfection reagent and gently mix before leaving at room temperature for 30 min. Following this, add 3 ml of HEK293 cell culture medium (without antibiotics), pipette the contents of the centrifuge tube up and down twice, transfer drop-wise onto the cells growing in a 10 cm tissue culture dish, and incubate for 48 hr at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator).

5. Wash the HEK293 cells gently with sterile PBS, pH 7.4, and dilute the transfected HEK293 cells into new 10 cm tissue culture dishes, in the following ratios: 1:3, 1:5, 1:7, 1:10, 1:15 and 1:20. This ensures that the cells will not become overly confluent.

6. Start selection of the HEK293 cells expressing both GABA<sub>R</sub> α1 and β3 subunits by adding 800 μg/ml of each antibiotic selection marker, G418, and Phleomycin D1, to the culture medium. Incubate the cells at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator) and replace the antibiotic-containing medium (10 ml) every 2 days.

7. When small white colonies start to form (usually after about 7 days), carefully select a single colony from each of the dishes and collect it using a sterile P1000 pipette tip. Transfer it to one well of a 24-well tissue culture plate containing 500 μl of medium and carefully resuspend by pipetting the medium up and down. Ensure that exactly one colony is transferred into each well (a total of 5-20 colonies is advised).

8. Incubate the cells at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator) and replace the antibiotic-containing medium every 2 days.

9. Once the colonies become 70-80% confluent, gently pipette the medium up and down to dislodge the cells from the bottom of the 24-well tissue culture plate. Transfer and divide the suspension of cells between 2 wells in a 6-well tissue culture plate. Incubate the cells at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator) and replace the antibiotic-containing medium every 2 days.

10. Dislodge and transfer only positive clones from the remaining wells to larger 6 cm tissue culture dishes. Incubate the cells at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator) and replace the antibiotic-containing medium every 2 days.

11. Gradually expand the colonies of cells under the antibiotic selection by transferring them to 10 cm tissue culture dishes and finally to tissue culture flasks (T-75 flasks). Incubate the cells at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator) and replace the antibiotic-containing medium every 2 days.

12. Plate 70,000 cells from each colony on glass coverslips (13 mm in diameter) and fix the cells to analyze the cell surface expression and co-localization of GABA<sub>R</sub> subunits by immunofluorescence.

13. Select the positive clone of HEK293 cells expressing high levels of both α1 and β3 subunits of GABA<sub>R</sub> receptors, plate 2 x 10<sup>5</sup> cells into a 10 cm sterile tissue culture dish and incubate at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator) overnight. Ensure that the cells are incubated in antibiotic-containing (G418 and Phleomycin D1) medium at all times.

14. The following day, transfet HEK293 cells with GABA<sub>R</sub> γ2s subunit cDNA in the pcDNA™ 3.1(+) expression vector incorporating Hygromycin B resistance gene using a non-liposomal lipid transfection reagent (Table 1).

NOTE: This transfection method allows better survival and higher expression efficiency of foreign proteins in slow growing stable cell lines which are under continuous selection with antibiotics.

15. In a sterile 15 ml centrifuge tube, add 250 μl of Enhancer and DNA condensation buffer (Table 1) and 1.4 μg of γ2s GABA<sub>A</sub> receptor subunit cDNA. Add 11.2 μl of Enhancer and vortex for 1 sec before leaving at room temperature for 5 min.

16. Add 35 μl of non-liposomal lipid transfection reagent and vortex for 10 sec before leaving at room temperature for 10 min. Add 3 ml of G418/Phleomycin D1-containing medium and pipette the content of the centrifuge tube up and down twice before transferring it drop-wise onto the cells growing in a 10 cm tissue culture plate. Incubate the cells for 48 hr at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator).

17. Wash the cells gently with sterile PBS and dilute them into new 10 cm tissue culture dishes, in the following ratios: 1:3, 1:5, 1:7, 1:10, 1:15 and 1:20.

18. Start selection of α1/β3-HEK293 cells expressing the γ2s subunit by adding 800 ng/ml of antibiotic selection marker Hygromycin B to the G418/Phleomycin D1-containing medium. Replace the old medium with the fresh G418/Phleomycin D1/Hygromycin B-containing medium (10 ml) every 2 days.

19. Repeat steps 2.7-2.12, under continuous selection in G418/Phleomycin D1/Hygromycin B-containing cell culture medium.

20. Store the positive clones at -140 °C in antibiotic-free cell culture medium and 10% dimethyl sulfoxide (DMSO) for the future use.

21. Test the level of expression of α1, β3 and γ2s GABA<sub>R</sub> subunits by immunoblotting and immunofluorescence in each clone following defrosting, because the expression can change due to reduced survival of cells under antibiotic selection.

3. Maintenance of HEK293 Cell Lines

1. Defrost a vial of control HEK293 cells or those expressing either α1/β3/γ2-GABA<sub>R</sub> (Table 1) or α1/β3/γ2-GABA<sub>R</sub> (described above) into 10 ml of cell culture medium in a 15 ml sterile centrifuge tube. Centrifuge at 440 x g for 5 min to remove excess DMSO.

2. Remove supernatant and resuspend cells in 1 ml of fresh cell culture medium.

3. First add 9 ml of fresh cell culture medium to a 10 cm tissue culture dish coated with poly-D-lysine (0.1 mg/ml) and then 1 ml of resuspended cells. Agitate gently, side-to-side, to disperse the cells and incubate at 37 °C in a humidified 5% CO<sub>2</sub> atmosphere (CO<sub>2</sub> incubator).
4. The following day, aspirate the medium to remove any cell debris and replace with 10 ml of fresh HEK293 cell culture medium.

5. Select stable cell lines with antibiotics to remove any cells which do not express GABA\(_A\)R subunits and therefore also lack the expression of antibiotic resistance markers. For the \(\alpha_1/\beta_3/\gamma_2\) stable cell line, replace normal medium with fresh cell culture medium containing G418 (800 \(\mu\)g/ml), Phleomycin D1 (800 \(\mu\)g/ml) and Hygromycin B (800 \(\mu\)g/ml). **CAUTION!** G418 is an irritant and Hygromycin B is corrosive, toxic and an irritant.

6. Passage the cells into a new tissue culture dish by seeding at a lower density once they achieve >70% confluence. Aspirate 10 ml of cell culture medium and wash twice briefly with PBS, pH 7.4. Add 1 ml of trypsin-EDTA solution, a solution of the protease trypsin (0.05% trypsin) and a \(\text{Ca}^{2+}\) chelator EDTA (0.02%) in PBS, pH 7.4, to detach the cells from the dish. **CAUTION!** Trypsin-EDTA solution is an irritant.

7. Add 10 ml of cell culture medium containing the correct antibiotics, to the dish and aspirate the cells. Centrifuge the cells at 440 \(x\) g for 5 min and resuspend them in 5 ml of the cell culture medium.

8. Passage the cells using a 1:10 dilution into a new tissue culture flask (T-75 flask) containing fresh cell culture medium and the correct antibiotics. Incubate the cells at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator) and replace the medium every two days. Passage the cells when >70% confluent (see step 2.8).

4. Preparation of GABAergic Medium Spiny Neuron Culture

1. Under sterile conditions prepare a 24-well plate with poly-L-lysine (0.1 mg/ml)-coated coverslips (13 mm in diameter) and incubate at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator).

2. The following day, aspirate with a pipette the excess poly-L-lysine and wash coverslips with two brief 10 sec and two 5 min long washes with sterile water. Add laminin (0.01 mg/ml) overnight and incubate at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator).

3. SANITIZE the dissection area with 70% ethanol and gather an array of dissecting tools such as curved and straight tissue forceps, scissors and tweezers, and place into 70% ethanol to fully sterilize the dissecting instruments.

4. Place the pregnant rat/mouse euthanized with CO\(_2\) on its back and clean the skin on its abdomen with 70% ethanol. Pinch the skin with tweezers and cut around the abdomen through the skin, muscle and peritoneum, to reveal the internal organs and uterus with embryos clearly. Extract the embryos (E16-17) from the uterus and place them in a Petri dish with chilled PBS.

5. Place the embryos in the laminar flow hood and decapitate them, collecting the heads in a new Petri dish with chilled HBSS.

6. UNDER A DISSECTING MICROSCOPE, disperse the heads using the curved and straight forceps. Place the brains into a new Petri dish containing chilled HBSS.

7. Separate the two cerebral hemispheres and carefully remove the meninges. Cut along the line of the hippocampus and peel back the cortex to reveal the striatum. Observe the striatum as a striated white structure at the anterior of the hemisphere.

8. DISSECT the striatum and cut it into very small pieces (1 - 2 mm in diameter) and use a fire-polished Pasteur pipette to collect the material into a sterile 15 ml centrifuge tube with a total volume of 1 ml. Fire-polishing ensures the material is collected without being damaged.

9. USING THE FIRE-POLISHED TIP of the Pasteur pipette, aspirate the cells and release them 8 - 10 times. Taking a new pipette with a fire-polish tip of approximately 30% of its original diameter (1 mm), triturate the solution a further 4 - 6 times until it appears homogeneous.

10. Filter the cells using a 100 \(\mu\)m nylon cell strainer into a fresh sterile centrifuge tube.

11. COUNT the cells with a hemocytometer and plate 70,000 cells into 500 \(\mu\)l of neuronal culture medium per well in a 24-well tissue culture plate. Agitate the wells left to right and incubate at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator) for 14 days in vitro (DIV).

12. After 7 days, check the purity of neuronal cultures and, if glia cells are present, add cytosine \(\beta\)-D-arabinoside (Ara-C; 5 \(\mu\)M) to the wells to stop their proliferation. To do this, remove 250 \(\mu\)l of neuronal culture medium (pH 7.4) from each well and add 250 \(\mu\)l of fresh medium containing Ara-C. **CAUTION!** Ara-C is an irritant.

5. Co-culture Preparation

1. ON DAY 11 of neuronal culture, coat a 6-well plate with poly-D-lysine (0.1 mg/ml) under sterile conditions and incubate overnight at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator).

2. The following day (day 12 of neuronal culture), aspirate the excess poly-D-lysine and wash wells 2x briefly and 2x for 5 min with sterile water before adding fresh cell culture medium (without antibiotics) to coat the wells with a small amount of serum from the medium.

3. ASPIRATE the culture medium from the tissue culture flask (T-75). Rinse the cells twice with PBS before adding 1 ml of Ca\(^{2+}\)/Mg\(^{2+}\)-chelating agent EDTA (0.48 mM) which gently and non-enzymatically dissociates cells. Agitate the cells to detach them from the bottom of the tissue flask.

4. ADD 10 ml of cell culture medium (without antibiotics as this may interfere with transfection), aspirate the cells and place in a sterile centrifuge tube. Pellet cells at 440 \(x\) g for 5 min using a low-speed bench-top centrifuge.

5. REMOVE the supernatant and resuspend the cells in 1 ml of fresh cell culture medium. Using a hemocytometer, count the cells and plate at a density of 3 \(\times\) 10\(^5\) cells per well in a 6-well plate. Agitate gently and incubate for 24 hr at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator).

6. THE FOLLOWING DAY (day 13 of neuronal culture) transiently transfect the HEK293 cells with mCherry cDNA in pcDNA3 expression construct using liposomal transfection reagent. Briefly, in a sterile microcentrifuge tube, add 500 \(\mu\)l of reduced serum medium and 5 \(\mu\)g of mCherry cDNA. Add 5 \(\mu\)l of liposomal buffering reagent and gently mix before leaving at room temperature for 5 min.

7. ADD 8.75 \(\mu\)l of liposomal transfection reagent and gently mix before leaving at room temperature for 30 min. Pipette the contents of microcentrifuge tube up and down twice, transfer drop-wise to each well on the 6-well tissue culture plate and incubate at 37 °C in a humidified 5% \(\text{CO}_2\) atmosphere (CO\(_2\) incubator).

8. THE FOLLOWING DAY (day 14 of neuronal culture), aspirate the HEK293 cell culture medium from each of the 6-wells and wash each well twice briefly with PBS (pH 7.4). Add 300 \(\mu\)l of Ca\(^{2+}\)/Mg\(^{2+}\)-chelating agent EDTA (0.48 mM) and 200 \(\mu\)l of trypsin-EDTA (0.02 - 0.48 mM) solution to each well and incubate at 37 °C for 5 min.

9. ADD 1 ml of fresh HEK293 cell culture medium per well (this quenches the trypsin) and aspirate the detached cells into a sterile 15 ml centrifuge tube. Centrifuge the cells at 440 \(x\) g for 5 min at room temperature and remove the supernatant. Resuspend the pellet in 500 \(\mu\)l of neuronal culture medium (pH 7.4).
10. Using a hemocytometer, count the cells and seed at a density of 30,000 cells per well in a 24-well tissue culture plate containing neurons. Agitate the plate to disperse the cells and incubate co-cultures at 37 °C in a humidified 5% CO₂ atmosphere (CO₂ incubator) for 24 hr.

6. Analysis of Synaptic Contacts and their Activity

1. After 23 hr in co-culture, investigate the formation of ‘active’ contacts between GABAergic medium spiny neurons and HEK293 cells using activity-dependent uptake of anti-synaptotagmin luminal domain-specific antibody conjugated with a fluorescent dye (Cy5, see Table 1).

   NOTE: The antibody will only gain access to the luminal domain of synaptotagmin, to which it binds when there is continuity between the presynaptic vesicle lumen and extracellular space. This specifically occurs during neurotransmitter release, making this antibody an excellent marker of the active presynaptic terminals.

2. Firstly rinse the co-cultures with neuronal medium (Neurobasal A medium, pH 7.4; see Table 1) and add Cy5-labeled mouse anti-synaptotagmin antibody, diluted 1:50 in neuronal medium (Neurobasal A medium, pH 7.4), to the cultures, for 30 min. Incubate the cells during this time at 37 °C in a humidified 5% CO₂ atmosphere (CO₂ incubator).

3. To remove the access of the antibody, wash the co-cultures briefly three times: first with cold normal PBS (pH 7.4), second with cold PBS (pH 7.4) containing 200 mM NaCl, and third with cold normal PBS (pH 7.4).

4. Fix the cells with 300 μl of 4% paraformaldehyde/4% sucrose in PBS (PFA, pH 7.4) for 10 min with agitation. Wash the cells briefly twice with PBS (pH 7.4) then with two longer 10 min washes.

5. Add glycine (0.3 M) to each well for 10 min with agitation to quench PFA.

6. Wash the cells briefly twice with PBS (pH 7.4) then with two longer 10 min washes before adding 300 μl of blocking solution (1% (w/v) bovine serum albumin (BSA) in PBS, pH 7.4) to reduce the non-specific binding of antibodies.

7. Aspirate the blocking solution and add the guinea pig anti-GABAₐR-γ2 antibody directed against the γ2 N-terminal domain (1:3,000 in PBS, pH 7.4) overnight at 4 °C.

8. The following day, aspirate the primary antibody from the wells and wash the cells briefly twice with PBS then with two longer 10 min washes. Permeabilize the cells using Triton X-100 (0.1 %) in blocking solution for 30 min at room temperature.

9. Permeabilize the cells using Triton X-100 (0.1 %) in blocking solution for 30 min at room temperature.

10. Wash the cells briefly twice with PBS (pH 7.4) then with two longer 10 min washes before adding either the mouse anti-glutamic acid decarboxylase (GAD) 65 antibody (1:4,000, Table 1) or the mouse anti-synapsin I antibody (1:1,000, Table 1) for 120 min at room temperature.

11. Wash the cells briefly twice with PBS (pH 7.4) then with two longer 10 min washes and then add blocking solution for 30 min at room temperature.

12. Centrifuge the appropriate secondary antibodies (typically goat anti-guinea pig IgG conjugated to Cy5, goat anti-mouse IgG conjugated to Alexa Fluor 488 or goat anti-mouse IgG Alexa Fluor 405, all at 2 μg/ml) to remove aggregates of antibodies at 21,910 x g for 10 min, and add antibodies (1:750) to blocking solution. Apply to appropriate wells for 1 hr and cover with aluminum foil to protect the fluorophores from light exposure and consequent photobleaching.

13. Finally, wash the cells briefly twice with PBS (pH 7.4) then with two longer 10 min washes to remove any unbound secondary antibody and mount the coverslips using mounting reagent (Prolong Gold, Table 1) with 10 μl per coverslip. Allow 24 hr to set at room temperature while protected from light before transferring to 4 °C for long term storage.

14. Analyze samples using a laser scanning confocal microscope with a 63X oil-immersion objective. Ensure the light levels and detector gain is adjusted to avoid saturation.

15. Observe potential synapse-like contacts as regions of co-localization between the presynaptic terminals positive for GAD65, synapsin I or Cy5-labelled anti-synaptotagmin and the postsynaptic HEK293 cells visualized by DIC or by the mCherry fluorescent indicator.

16. Count potential synapse-like contacts using Z-stack series of optical sections (8 - 10) through a depth of 4 - 5 μm per cell using the imaging software.

Representative Results

The protocol for this neuron-HEK293 cell co-culture model system has been finely tuned to allow optimal cell survival. In this system, formation of synapse-like contacts and their analysis relies on stable and consistent expression of all three GABAₐR subunits which assemble into a functional receptor. It is therefore important to use immunocytochemical analysis to test for subunit expression at the surface of HEK293 cells before adding them to neuronal cultures. In these experiments, cell surface expression of α1, β2 and γ2 subunits (Figure 1A), or α1, β3 and γ2 subunits (Figure 1B), was detected using subunit-specific antibodies which bind to the extracellular epitopes of these subunits. A high degree of co-localization between these subunits at the HEK293 surface was demonstrated.

After confirming the surface expression and co-localization of GABAₐR subunits in HEK293 cells, co-cultures were prepared using HEK293 cells expressing α1/β2/γ2 GABAₐR subunits and medium spiny neurons cultured for 14 days (14 days in vitro (DIV)). Cells in co-culture were incubated for 24 hr, fixed and analyzed using immunocytochemistry and confocal microscopy. Analysis of contacts indicated that GAD65-positive GABAergic axon terminals formed only sporadic contacts with the control HEK293 cells (Figure 2A, 2B). The number of contacts detected at 4 hr was 7.3 ± 0.9 per HEK293 cell, and this number was reduced to 5.5 ± 0.5 connections (mean ± SEM) per HEK293 cell at 24 hr after adding HEK293 cells to the cultured neurons. In contrast, GAD65-positive GABAergic axon terminals formed numerous synapse-like contacts with HEK293 cells expressing GABAₐR. The number of contacts obtained at 4 hr after adding HEK293 cells was 28.3 ± 4.7 per HEK293 cell, and this number was further increased to 52.1 ± 6.3 (mean ± SEM) per HEK293 cell at 24 hr in co-culture (Figure 2A, 2B).

To determine whether these synapse-like contacts were ‘active,’ i.e. supported vesicular transmitter release, a vesicle-luminal domain-specific anti-synaptotagmin Cy5-conjugated antibody was added to the co-culture medium after 23 hours of incubation. This antibody is only incorporated into presynaptic nerve terminals when a pore forms between the synaptic vesicle lumen and the extracellular fluid in the synaptic cleft during neurotransmitter release. Following release, the pore closes, leaving the synaptotagmin Cy5-conjugated fluorescent antibody attached to synaptotagmin inside the vesicle. In this way, only the vesicles actively engaged in neurotransmitter release were labeled with the antibody. In these experiments fewer if any contacts between the control HEK293 cells and the medium spiny neuron terminals were ‘active’ as shown by the lack of co-localization between the presynaptic GAD65/synaptotagmin fluorescence and mCherry fluorescence in HEK293 cells (Figure 3A).
In contrast, many ‘active’ contacts were formed between the medium spiny neuron terminals and α1/β2/γ2-expressing HEK293 cells, as revealed by a high degree of co-localization between GAD65/synaptotagmin and mCherry, expressed specifically in HEK293 cells (Figure 3B).

To test whether a different subtype of GABA<sub>A</sub>R can also promote synapse-like formation in vitro, we have co-cultured α1/β3/γ2 expressing HEK293 cells with medium spiny neurons. Again, control HEK293 cells rarely received contacts with synapsin-positive presynaptic terminals reaching 10.8 ± 0.48 (mean ± SEM) contacts per HEK293 cell after 24 hr in co-culture (Figure 4A left, 4B). However, HEK293 cells expressing α1/β3/γ2 GABA<sub>A</sub>Rs form significantly more synapse-like contacts with synapsin-positive presynaptic terminals of medium spiny neurons reaching 25.3 ± 0.27 (mean ± SEM) contacts per HEK293 cell after 24 hr in co-culture (Figure 4A right, 4B). This indicates that α1/β3/γ2 GABA<sub>A</sub>Rs expressed in HEK293 cells are also able to promote synaptic contact formation, albeit their potency is lower than the potency of α1/β2/γ2-containing GABA<sub>A</sub>Rs.

These experiments indicate that the co-culture model system developed in our laboratory permits quantitative analysis of synaptic contact formation in vitro as well as evaluation of the efficacy of different subtypes of GABA<sub>A</sub>Rs in this process. These experiments further demonstrate that GABA<sub>A</sub>Rs, in addition to being critical functional components of GABAergic synapses, may play a key role in the process of recognition and formation of synaptic contacts between inhibitory neurons and the appropriate neuronal target cells, independently of other synaptic adhesion proteins.

Figure 1. Immunocytochemical analysis of expression of GABA<sub>A</sub>R α1/β2/γ2 or α1/β3/γ2 in stable HEK293 cell lines. Antibodies recognizing the extracellular domains of GABA<sub>A</sub>R subunits were used to label receptors expressed at the cell surface. (A) HEK293 cell line expressing α1 (Alexa Fluor 488), β2 (Alexa Fluor 555) and γ2 (Cy5) at high levels. (B) HEK293 cell line expressing α1 (Alexa Fluor 488), β3 (Alexa Fluor 555) and γ2 (Cy5) subunits at high levels. Scale bar: 10 μm. Please click here to view a larger version of this figure.
Figure 2. GABAergic medium spiny neurons form synapse-like contacts with α1/β2/γ2-expressing HEK293 cells in co-culture. (A) Fluorescent labeling of presynaptic terminals with anti-GAD65 antibodies (in green) and HEK293 cells with mCherry (in red, left) or the GABA_A_R γ2 subunit (in blue, right), revealed points of co-localization between these markers indicating formation of synapse-like contacts after 4 or 24 hr in co-culture. Scale bar: 10 μm. (B) Quantitative analysis of synapse-like contacts. HEK293 cells were identified based on their shape as revealed by DIC imaging and/or mCherry expression, and the number of contacts between GAD-65 positive puncta (in green) and the surface of HEK293 cells was counted by eye in each optical section of a Z-stack series (8 - 10) per cell using the imaging software, and expressed as the number of contacts/cell. The graph shows the number of contacts between medium spiny neurons and control HEK293 cells (light grey) or α1/β2/γ2-HEK293 cells (black) after 4 and 24 hr in co-culture (mean ± SEM, n = 8 in each condition from two independent experiments). This figure has been modified from Fuchs et al. (2013)30. Please click here to view a larger version of this figure.
Figure 3. GABA<sub>Rs</sub> promote formation of active synaptic contacts. Immunolabeling of synapse-like contacts formed after 24 hours in coculture between medium spiny neuron terminals positive for GAD65 (Alexa Fluor 405 cyan) and (A) control HEK293 cells, or (B) HEK293-α1/β2/γ2 cells, both transiently transfected with mCherry construct (red). Active contacts are identified by co-localization between the vesicle luminal domain-specific anti-synaptotagmin antibody (Cy5) and GAD65-specific antibody both in presynaptic terminals, and mCherry expressed in HEK293 cells. Scale bar: 10 μm. Please click here to view a larger version of this figure.
Discussion

Although this protocol is not technically difficult to perform, there are several critical steps that must be followed to achieve the most accurate and repeatable co-culture assays. Firstly, cultured medium spiny neurons must be seeded at an optimal density. If seeded too sparsely, neurons tend to develop very slowly and survival is greatly reduced. On the other hand, if seeded too densely, neurons tend to aggregate which compromises the analysis of contacts with HEK293 cells. Secondly, it is recommended to transiently express a fluorescent reporter, GFP or mCherry, in HEK293 cells stably expressing GABA<sub>A</sub>Rs, prior to plating them into the co-culture. This allows reliable recognition of HEK293 cells, which can be compromised by similarity in shape and size between these cells and rare surviving glia cells in neuronal cultures. To achieve the efficient transfection with GFP or mCherry cDNA, HEK293 cell lines have to be in the exponential growth phase and seeded at the appropriate density in 6-well plates. Sparse seeding followed by transfection will cause cells to grow poorly, while over-seeding will prevent the cells from taking up the cDNA. Ideally, cells should be seeded so that they are between 70 - 90% confluent on the day of transfection. Thirdly, transfection must be optimized for each cell line used, as some cell lines are more sensitive than the others. This is because constitutive GABA<sub>A</sub>Rs expression in HEK293 cells reduces cell survival and the ability of cells to recover after transfection. Moreover, survival depends on the type of GABA<sub>A</sub>Rs expressed in HEK293 cells, with some cell lines being significantly more sensitive than the others. Transfection using liposomal reagent is an optimal method for expressing foreign proteins in fast growing cell lines, providing both the high transfection efficiency and level of expression. However, this reagent causes too much damage to slowly growing cell lines, for which we regularly use a non-liposomal transfection reagent. This works in a similar way to the liposomal reagent but the amount of DNA required for efficient transfection is significantly reduced. This allows greater cell survival (roughly 80 - 90% compared with 60% using liposomal reagent) but with lower transfection efficiency (60%). Lastly, the number of control HEK293 or α1/β2/γ2 HEK293 expressing cells added to neuronal cultures needs to be optimized. Adding too few cells...
comprises the successful analysis of contacts between HEK293 cells and neurons, because they become very rare. Conversely, adding too many HEK293 cells causes neuronal cell death within few hours.

Embryonic medium spiny neuron cultures should ideally be prepared using striatal tissue dissected from embryonic age 15 - 17. However, it often happens that embryos are slightly younger or older than the optimal age. In this case, the number of neurons seeded in culture will need to be varied. Tissue that is younger than E15 may need to be seeded at a slightly lower density, whilst tissue that is older than E17 may need to be seeded at a higher density, to allow optimal cell survival. Furthermore, cytosine arabinoside (Ara-C) may need to be added to older cultures to prevent growth of glia, which is more abundant in older tissue.

When creating co-cultures, it is important to plate the optimized number of transfected HEK293 or α1/β2/γ2 HEK293 expressing cells, as mentioned above. However, it may be necessary to determine this for each individual cell line, because of differences in their survival. Typically 30,000 cells in a maximum volume of 50 μl should be added to each well of a 24-well dish, which already contains 500 μl of neuronal culture medium, as this ensures that the conditioned neuronal medium is not diluted too much and that the conditions within each well remain fairly constant, e.g. the concentration of growth factors. Adding volumes greater than 50 μl to each well would generally kill the neurons.

One of the major disadvantages of the co-culture technique is that the neuronal cultures are created from dissociated cells grown as a monolayer, which means that the neurons have been removed from their normal microenvironment and are unable to establish their normal anatomical organization. Therefore they lack the appropriate connections, inputs and secreted molecules from other cells that may influence the initial stages of synapse development. For example, in vivo medium spiny neurons are densely innervated by glutamatergic inputs from the cortex, thalamus and other brain regions84, however, in our neuronal cultures glutamatergic synapses do not form because these inputs are damaged during dissection of the striatal tissue. How the absence of functional glutamatergic synapses in cultured medium spiny neurons affects their ability to form GABAergic synapses with each other and/or HEK293 cells expressing GABAARs remains an open question. This question could be easily addressed by culturing medium spiny neurons together with cortical glutamatergic neurons thereby allowing them to form functional synapses30 prior to the addition of HEK293 cells. An alternative approach would be to design a co-culture model system based on organotypic slice cultures, which maintain some of the cytoarchitecture which may be important for maturation and synapse formation. However, organotypic slice cultures have dense and heterogeneous neuropil which can compromise the analysis performed here. Another important disadvantage of using co-culture assays is that GABAARs expressed at the surface of HEK293 cells are not clustered as they are in neurons, although this appears not to be necessary for synapse formation given a high enough surface expression32. For example, in the rodent brain and in hippocampal cultures, the α1 GABAAR subunit is found in most GABAergic synapses on all postsynaptic domains of pyramidal cells. However, the δ2 is specifically located in a subset of synapses on the somata and dendrites but is highly enriched in the axon initial segment, as revealed by immunofluorescence and electron microscopy34. Given that synapse formation in the co-cultures can still be reliably detected and analyzed30, this suggests that the density of GABAδ2Rs at the cell surface of HEK293 cells may be similar to, or even higher than the density of these receptors within synaptic clusters in neurons. This can explain, at least in part, why synaptic adhesion proteins, such as neurexin, and postsynaptic density proteins, such as gephyrin, are not necessary for synapse formation in the co-cultures, if the appropriately assembled GABAδ2Rs are present at sufficient density.

It is well documented that GABAδ2Rs are structurally and functionally heterogeneous, and that the receptor subunit composition determines their subcellular localization and pharmacological properties. For example, incorporation of the 2 subunits is known to be a prerequisite for the synaptic localization of GABAδ2Rs while the subunit is almost exclusively present in extrasynaptic GABAδ2Rs. The receptors that incorporate only δβ combinations are also thought to be predominantly localized to the extrasynaptic domains12-14. Whether this specificity is maintained in our co-culture system can be easily tested by transiently transfecting 2 or subunit cDNAs into HEK293 cell lines stably expressing α and β subunits, before adding them to neuronal cultures. Our preliminary experiments using this approach have suggested that synaptic contacts are readily formed only in the presence of the 2 subunit, indicating that the specificity observed in vivo is likely to be preserved in vitro (data not shown).

Furthermore, GABAδ2Rs incorporating different α subunits are selectively localized to synaptic contacts formed with specific types of presynaptic neurons. For example, in the globus pallidus, the α1-GABAδ2Rs are generally found at striatopallidal (st-P) and pallidopallidal (GP-GP) synapses, which are located on dendrites and somatic contacts of the medium spiny neurons, respectively. The α3-GABAδ2Rs are located in perisomatic regions of medium spiny neurons and are contacted by local GP axon collaterals, whilst the α2-GABAδ2Rs are located on distal dendrites of these neurons and contacted primarily by inputs from the stratum32. Expression of specific α subunits in different types of synapses and in different neuronal compartments has also been demonstrated in other brain areas such as hippocampus32 and neocortex16-20. These findings raise the question as to how the specific inhibitory synapses are formed in the brain. Does the adhesion of a specific type of presynaptic terminal induce the insertion of specific GABAδ2R subtypes into the postsynaptic density? Are the receptors trafficked to specific subcellular locations according to their subunit composition, where their plasma membrane insertion is a prerequisite for the adhesion of axonal terminals of specific origin? To date, these questions remain unanswered. The use of reduced model systems such as the co-culture model system, allow us to start answering this complex questions because the system is easily amenable to transfection of DNA constructs and application of reagents and, importantly, it is suitable for live cell imaging analysis30. Thus, using this model system we can start testing the role of individual molecules, including different types of GABAδ2Rs, known to be present at synaptic contacts. Another advantage is that synapses in this model system form rapidly, within minutes to hours, reducing the duration of experiments. Similar co-culture model systems were successfully employed in the past to screen for the novel synaptogenic molecules27-30. Further development of co-culture models is important because they have the potential to advance
our knowledge of the molecular mechanisms which guide the ‘normal’ brain development and thus increase our understanding of how these mechanisms are altered in many neurodevelopmental diseases, such as epilepsy, schizophrenia, autism spectrum disorders and many others.

**Disclosures**

The authors declare that they have no competing financial interests.

**Acknowledgements**

We would like to acknowledge financial support from the MRC UK (G0800498). We would also like to thank Professor J-M Fritschy, University of Zurich, for providing the GABA<sub>A</sub>-R subunit-specific y2 antibody and Professor R. Harvey, UCL School of Pharmacy, for providing the pcDNA 3.1<sup>®</sup> expression vectors containing antibiotic resistance genes for production of stably transfected HEK293 cell lines.

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