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Anatomical Characterization of the Type-1 cannabinoid receptors in specific brain cell populations of mutant mice

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THÈSE EN COTUTELLE PRÉSENTÉE

POUR OBTENIR LE GRADE DE

DOCTEUR DE

L'UNIVERSITÉ DE BORDEAUX

ET DE L'UNIVERSITÉ DE PAYS BASQUE (UPV/EHU)

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ÉCOLE DOCTORALE DU UNIVERSITÉ DU PAYS BASQUE

SPÉCIALITÉ NEUROSCIENCES

Par Ana GUTIÉRREZ RODRÍGUEZ

**ANATOMICAL CHARACTERIZATION OF THE TYPE-1
CANNABINOID RECEPTORS IN SPECIFIC BRAIN CELL
POPULATIONS OF MUTANT MICE**

Sous la direction de Giovanni MARSICANO
et de Pedro GRANDES MORENO

Soutenue le 2 décembre 2016

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Titre : Caractérisation anatomique des récepteurs cannabinoïdes de type 1 dans des populations de cellules cérébrales spécifiques de souris mutantes.

Résumé : Dans cette thèse l'expression du récepteur CB₁ dans l'hippocampe de souris mutantes ré-exprimant spécifiquement le gène spécifiquement dans certains types cellulaires du cerveau tels que : les neurones glutamatergiques du télencéphale dorsal, et les neurones GABAergiques a été analysé. De plus, dans le but de connaître la distribution anatomique exacte des récepteurs CB₁ astrogliaux par rapport aux synapses excitatrices et inhibitrices, j'ai étudié l'expression des récepteurs CB₁ dans les astrocytes de souris exprimant le récepteur CB₁ seulement dans les astrocytes et une souris mutante ciblée pour exprimer la protéine cytoplasmique hrGFP diffusible dans les cellules astrogliales ce qui permet une meilleure détection des prolongements astrocytaires. Les conclusions de ce travail de thèse sont les suivantes: La distance la plus commune entre le récepteur CB₁ astroglial et la synapse la plus proche est de 400 à 800 nm. La majorité des synapses entourées par des astrocytes immunopositifs pour le récepteur CB₁ dans l'hippocampe, est de nature excitatrice. Les souris mutantes réexprimants le récepteur CB₁ caractérisées dans ce travail de thèse montre : 1) expression du récepteur CB₁ dans différents types cellulaires, 2) la réexpression est limitée à une population neuronale particulière ou aux astrocytes, 3) les niveaux endogènes de récepteurs CB₁ sont maintenus dans les différents types cellulaires ré-exprimants le récepteur CB₁. De façon générale, ces résultats nous montrent que les souris mutantes ré-exprimants le récepteur CB₁ sont d'excellents outils pour l'étude fonctionnelle et translationnelles sur le rôle de ce récepteur dans le cerveau sain ou pathologique.

Mots clés : Système endocannabinoïde, synapses, microscopie électronique, hippocampe, souris mutantes.

Title : Anatomical Characterization of the Type-1 cannabinoid receptors in specific brain cell populations of mutant mice

Abstract : The Cannabinoid Type I receptor protein (CB₁) expression in the hippocampus of rescue mice modified to express the gene exclusively in specific brain cell types : such as dorsal telencephalic glutamatergic neurons, or GABAergic neurons have been analysed. Furthermore, aiming at knowing the exact anatomical distribution of the astroglial CB₁ receptors with respect to the excitatory and inhibitory synapses, the CB₁ receptor expression in astrocytes of mouse expressing CB₁ receptor only in astrocytes and mutant mouse expressing the protein hrGFP into astrocytes (that allows for better detection of the astrocytic processes) have been also investigated. The results showed that the majority of the hippocampal synapses surrounded by CB₁ receptor immunopositive astrocytes in the 400-800 nm range are of excitatory nature. Moreover, the CB₁ receptor rescue mutant mice characterized in this Doctoral Thesis have proven 1) to express CB₁ receptors in specific brain cell types; 2) the re-expression is limited to the particular brain cell populations; 3) the endogenous levels of CB₁ receptors are maintained in the brain cell types re-expressing the receptor. Which makes this mutant mice excellent tools for functional and translational investigations on the role of the CB₁ receptors in the normal and diseased brain.

Keywords : Endocannabinoid system, synapses, electron microscopy, hippocampus, mutant rodents.

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1 SUMMARIES

1.1 FRENCH SUMMARY

INTRODUCTION

Le système endocannabinoïde est principalement formé par des ligands endogènes lipides (endocannabinoïdes), le Récepteur au cannabinoïde de type 1 (CB₁) et de type 2 (CB₂), ainsi que leurs protéines de synthèse, de transport et de dégradation (Piomelli, 2003; Marsicano and Lutz, 2006; Katona and Freund, 2012; Pertwee, 2015; Lu and Mackie, 2016). Ce système est impliqué dans plusieurs fonctions cérébrales, le développement prénatal du cerveau et de nombreuses fonctions cognitives (Marsicano and Kuner, 2008; Bellocchio et al., 2010; Katona and Freund, 2012). L'expression du récepteur CB₁ est très élevée dans les neurones inhibiteurs GABAergiques concentrée sur leurs terminaisons synaptiques (Katona et al., 1999; Kawamura et al., 2006; Ludányi et al., 2008; Marsicano and Kuner, 2008; Katona and Freund, 2012; De-May and Ali, 2013; Steindel et al., 2013; Hu and Mackie, 2015) et faible dans les neurones glutamatergiques avec une localisation rare des récepteurs sur leurs boutons axonaux (Marsicano et al., 2003; Domenici et al., 2006; Takahashi and Castillo, 2006; Katona et al., 2006; Monory et al., 2006; Kamprath et al., 2009; Bellocchio et al., 2010; Puente et al., 2011; Reguero et al., 2011; Steindel et al., 2013; Ruehle et al., 2013; Soria-Gómez et al., 2014). Les récepteurs CB₁ sont aussi localisés dans les membranes mitochondriales dans le cerveau et les muscles (Bénard et al., 2012; Hebert-Chatelain et al., 2014a; b; Koch et al., 2015) et dans les astrocytes où ils jouent un rôle clé dans le contexte de la synapse tripartite (Navarrete and Araque, 2008, 2010; Perea et al., 2009; Han et al., 2012; Gómez-Gonzalo et al., 2014; Navarrete et al., 2014; Metna-Laurent and Marsicano, 2015; Viader et al., 2015; Oliveira da Cruz et al., 2016). En raison de la très grande différence de niveau d'expression entre les différents types de cellules (Marsicano and Lutz, 1999), il est extrêmement difficile d'identifier les faibles niveaux du récepteur CB₁ dans ces types cellulaires et/ou dans les compartiments subcellulaires de cerveaux de souris de type sauvage (Busquets-Garcia et al., 2015). En effet, auparavant, de faibles niveaux d'expression du récepteur CB₁ ont été considérés comme bruit de fond (Freund et al., 2003). Pour établir correctement la distribution anatomique des récepteurs CB₁ dans des types cellulaires spécifiques, indépendamment des niveaux d'expression endogènes, de nouvelles techniques d'analyse sont nécessaires. Par conséquent, la stratégie génétique de "sauvetage" (rescue) basée sur l'utilisation du système Cre/loxP (Balthasar et al., 2005) est apparue comme un excellent outil pour la détermination de l'expression du récepteur CB₁ endogène dans des types cellulaires spécifiques (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015). Ceci est particulièrement utile pour élucider les niveaux réels des récepteurs CB₁ dans des types cellulaires spécifiques exprimant de faibles niveaux de la protéine CB₁, ce qui élimine les interférences causées par des cellules qui expriment des niveaux élevés de récepteurs CB₁. Cependant, il est très important de vérifier que la ré-expression est limitée aux types de cellules recherchées et qu'elle maintient les niveaux des récepteurs CB₁ endogènes, où les séquences régulatrices du récepteur sont normalement activées.

J'ai examiné dans ma thèse l'expression du récepteur CB₁ dans l'hippocampe de souris mutantes ré-exprimant spécifiquement le gène spécifiquement dans

certains types cellulaires du cerveau tels que : les neurones glutamatergiques du télencéphale dorsal (souris Glu- CB_1 -RS), et les neurones GABAergiques (souris GABA- CB_1 -RS).

De plus, dans le but de connaître la distribution anatomique exacte des récepteurs CB_1 astrogliaux par rapport aux synapses excitatrices et inhibitrices, j'ai également étudié l'expression des récepteurs CB_1 dans les astrocytes de souris GFAP- CB_1 -RS (exprimant le récepteur CB_1 seulement dans les astrocytes) et une souris mutante ciblée pour exprimer la protéine cytoplasmique hrGFP diffusible dans les cellules astrogliales (GFAPhrGFP- CB_1 -WT) ce qui permet une meilleure détection des prolongements astrocytaires.

MATERIEL ET METHODES

Souris: CB_1 -WT, CB_1 -KO (absence des récepteurs CB_1), GABA- CB_1 -RS, Glu- CB_1 -RS, GABA- CB_1 -KO (portant une délétion sélective du récepteur CB_1 dans les neurones GABAergiques), Glu- CB_1 -KO (portant une délétion sélective du récepteur CB_1 dans les neurones glutamatergiques du télencéphale dorsal), GFAP- CB_1 -RS, GFAP- CB_1 -KO (portant une délétion sélective du récepteur CB_1 dans les astrocytes), GFAPhrGFP- CB_1 -WT et GFAPhrGFP- CB_1 -KO (absence des récepteurs CB_1 dans les astrocytes). Les animaux ont été profondément anesthésiés et perfusés par voie transcardiaque.

Une méthode de pré-enrobage immunogold pour la microscopie électronique a été appliquée aux coupes hippocampales des souris CB_1 -WT, CB_1 -KO, GABA- CB_1 -RS, Glu- CB_1 -RS, GABA- CB_1 -KO et Glu- CB_1 -KO. Egalement, une méthode combinant un pré-enrobage immunogold et une immunoperoxidase a été utilisée pour les coupes hippocampales des souris GFAP- CB_1 -RS, GFAP- CB_1 -KO, GFAPhrGFP- CB_1 -WT et GFAPhrGFP- CB_1 -KO. Les expériences ont toujours été réalisées sur trois animaux différents pour chaque condition. Afin de déterminer statistiquement la distribution des récepteurs CB_1 dans les différents types de cellules et compartiments cellulaires étudiés, un test de normalité a été appliqué (test de normalité de Kolmogorov-Smirnov) avant l'exécution des analyses statistiques. Les données ont été analysées à l'aide de tests non-paramétriques (Man-Whitney U test lorsque $k=2$ ou Kruskal-Wallis test lorsque $k>2$).

RESULTATS

La proportion de récepteurs CB_1 immunopositifs au niveau des terminaisons synaptiques excitatrices du stratum radiatum de la région CA1 de l'hippocampe des Glu- CB_1 -RS ($21.89\% \pm 1.21$), CB_1 -WT et GABA- CB_1 -KO n'est pas statistiquement différente. Cependant, le pourcentage de terminaisons immunopositives disparaît virtuellement chez les animaux Glu- CB_1 -KO, GABA- CB_1 -RS et CB_1 -KO. En outre, aucune différence statistique n'est trouvée pour la densité de récepteur CB_1 , exprimée comme étant le nombre de particules par μm du périmètre de la cellule (particules/ μm), entre les terminaisons synaptiques excitatrices des Glu- CB_1 -RS (0.45 ± 0.02), CB_1 -WT et GABA- CB_1 -KO.

L'analyse du tiers interne de la couche dentée moléculaire montre des pourcentages de récepteurs immuno-positifs au niveau des terminaisons

synaptiques excitatrices comparables chez les Glu- CB_1 -RS ($53.19\% \pm 2.89$), CB_1 -WT et GABA- CB_1 -KO. Cette proportion est quasi nulle chez les Glu- CB_1 -KO, GABA- CB_1 -RS et CB_1 -KO. La densité d'immunoparticules de récepteurs CB_1 est statistiquement similaire chez les Glu- CB_1 -RS (0.39 ± 0.02) CB_1 -WT et GABA- CB_1 -KO.

Le nombre de récepteurs CB_1 immunopositifs au niveau des terminaisons neuronales des synapses inhibitrices chez les GABA- CB_1 -RS ($77.92\% \pm 2.63$), CB_1 -WT et Glu- CB_1 -KO est maintenu dans le stratum radiatum de CA1. Cependant, la proportion disparaît chez les GABA- CB_1 -KO, Glu- CB_1 -RS et CB_1 -KO. Il n'existe pas de différence significative pour la densité des immunoparticules du récepteur CB_1 entre les terminaisons des synapses inhibitrices des GABA- CB_1 -RS (4.33 ± 0.11), CB_1 -WT et Glu- CB_1 -KO.

La proportion de récepteurs CB_1 immunopositifs au niveau des terminaisons des synapses inhibitrices du tiers interne de la couche dentée moléculaire est similaire entre les GABA- CB_1 -RS ($85.07\% \pm 1.76$), CB_1 -WT et Glu- CB_1 -KO. Ces valeurs disparaissent quasiment chez les GABA- CB_1 -KO, Glu- CB_1 -RS et CB_1 -KO. Il n'existe pas de différence significative pour la densité de récepteurs CB_1 au niveau des synapses inhibitrices des GABA- CB_1 -RS (7.47 ± 0.14), CB_1 -WT et Glu- CB_1 -KO.

La proportion de récepteurs CB_1 immunopositifs au niveau des prolongements astrocytaires du stratum radiatum n'est pas significativement différente entre les animaux CB_1 -WT et GFAP- CB_1 -RS ($37.12\% \pm 3.79$)

Chez les animaux GFAPhrGFP- CB_1 -WT, la proportion d'éléments astrocytaires immunomarqués pour le récepteur CB_1 augmente de $59.91\% \pm 3.29$, devenant statistiquement différent comparé au CB_1 -WT. A l'inverse, seuls de rares récepteurs CB_1 sont marqués dans les astrocytes des souris GFAP- CB_1 -KO, CB_1 -KO et chez les souris GFAPhrGFP- CB_1 -KO. La densité d'immunoparticule pour le récepteur CB_1 au niveau de la membrane astrocytaire (particules/ μm) est également analysée. Nos données montrent qu'il n'y a pas de différence significative entre les CB_1 -WT et les GFAP- CB_1 -RS (0.128 ± 0.020), alors que nous avons une plus forte densité chez les GFAPhrGFP- CB_1 -WT (0.384 ± 0.039).

L'analyse du tiers interne de la couche dentée moléculaire montre des pourcentages de récepteurs immuno-positifs au niveau des prolongements astrocytaires comparables chez les GFAP- CB_1 -RS ($39.84\% \pm 3.50$) et les CB_1 -WT alors que cette proportion augmente de manière significative ($59.99\% \pm 3.37$) chez les GFAPhrGFP- CB_1 -WT. Seuls des particules résiduelles sont observées chez les souris GFAP- CB_1 -KO, GFAPhrGFP- CB_1 -KO et CB_1 -KO. Enfin, aucune différence significative de densité du récepteur CB_1 n'est observée dans les prolongements astrocytaires des souris CB_1 -WT et GFAP- CB_1 -RS (0.138 ± 0.016). On observe une différence significative de densité du récepteur CB_1 dans les prolongements astrocytaires des souris GFAPhrGFP- CB_1 -WT comparée au CB_1 -WT (0.334 ± 0.033).

Grâce à une meilleure détection des astrocytes obtenue avec la protéine hr-GFP en microscopie électronique, nous avons mesuré la distance entre les immunoparticules du récepteur CB_1 astroglial et le point central de la synapse la plus proche entourée par l'élément astrocytaire immunopositif chez les GFAPhrGFP- CB_1 -WT. Dans le CA1, $51.07\% \pm 3.79$ de toutes les synapses

entourées par un prolongement astrocytaire se trouvent à une distance de 400 à 800 nm de la particule astrocytaire la plus proche. Parmi celles-ci, $43.33\% \pm 1.01$ sont des synapses excitatrices alors que $9.91\% \pm 1.93$ sont des synapses inhibitrices. Dans la couche moléculaire dentée, $57.25\% \pm 3.19$ des synapses entourées par un prolongement astrocytaire se trouvent entre 400 et 800 nm de la particule du récepteur CB₁ astroglial. $46.67\% \pm 2.17$ sont au niveau de synapses excitatrices alors que $10.59\% \pm 2.18$ se trouvent au niveau de synapses inhibitrices.

CONCLUSIONS

Les conclusions de ce travail de thèse sont les suivantes:

1. Le niveau d'expression et la localisation des récepteurs CB₁ des neurones glutamatergiques de l'hippocampe de souris Glu-CB₁-RS est similaire à celui de souris CB₁-WT.
2. Le niveau d'expression et la localisation des récepteurs CB₁ des neurones GABAergiques de l'hippocampe de souris GABA-CB₁-RS est similaire à celui des CB₁-WT.
3. Le niveau d'expression et la localisation des récepteurs CB₁ dans les astrocytes de l'hippocampe de souris GFAP-CB₁-RS est similaire à celui des CB₁-WT.
4. La détection des récepteurs CB₁ dans les astrocytes de l'hippocampe de souris GFAPhrGFP-CB₁-WT est significativement supérieure comparée aux animaux CB₁-WT avec un marqueur GFAP.
5. La distance la plus commune entre le récepteur CB₁ astroglial et la synapse la plus proche est de 400 à 800 nm.
6. La majorité des synapses entourées par des astrocytes immunopositifs pour le récepteur CB₁ dans la région CA1 et de la couche moléculaire dentée, est de nature excitatrice.
7. Les souris mutantes réexprimants le récepteur CB₁ caractérisées dans ce travail de thèse montre: 1) expression du récepteur CB₁ dans différents types cellulaires, 2) la réexpression est limitée à une population neuronale particulière ou aux astrocytes, 3) les niveaux endogènes de récepteurs CB₁ sont maintenus dans les différents types cellulaires ré-exprimants le récepteur CB₁.
8. De façon générale, ces résultats nous montrent que les souris mutantes ré-exprimants le récepteur CB₁ sont d'excellents outils pour l'étude fonctionnelle et translationnelles sur le rôle de ce récepteur dans le cerveau sain ou pathologique.

1.2 SPANISH SUMMARY

INTRODUCCIÓN

El sistema endocannabinoide está compuesto, principalmente, por un grupo de ligandos lipídicos endógenos denominados endocannabinoides, los Receptores Cannabinoide de Tipo 1 (CB₁) y Tipo 2 (CB₂), así como las proteínas que participan en la síntesis, transporte y la degradación de los endocannabinoides (Piomelli, 2003; Marsicano and Lutz, 2006; Katona and Freund, 2012; Pertwee, 2015; Lu and Mackie, 2016). Este sistema se encuentra involucrado en numerosos mecanismos cerebrales, en el desarrollo del sistema nervioso prenatal y en diversas funciones cognitivas (Marsicano and Kuner, 2008; Bellocchio y coils., 2010; Katona and Freund, 2012). Respecto a la expresión del receptor CB₁, se ha descrito como los terminales inhibidores de neuronas GABAérgicas presentan elevados niveles de expresión de este receptor (Katona y coils., 1999; Kawamura y coils., 2006; Ludányi y coils., 2008; Marsicano and Kuner, 2008; Katona and Freund, 2012; De-May and Ali, 2013; Steindel y coils., 2013; Hu and Mackie, 2015) mientras en los terminales axónicos de neuronas glutamatérgicas encontramos niveles de expresión más bajos (Marsicano y coils., 2003; Domenici y coils., 2006; Takahashi and Castillo, 2006; Katona y coils., 2006; Monory y coils., 2006; Kamprath y coils., 2009; Bellocchio y coils., 2010; Puente y coils., 2011; Reguero y coils., 2011; Steindel y coils., 2013; Ruehle y coils., 2013; Soria-Gómez y coils., 2014). Por otro lado, a nivel subcelular, se ha descrito la expresión del receptor CB₁ en las membranas de las mitocondrias del cerebro y del músculo (Bénard y coils., 2012; Hebert-Chatelain y coils., 2014a; b; Koch y coils., 2015). Así mismo, se ha descrito como el receptor CB₁ está también localizado en los astrocitos en donde desempeña un papel fundamental en el contexto de la sinapsis tripartita (Navarrete and Araque, 2008, 2010; Perea y coils., 2009; Han y coils., 2012; Gómez-Gonzalo y coils., 2014; Navarrete y coils., 2014; Metna-Laurent and Marsicano, 2015; Viader y coils., 2015; Oliveira da Cruz y coils., 2016).

Las diferencias existentes en los niveles de expresión de este receptor entre los diversos tipos celulares (Marsicano and Lutz, 1999), hacen que resulte extremadamente difícil detectar su expresión en tipos de células y / o en compartimentos sub-celulares donde su expresión es fisiológicamente muy baja (Busquets-Garcia y coils., 2015). En ese sentido, en el pasado, la presencia de niveles bajos de receptor CB₁ fue interpretada erróneamente como mera tinción de fondo en análisis anatómicos de ratones de silvestres (CB₁-WT) (Freund y coils., 2003).

De esta manera, para la correcta caracterización anatómica de la distribución subcelular de los receptores CB₁ en cualquier tipo celular, independientemente de sus niveles fisiológicos de expresión, se hace necesarias mejores técnicas de análisis. En ese sentido, las estrategias de "rescate" genético mediante el uso del sistema Cre/LoxP (Balthasar y coils., 2005) suponen una gran herramienta de estudio, ya que permiten analizar la expresión del receptor CB₁ endógeno exclusivamente en tipos celulares determinados (Ruehle y coils., 2013; Soria-Gómez y coils., 2014; de Salas-Quiroga y coils.). En concreto, esta estrategia resulta particularmente útil para determinar el patrón de expresión del receptor

CB₁ en tipos específicos de células donde sus niveles de expresión son bajos y puedan encontrarse enmascarados por la enorme expresión de este receptor en otros tipos de células. Sin embargo, es de gran importancia comprobar que la re-expresión del receptor CB₁ mantiene sus niveles endógenos y se limita a las células deseadas.

En mi tesis doctoral se pretende examinar la expresión receptor CB₁ en el hipocampo de ratones rescatados genéticamente los cuales expresan el gen del receptor CB₁ exclusivamente en determinados tipos celulares: neuronas glutamatérgicas telencefálicas dorsales (ratones Glu-CB₁-RS) y neuronas GABAérgicas (ratones GABA-CB₁-RS).

Así mismo, con el objetivo de conocer la distribución del receptor CB₁ de astrocitos, respecto a las sinapsis excitadoras e inhibitoras, se realizará el análisis de la expresión del CB₁ en ratones que expresan el receptor CB₁ únicamente en astrocitos, (GFAP-CB₁-RS) y en ratones transgénicos que expresan en astrocitos la proteína citoplasmática difusible hrGFP (ratones GFAPhrGFP-CB₁-WT) que permite una mejor detección de la superficie de los astrocitos.

MATERIAL Y MÉTODOS

Ratones: CB₁-WT, CB₁-KO (no expresan receptor CB₁), GABA-CB₁-RS, Glu-CB₁-RS, GABA-CB₁-KO (no expresan el receptor CB₁ en neuronas GABAérgicas), Glu-CB₁-KO (no expresan el receptor CB₁ en neuronas glutamatérgicas telencefálicas dorsales), GFAP-CB₁-RS, GFAP-CB₁-KO (no expresan el receptor CB₁ en astrocitos), GFAPhrGFP-CB₁-WT y GFAPhrGFP-CB₁-KO (no expresan el receptor CB₁), fueron profundamente anestesiados y sometidos a perfusión transcardiaca.

La técnica de inmuno-oro preinclusión para Microscopía Electrónica fue aplicada a secciones del hipocampo de ratones CB₁-WT, CB₁-KO, GABA-CB₁-RS, Glu-CB₁-RS, GABA-CB₁-KO y Glu-CB₁-KO. Mientras que a las secciones de hipocampo de ratones GFAP-CB₁-RS, GFAP-CB₁-KO, GFAPhrGFP-CB₁-WT y GFAPhrGFP-CB₁-KO se les aplicó la técnica de inmuno-oro e inmunoperoxidasa preinclusión para Microscopía Electrónica. En todos los casos los análisis se llevaron a cabo utilizando tres animales de cada tipo. En estos animales se analizó la distribución anatómica del receptor CB₁ en los diferentes tipos celulares y compartimentos subcelulares. Los datos obtenidos fueron analizados aplicando primero la prueba de normalidad de Kolmogorov-Smirnov para posteriormente analizar estadísticamente los datos mediante test no paramétricos de contraste de medias (prueba de Man-Whitney cuando k = 2 o la prueba de Kruskal-Wallis cuando k > 2).

RESULTADOS

La comparación del porcentaje de terminales excitadores marcados con CB₁ en el stratum radiatum del CA1 en el ratón Glu-CB₁-RS (21,89% ± 1,21) en el ratón CB₁-WT y en el ratón GABA-CB₁-KO no fue estadísticamente significativo. Sin embargo, este valor prácticamente desaparece en los ratones Glu-CB₁-KO, GABA-CB₁-RS and CB₁-KO. Además, no se encontraron diferencias significativas entre el valor de la densidad, expresada como el número de partículas del receptor

CB₁ por μm del perímetro de la membrana (partículas/ μm), entre los terminales excitadores de Glu-CB₁-RS (0.45 ± 0.02), CB₁-WT y GABA-CB₁-KO.

En el tercio interno de la capa molecular del giro dentado similares porcentajes de terminales excitadores que presentan receptor CB₁ se encontraron en Glu-CB₁-RS ($53.19\% \pm 2.89$), CB₁-WT y GABA-CB₁-KO. Desapareciendo prácticamente esta proporción en los ratones Glu-CB₁-KO, GABA-CB₁-RS and CB₁-KO. La densidad de inmunopartículas de CB₁ fue estadísticamente similar entre Glu-CB₁-RS (0.39 ± 0.02) CB₁-WT y GABA-CB₁-KO.

El valor del porcentaje de terminales inhibidores marcados positivamente para el receptor CB₁ en los ratones GABA-CB₁-RS ($77.92\% \pm 2.63$), CB₁-WT y Glu-CB₁-KO se mantuvo en el stratum radiatum de la región CA1. Sin embargo, esta proporción prácticamente desaparece en los animales GABA-CB₁-KO, Glu-CB₁-RS y CB₁-KO. No se encontraron diferencias significativas tras la comparación de la densidad de partículas de CB₁ en los terminales inhibidores de los ratones GABA-CB₁-RS (4.33 ± 0.11 partículas/ μm), CB₁-WT y Glu-CB₁-KO.

En el tercio interno de la capa molecular del giro dentado, no se obtuvieron diferencias significativas comparando los valores obtenidos para el porcentaje de terminales inhibidores marcados con CB₁ en ratones GABA-CB₁-RS ($85.07\% \pm 1.76$), CB₁-WT y Glu-CB₁-KO. Solamente valores residuales del receptor CB₁ se hallaron en animales GABA-CB₁-KO, Glu-CB₁-RS y CB₁-KO. No hubo diferencias significativas tras el análisis estadístico de la densidad de partículas de CB₁ en los terminales inhibidores entre los ratones GABA-CB₁-RS (7.47 ± 0.14), CB₁-WT y Glu-CB₁-KO.

En el stratum radiatum de la región CA1 el análisis de los procesos astrocíticos que presentaban un marcaje positivo para CB₁ no resultó estadísticamente significativo al comparar los valores obtenidos en el ratón CB₁-WT y el ratón GFAP-CB₁-RS ($37.12\% \pm 3.79$). Sin embargo, la proporción de procesos astrocíticos positivos para CB₁ en el ratón GFAPhrGFP-CB₁-WT aumentó hasta alcanzar el valor de $59.91\% \pm 3.29$, un aumento que resultó estadísticamente significativo al compararlo con CB₁-WT. Por otra parte, sólo niveles residuales del receptor CB₁ se encontraron en los astrocitos de los animales GFAP-CB₁-KO, CB₁-KO y GFAPhrGFP-CB₁-KO.

Así mismo, la densidad de partículas de CB₁ en la membrana de los procesos astrocíticos (partículas/ μm) fue similar entre los valores del ratón CB₁-WT y del ratón GFAP-CB₁-RS (0.128 ± 0.020). Por otro lado, un gran aumento estadísticamente significativo en la densidad del receptor se obtuvo en el ratón GFAPhrGFP-CB₁-WT (0.384 ± 0.039).

En la capa molecular del giro dentado del hipocampo, tras analizar el porcentaje de procesos astrocíticos positivos para CB₁ no se obtuvieron diferencias significativas entre los valores del ratón GFAP-CB₁-RS ($39.84\% \pm 3.50$) y el ratón CB₁-WT, mientras que la proporción aumentó significativamente hasta ser del $59.99\% \pm 3.37$ en el ratón GFAPhrGFP-CB₁-WT. Por otra parte, solamente se encontraron partículas de oro residuales en los ratones GFAP-CB₁-KO, GFAPhrGFP-CB₁-KO y CB₁-KO. Además, no hubo diferencias significativas tras la comparación de la densidad del receptor en los astrocitos del ratón CB₁-WT y el ratón GFAP-CB₁-RS (0.138 ± 0.016). Sin embargo, se obtuvieron diferencias

significativas al comparar estos valores con el obtenido en el ratón GFAPhrGFP- CB_1 -WT (0.334 ± 0.033).

Una vez demostrado que una mejor detección del CB_1 en astrocitos es posible al utilizar la proteína hr-GFP como marcador en el ratón GFAPhrGFP- CB_1 -WT, se analizó la distancia existente entre el receptor CB_1 astroglial y el punto medio de la sinapsis más cercana. Así, en el stratum radiatum de CA1, el $51,07\% \pm 3,79$ de las sinapsis totales que rodean a un proceso astrocítico estaban situadas a una distancia entre 400 y 800 nm del receptor CB_1 astroglial. Por otra parte, el $43\% \pm 1,01$ de estas sinapsis fueron identificados como sinapsis excitadoras mientras que sólo el $9,91\% \pm 1,93$ eran inhibitoras. En la capa molecular del giro dentado el $57,25\% \pm 3,19$ del total de las sinapsis que rodean a un proceso astrocítico se encontraron entre 400 y 800 nm de distancia del receptor CB_1 de los astrocitos. De estas sinapsis el $46,67\% \pm 2,17$ fueron identificadas como excitadoras mientras que el $10,59\% \pm 2,18$ eran sinapsis inhibitoras.

CONCLUSIONES

Las conclusiones obtenidas en esta Tesis son las que siguen:

- 1- En el ratón Glu- CB_1 -RS se mantienen los mismos niveles de expresión de los receptores CB_1 hipocampal que en los ratones tipo silvestre CB_1 -WT en las neuronas glutamatérgicas del telencéfalo dorsal.
- 2- En el ratón GABA- CB_1 -RS se mantienen los mismos niveles de expresión de los receptores CB_1 hipocampal que en los ratones tipo silvestre CB_1 -WT únicamente en neuronas GABAérgicas.
- 3- En el ratón GFAP- CB_1 -RS se mantienen los mismos niveles de expresión de los receptores CB_1 hipocampal que en procesos astrocíticos de los ratones tipo silvestre CB_1 -WT.
- 4- La utilización de hrGFP como marcador en el ratón mutante GFAPhrGFP- CB_1 -WT permitió la detección de niveles significativamente mayores de expresión del receptor CB_1 en los procesos astrocíticos hipocampales que los encontrados en los ratones tipo silvestre CB_1 -WT utilizando como marcador GFAP.
- 5- En el hipocampo, la distancia más común entre el receptor CB_1 de astrocitos y la sinapsis más cercana se encuentra en el rango de los 400-800 nm, siendo la mayoría de ellas excitadoras.
- 6- La mayoría de las sinapsis observadas en el CA1 y en la capa molecular del giro dentado rodeadas de astrocitos inmunopositivos para el receptor CB_1 en el rango de 400-800 nm son de naturaleza excitatoria.
- 7- El estudio de los ratones mutantes rescatados para el receptor CB_1 caracterizados en esta Tesis Doctoral ha demostrado que: 1) la expresión de los receptores CB_1 se produce en tipos celulares específicos; 2) la re-expresión se encuentra limitada a poblaciones neurales concretas o a astrocitos; 3) los niveles endógenos de receptores CB_1 se mantienen en los tipos celulares que re-expresan el receptor.
- 8- En conjunto, los ratones mutantes estudiados rescatos para el receptor CB_1 son herramientas excelentes para la realización de estudios funcionales y translacionales relativas al papel de los receptores CB_1 en el cerebro en condiciones normales y patológicas.

2 INTRODUCTION

2.1 THE ENDOCANNABINOID SYSTEM

The Cannabis plant (*Cannabis sativa* L.) has been used since antiquity, mostly for the manufacturing and fabrication of cordages but also by its psychotropic, analgesic and antispasmodic effects (Piomelli, 2003). Cannabis plants produce a mixture of chemical constituents, which are collectively referred as phytocannabinoids. Cannabinoids have been shown to produce a unique plethora of effects on the behavior of humans and animals which include euphoria, excitement, cognitive impairments, hypothermia, errors of time and space, enhancement of the sense of hearing, fluctuations of emotions, discoordination, illusions and hallucinations, disruption of short-term memory, relaxation, tachycardia, a decreased ability to focus attention, and sleepiness (Block et al., 1992; Chait and Perry, 1994; Heishman et al., 1997; Court, 1998; Mechoulam and Parker, 2011).

Among these compounds, delta (9) - tetrahydrocannabinol (Δ^9 -THC or THC) was first isolated in 1964 (Gaoni and Mechoulam, 1964), and was shown to account for the psychotropic effects of cannabis. Δ^9 -THC was described to couple with a group of receptors later named as the cannabinoid receptors, which consequently represent the target of a type of endogenous lipidic messengers, the endocannabinoids. Thus, the endocannabinoid system is comprised by the cannabinoid receptors, their endogenous lipid ligands (the endocannabinoids) and also the enzymes for their synthesis and degradation (Piomelli, 2003, 2014; De Petrocellis et al., 2004; Marsicano and Lutz, 2006; Kano et al., 2009; Katona and Freund, 2012; Pertwee, 2015; Lu and Mackie, 2016).

This system is involved in many brain processes ranging from food intake to cognition (Marsicano and Kuner, 2008; Bellocchio et al., 2010; Katona and Freund, 2012). More specifically, endocannabinoids participate in the complex cellular and molecular mechanisms involved in endocannabinoid-mediated synaptic plasticity (Castillo, 2012) which provide important computational properties to brain circuits, such as coincidence detection and input specificity, critical for high brain functions (Castillo et al., 2012).

2.1.1 ENDOCANNABINOID COMPOUNDS

Endocannabinoids are endogenous lipid compounds synthesized from the cellular membranes. Among them, there are two main endocannabinoids which are better identified and characterized, i.e. *N*-arachidonylethanolamine (anandamide, AEA) (Devane et al., 1992) and 2-arachidonoylglycerol (2-AG) (Mechoulam et al., 1995). AEA is known by triggering the “tetrad” effects (i.e., catalepsy, antinociception, hypolocomotion, and hypothermia) of cannabinoids in rodents (Fride and Mechoulam, 1993) whereas 2-AG plays a key role in most of the Cannabinoid Type I (CB₁) receptor mediation of synaptic transmission (Kano et al., 2009).

Endocannabinoids are synthesized and released *on demand* following physiological and pathological stimuli and, for the best of our knowledge no evidence have been found for their storage in secretory vesicles (De Petrocellis and Di Marzo, 2009a; Pertwee et al., 2010).

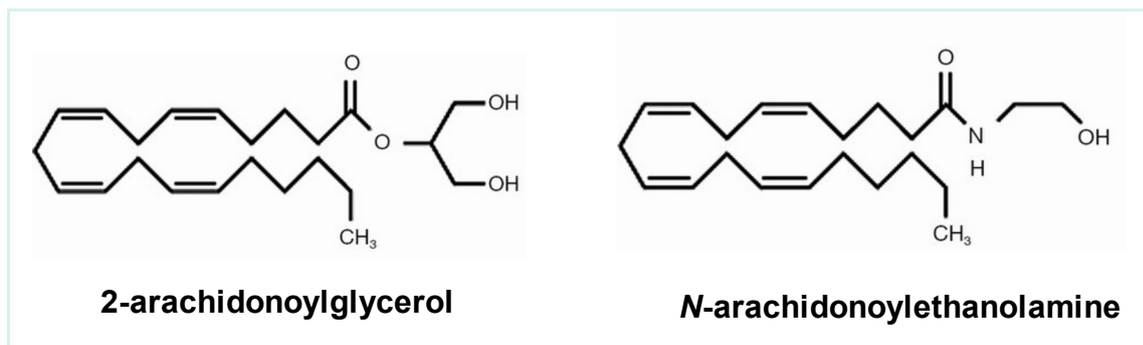


Figure 1. Chemical structures of the two major endocannabinoids. N-arachidonylethanolamide (AEA; “anandamide”) and 2-arachidonoylglycerol (2-AG).

Interestingly, other *N*-acylethanolamines such as oleoylethanolamine (OEA) (Fu et al., 2003; Wang et al., 2005; Thabuis et al., 2008) and palmitoylethanolamine (PEA) (Di Marzo et al., 2001; Costa et al., 2008) exhibit a variety of similar biological activities and share with the endocannabinoids certain metabolic pathways. However, it has not been well established if these molecules also share with endocannabinoids their molecular targets (De Petrocellis and Di Marzo, 2009a).

2.1.2 METABOLISM OF ENDOCANNABINOIDS

A wide variety of biochemical pathways for the synthesis, transport, release and degradation of endocannabinoids have been described to date (Piomelli, 2014). In this sense, neuronal membrane depolarization or the activation of G_q-coupled GPCRs triggers the synthesis of 2-AG (Hashimoto et al., 2005; Kano et al., 2009). It is thought that phospholipase C, β or δ hydrolyses membrane phospholipids to obtain diacylglycerols (DGs), and then the degradation of these DGs by diacylglycerol lipases (DGLs) is driven to the synthesis 2-AG (Gao et al., 2010; Tanimura et al., 2010).

Similarly, it has been described that the Ca²⁺-dependent enzyme *N*-acyltransferase (NAT) drives the transfer of arachidonic acid (AA) from phosphatidylcholine to phosphatidylethanolamine, consequently forming the AEA precursor *N*-arachidonoyl phosphatidylethanolamine (NAPE) (Cadas et al., 1996). Afterwards, the *N*-acyl phosphatidylethanolamine specific phospholipase D (NAPE-PLD) (Okamoto et al., 2004) catalyzes the release of AEA.

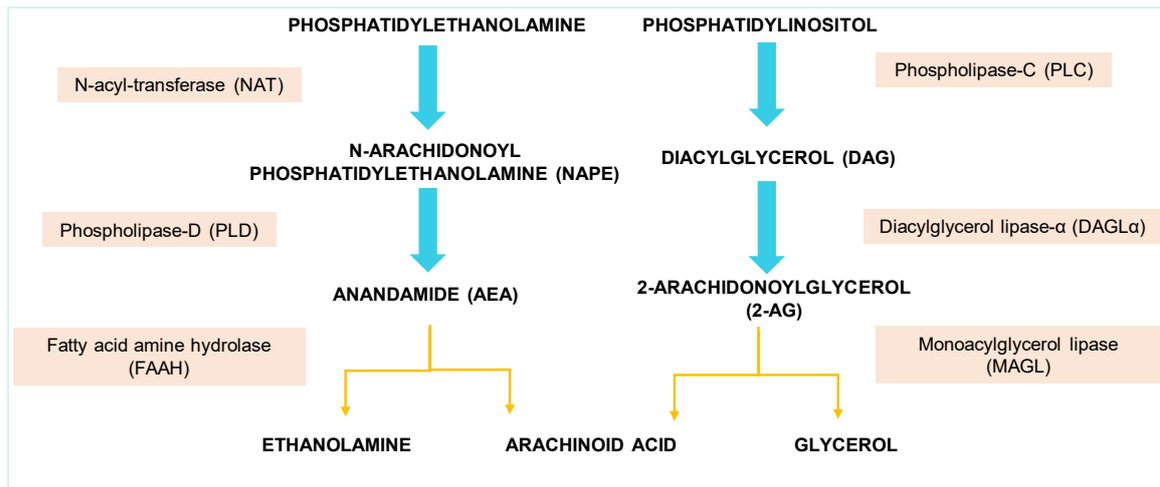


Figure 2. Major pathways of anandamide and 2-AG synthesis and degradation. Modified from Lee et al., 2015.

Nevertheless, an alternative pathway for the release of AEA includes the hydrolysis of NAPE to phosphoanandamide by the action of the Phospholipase C (PLC) followed by dephosphorylation mediated by phosphatase, such as PTPN22 (Liu et al., 2006).

Conversely, monoacylglycerol lipase (MAGL) and fatty acid amide hydrolase (FAAH) represent the two main catabolic enzymes for the hydrolysis of 2-AG and AEA respectively (Dinh et al., 2002; Ueda, 2002). In this sense, MAGL is mainly found on presynaptic terminals, where it hydrolyses 2-AG molecules bound to presynaptic CB₁ receptors (Straiker et al., 2009; Kano et al., 2009). Furthermore, two other hydrolases, ABHD6 and ABHD12 (α/β -hydrolase domain containing 6 and 12) are also capable of hydrolyzing 2-AG (Blankman et al., 2007). On the other hand, FAAH showing overlap with CB₁ in many brain regions (Egertová et al., 2003; Kano et al., 2009). Moreover, FAAH has also been described to be involved in the metabolism of other bioactive N-acylethanolamines such as OEA and PEA (Di Marzo et al., 2001).

2.1.3 CANNABINOID RECEPTORS

Endocannabinoids receptors are known to be present in many vertebrate species, including rodents and primates, both monkeys and humans, (Elphick and Egertová, 2005). To date, two cannabinoid receptors have been cloned, i.e. the Type I Cannabinoid receptor (CB₁ receptor) (Matsuda et al., 1993) and the Type II Cannabinoid receptor (CB₂ receptor) (Munro et al., 1993).

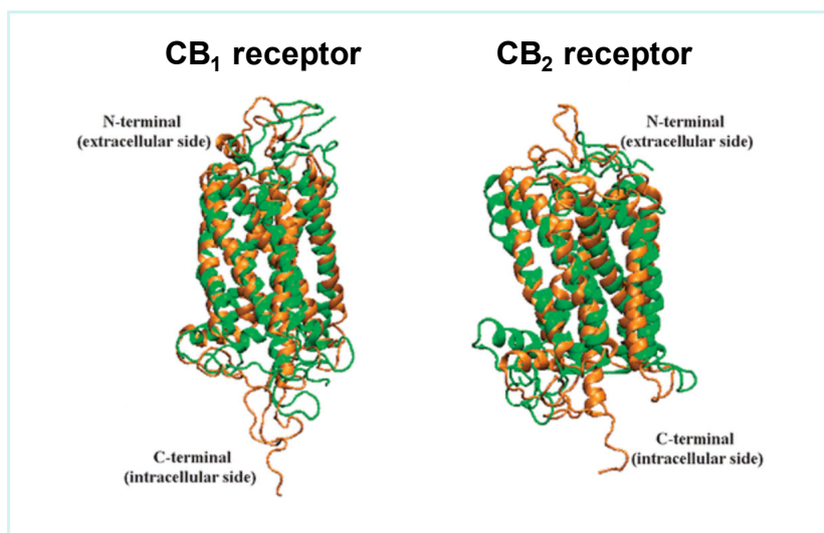


Figure 3. Three-dimensional representation of CB₁ receptor and CB₂ receptor structure. Modified from Ramos et al., 2011.

The CB₁ and CB₂ cannabinoid receptors belonging to the G-protein-coupled receptors (GPCRs) superfamily, the largest family of trans-membrane proteins in the human genome, and crucial for many essential physiological processes. GPCRs are characterized by seven hydrophobic transmembrane segments connected by intracellular and extracellular loops, an N-terminal extracellular domain that possesses glycosylation sites and a C-terminal intracellular domain coupled to a G_{i/o} protein (Howlett et al., 2002).

In the cell, the activation of CB₁ or CB₂ receptor by endocannabinoids triggers intracellular signaling events coupled to these two G-protein-coupled receptors (GPCRs) (Pertwee, 1997). Firstly, stimulation of CB₁ or CB₂ receptor leads to the inhibition of stimulus-induced adenylate cyclase (AC) and, consequently, modulates protein kinase-A-dependent cascades. Moreover, the activation of these receptors provokes the stimulation of mitogen-activated protein kinase signaling. More specifically, the activation of CB₁ receptor also exerts the modulation of ion channels, inducing inhibition of P, Q and N-type voltage-sensitive Ca²⁺ channels and stimulation of rectifying G-protein-coupled K⁺ channels. Also, just in the case of CB₁ receptors, the stimulation of this receptor induces the activation of PLC- γ , by the $\beta\gamma$ subunits of G_{i/o} proteins stimulates phosphatidylinositol 3-kinase and intracellular Ca²⁺ mobilization (Devane et al., 1988; Howlett et al., 1990, 2002; De Petrocellis et al., 2004; Pertwee, 2015).

CB₁ receptor is predominantly expressed in the central nervous system (CNS) particularly in brain cortex, basal ganglia, hippocampus, and cerebellum (Hu and Mackie, 2015). Besides, minor levels of expression of CB₁ receptors have been also document in various peripheral tissues (Pertwee, 2000, 2001). On the contrary, CB₂ receptor are localized primarily in immune cells and tissues derived from the immune system (Ameri, 1999; Cabral et al., 2015).

Although CB₁ and CB₂ are well established as endocannabinoid receptors, there exist other receptors that could be mediating the effects of endocannabinoids (Pertwee, 2015), such as the Transient Receptor Potential Vanilloid 1, also known as TRPV1, wich can be activated by anandamide (Maccarrone et al., 2008; De Petrocellis and Di Marzo, 2009b; Tóth et al., 2009; Alhouayek et al., 2014; Rossi et al., 2015), as well as for Transient Receptor Potential Ankyrin 1 (TRPA1) receptors (De Petrocellis et al., 2008), Peroxisome Proliferator-Activated

Receptors, namely PPAR- α (Sun et al., 2006; Alhouayek et al., 2014) and non-CB₁/CB₂ GPCRs such as G protein-coupled receptor 55 (GPR55) (Ryberg et al., 2007).

2.1.4 PHYSIOLOGICAL ACTIONS OF ENDOCANNABINOIDS

Endocannabinoids are released from the postsynaptic neurons after its depolarization and/or receptor activation. Once released, endocannabinoids bind to the presynaptically located CB₁ (Robbe et al., 2002; Chevaleyre and Castillo, 2003; Kano, 2014). Thus, the binding of the endocannabinoid to its receptor, is able to suppress the elevation of the intracellular free Ca²⁺ concentration induced by the depolarization of the postsynaptic neuron (Sugiura et al., 1997) and consequently regulate the release of different neurotransmitters such as: glutamate (Shen et al., 1996), and gamma-Aminobutyric acid (GABA) (Katona et al., 1999), acetylcholine (Gifford et al., 1997a; b) or noradrenaline (Schlicker et al., 1997). The transient suppression of transmitter release on the presynaptic neuron following the release of endocannabinoid from postsynaptic neurons is known as endocannabinoid-mediated short-term depression or eCB-STD. The transient suppression has been consistently demonstrated to mediate both, depolarization-induced suppression of inhibition (DSI) (Ohno-Shosaku et al., 2001; Wilson and Nicoll, 2001) and of excitation (DSE) (Kreitzer and Regehr, 2001), mechanisms by which short-term depression of GABA or glutamate release from axon terminals is induced by the depolarization of a postsynaptic neuron (Katona and Freund, 2012; Kano, 2014).

On the other hand, long-term depression (LTD) of GABA or glutamate release is mediated by CB₁ receptor in response to long-lasting synaptic activity, which triggers postsynaptic endocannabinoid releasement and retrograde signaling (endocannabinoid-mediated long-term depression or eCB-LTD) in different types of synapses throughout nervous system (Chevaleyre and Castillo, 2003; Chevaleyre et al., 2006; Castillo, 2012).

In the past years, endocannabinoid signaling have been demonstrated to play a key role in the potentiation of glutamatergic transmission through neuron-glia communication, also known as tripartite synapses, that includes the activation of astroglial CB₁ receptors (Navarrete and Araque, 2008, 2010; Navarrete et al., 2013; Metna-Laurent and Marsicano, 2015; Oliveira da Cruz et al., 2016).

In general, most of the actions of (endo)cannabinoids are mediated by CB₁ receptor ability to modulate synaptic transmission (Kano et al., 2009; Castillo, 2012; Katona and Freund, 2012; Kano, 2014). More specifically, the regulation of synaptic plasticity mediated by CB₁ receptor has been related to the memory-impairment effects of cannabinoids by means of the induction of LTD and the inhibition of LTP in the hippocampus (Stella et al., 1997; Han et al., 2012).

In addition, endocannabinoids have been described to also be involved in cell survival and differentiation processes (Galve-Roperh et al., 2013). In this sense, during embryony development, the activation of CB₁ and CB₂ receptors participate in the regulation of neuronal progenitor-stem cells self-renewal, proliferation and differentiation. Particularly, the activation of these receptors have been reported to trigger many different signaling pathways, such as mitogen-activated protein kinase signaling pathways, focal adhesion kinase cascade, phosphatidylinositol 3-kinase, extracellular-signaling regulated protein kinase, and

ceramide signaling, among others (Bouaboula et al., 1995b; Derkinderen et al., 1996; Aguado et al., 2005; Ortega-Gutiérrez et al., 2005; Galve-Roperh et al., 2013).

Endocannabinoids are also able to develop neuroprotective roles acting through CB₁ cannabinoid receptors. In this sense, it has been shown that the role developed by the eCBs in retrograde signaling system in glutamatergic synapses (Ohno-Shosaku and Kano, 2014) in the brain could be the origin of its protective role (Wallace et al., 2003; Kano et al., 2009). Importantly, the eCB system would be a pharmacological target for the treatment of diseases characterized by abnormal glutamate homeostasis, such as neurodegenerative disorders (Fernández-Ruiz, 2010; Fernández-Ruiz et al., 2015).

2.1.5 EXPRESSION OF CB₁ RECEPTOR IN THE BRAIN

The CB₁ receptor is the main endocannabinoid signaling protein regarding the CNS (Pertwee, 2009), and, more specifically, the CB₁ receptors represent one of the most abundant GPCRs in the brain (Herkenham et al., 1991; Tsou et al., 1998; Moldrich and Wenger, 2000). Its expression is widespread and heterogeneous, developing crucial roles in brain function, dysfunction and cognition (Marsicano et al., 2002; Monory et al., 2006; Bellocchio et al., 2010; Puente et al., 2011; Castillo, 2012; Katona and Freund, 2012; Steindel et al., 2013; Ruehle et al., 2013; Soria-Gómez et al., 2014, 2015; Hu and Mackie, 2015; Katona, 2015; Martín-García et al., 2015), and it has been reported to palliate numerous pathologies, including Alzheimer's disease, pain, obesity, and cancer, (Pertwee, 2009). More specifically, the highest levels of CB₁ receptors are present in the substantia nigra, globus pallidus, cerebellum, hippocampus and cortex (Howlett et al., 1990).

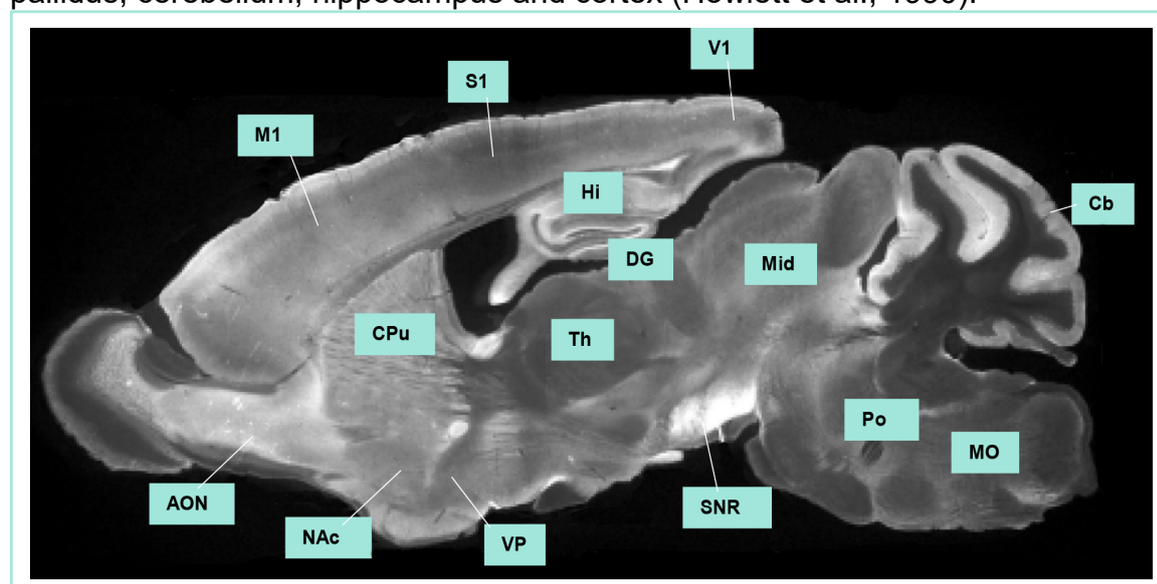


Figure 4. Distribution of CB₁ receptors in the central nervous system of adult mice. Primary somatosensory cortex (S1), primary motor cortex (M1), primary visual cortex (V1), hippocampus (Hi), dentate gyrus (DG), cerebellar cortex (Cb), midbrain (Mid), caudate putamen (CPu), thalamus (Th), anterior olfactory nucleus (AON), nucleus accumbens (NAc), ventral pallidum (VP), substantia nigra pars reticulata (SNR), pons (Po), medulla oblongata (MO). Modified from Kano et al., 2009.

At the ultrastructural level, CB₁ receptor expression is very high in inhibitory GABAergic synaptic terminals (Katona et al., 1999; Kawamura et al., 2006; Ludányi

et al., 2008; Marsicano and Kuner, 2008; Katona and Freund, 2012; De-May and Ali, 2013; Steindel et al., 2013; Hu and Mackie, 2015) but, to a lower extent, CB₁ receptors are also present in glutamatergic neurons (Marsicano et al., 2003; Domenici et al., 2006; Takahashi and Castillo, 2006; Katona et al., 2006; Monory et al., 2006; Kamprath et al., 2009; Bellocchio et al., 2010; Puente et al., 2011; Reguero et al., 2011; Ruehle et al., 2013; Soria-Gómez et al., 2014) and in brain astrocytes (Navarrete and Araque, 2008, 2010; Stella, 2010; Han et al., 2012; Bosier et al., 2013; Metna-Laurent and Marsicano, 2015; Viader et al., 2015; Oliveira da Cruz et al., 2016). CB₁ receptors are also constitutively expressed by oligodendrocytes (Molina-Holgado et al., 2002a; Benito et al., 2007; Garcia-Ovejero et al., 2009; Mato et al., 2009; Gomez et al., 2010). Inside the cells, CB₁ receptors, can be subcellularly localized at plasma membranes and intracellular organelles, such as brain mitochondria (Bénard et al., 2012; Hebert-Chatelain et al., 2014a; b; Koch et al., 2015).

Because of the hugely different levels of expression among different cell types (Marsicano and Lutz, 1999), it is extremely difficult to identify low CB₁ receptor expression in cell types and/or in subcellular compartments of wild-type brains (Busquets-Garcia et al., 2015). Actually, former anatomical studies often misinterpreted low CB₁ receptor expression as background staining (Freund et al., 2003).

Pharmacological blockade or genetic ablation of the receptor and its eventual behavioral and neuronal consequences are clear indications of the role of endocannabinoids in the brain. Hence, based on methodological “loss of function” approaches, the use of mutant mice lacking CB₁ receptors in specific cell populations helped to identify the necessary role of the receptor in different brain functions (Marsicano et al., 2003; Monory et al., 2006, 2015; Martín-García et al., 2015; Zimmer, 2015). Moreover, neuroanatomical analysis of these mutant animal models determined the presence of the CB₁ receptor at sites previously not known (e.g. cortical glutamatergic neurons or neuronal mitochondria).

However, these approaches do not provide definitive information concerning the possibly sufficient role of the cell-type expression of CB₁ receptors for a given function. Indeed, distinguishing between low levels (wild-type) and no-expression (mutant mice lacking CB₁ receptors in specific cell populations) is very difficult. To establish precise causal relationships and to identify low levels of expression, “rescue” strategies utilizing the endogenous genetic locus are needed in order to analyze the function of endogenous CB₁ receptor expression exclusively in specific cell types. To reach this aim the Cre/LoxP system have been previously applied to restore CB₁ receptor expression in a CB₁ receptor-null mutant only in specific cell types (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015). In this sense, the rescue approach was first applied to the hypothalamic expression of the melanocortin 4 receptor (Balthasar et al., 2005) using the Cre/loxP system, and was recently applied to the CB₁ gene (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015). Briefly, a “stop cassette” flanked by LoxP sites was introduced immediately upstream of the coding sequence of the CB₁ gene by homologous recombination to generate a mutant mouse line. The mice obtained by this approach bore no expression of the CB₁ receptor protein (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015), making these mice phenotypically very similar to conventional CB₁ knock-out mice (CB₁-KO). However, the crossing of these mice with Cre-

expressing transgenic mouse lines or the local injection of viruses expressing the Cre recombinase induces a Cre-mediated deletion of the “stop cassette”, causing a specific rescue of expression of the protein in specific cell populations and/or brain regions (Balthasar et al., 2005; Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015).

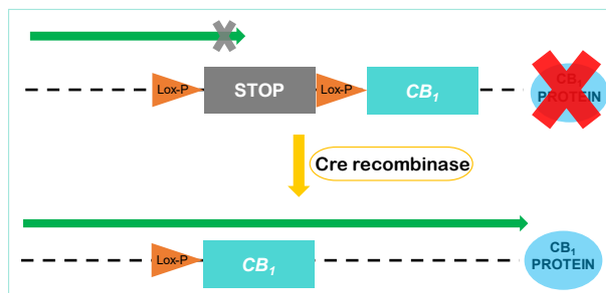


Figure 5. Schematic representation of rescued CB₁ receptor allele after excision of the stop cassette mediated by Cre recombinase.

Moreover, this re-expression is limited to the cell types where the regulatory sequences of the CB₁ receptor are normally activated and the expression levels are the same as in wild-type animals (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015).

Furthermore, these rescue mice allowed determining the sufficient causal role of the CB₁ receptor in the anterior olfactory cortex area as well as in dorsal telencephalic glutamatergic neurons (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015). However, no studies have addressed so far the subcellular localization of CB₁ receptor in these mutant brains at the electron microscopic level. This is particularly interesting due to the huge expression of CB₁ receptors in specific cell types (e.g. hippocampal GABAergic interneurons) which hinder to appreciate the detailed pattern of expression of the CB₁ receptor in other cell types (e.g. hippocampal glutamatergic pyramidal neurons) in wild-type mouse samples (Freund et al., 2003). Therefore, the conditional rescue mutants emerge as key tools for the detailed anatomical characterization of the subcellular distribution of the receptor in specific cell types, independently of its level of expression.

2.1.6 ASTROCYTES AND THE CB₁ RECEPTOR

Presence of CB₁ receptors in astrocytes has been long debated. Its presence in astrocytes was firstly reported in human and rodent cell cultures (Bouaboula et al., 1995a; Sánchez et al., 1998; Molina-Holgado et al., 2002b; Sheng et al., 2005). However, incongruous results were obtained in mice (Sagan et al., 1999; Walter and Stella, 2003) suggesting that CB₁ receptor expression in astrocytes could vary among different animal species, strains or different cell culture conditions. In addition, low levels of expression of astrocytic CB₁ receptor made it remarkably arduous to detect astroglial CB₁ mRNA or protein expression by means of in situ hybridization or immunohistochemistry. Fortunately, electron microscopy immunodetection allowed to reveal the presence of astrocytic CB₁ receptor (Rodríguez et al., 2001; Han et al., 2012; Bosier et al., 2013). Moreover, the enzymes for synthesis and degradation of the main endocannabinoids co-localize with Glial Fibrillary Acidic Protein (GFAP) (Suárez et al., 2010). Finally, in light of

many different works published during the last years (Navarrete and Araque, 2008, 2010; Stella, 2010; Han et al., 2012; Bosier et al., 2013; Gómez-Gonzalo et al., 2014; Navarrete et al., 2014; Metna-Laurent and Marsicano, 2015; Viader et al., 2015; Oliveira da Cruz et al., 2016) the presence of the CB₁ receptor in astrocytes and its functional importance have been demonstrated.

In this sense, different authors have proposed various signaling pathways in astrocytes after the CB₁ receptor activation. Thus, some authors have proposed the coupling of astrocytic CB₁ to Gα_{i/o}-protein (Guzmán et al., 2001) whereas other authors suggest that astrocytic CB₁ receptor is able to modulate Ca²⁺ levels in astrocytes and associated signaling by means of Gα_q proteins activation (Navarrete and Araque, 2008; Navarrete et al., 2014). In this sense, the development of different intracellular astrocytic signaling pathways in response to endocannabinoids provides to endocannabinoid-mediated responses with a particular important flexibility in astrocytes (Metna-Laurent and Marsicano, 2015).

Otherwise, astrocytes have long been known for their role as protective elements against injury in the central nervous system and as energetics providers for neurons (Magistretti and Pellerin, 1999; Bélanger and Magistretti, 2009; Yi et al., 2011), not taking a relevant part in the transmission of information for being non-electrically excitable cells. However, this last consideration has already changed, owing to the fact that many signaling systems that are capable to modulate neuronal activity have been described to also regulate astrocytic-neuronal communication (Araque et al., 1999; Haydon and Carmignoto, 2006; Perea et al., 2009; Gómez-Gonzalo et al., 2014). Moreover, these findings lead to the establishment of a new concept: the “tripartite synapse”, which is formed by a presynaptic and a postsynaptic neuronal elements and by the surrounding astroglial processes (Araque et al., 1999) (Fig. 6). In this context, astrocytic CB₁ receptors have been demonstrated to play a key role in the communication between neurons and astrocytes (Navarrete and Araque, 2008, 2010; Navarrete et al., 2013, 2014; Gómez-Gonzalo et al., 2014).

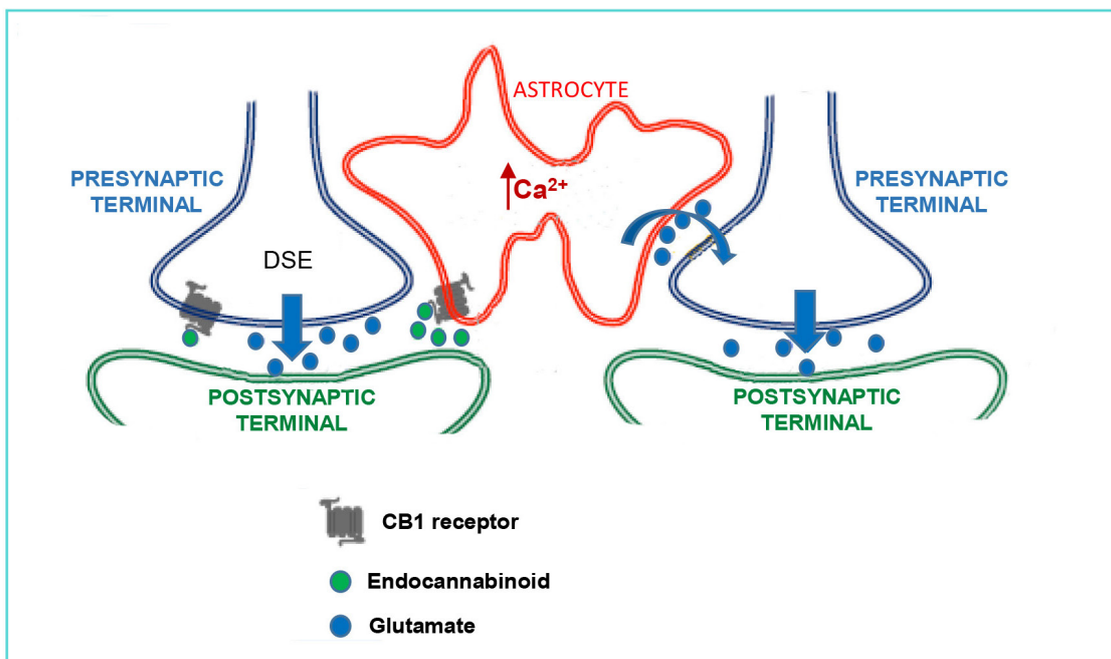


Figure 6. Signaling mechanisms underlying synaptic regulation by endocannabinoids through stimulation of astrocytes. Endocannabinoids are released by a postsynaptic neuron acting on presynaptic CB_1 receptors. As a response, a DSE is triggered in homoneuronal synapses. In addition, the released endocannabinoids also are able to activate astrocytic CB_1 receptors. The activation of astrocytic CB_1 receptors increases the intracellular calcium and triggers the release of glutamate by the astrocyte, which stimulates the release of neurotransmitter in hetero-neuronal synapses. Modified from Navarrete et al., 2014.

Moreover, glutamate release is promoted in neurons by the activation of CB_1 receptors in astrocytes (Navarrete and Araque, 2008), and therefore this activation could be improving synaptic transmission (Navarrete and Araque, 2010). Thus, in this context, it has been suggested that the CB_1 receptor would mediate in the development of epileptic seizures (Coiret et al., 2012). In this sense, it has been recently demonstrated that the CB_1 receptor expression in astrocytes is increased in sclerotic hippocampus, which is characterized by refractory temporal lobe epilepsy and both neuronal loss and astrocyte proliferation (Meng et al., 2014).

In astrocytes, GFAP is the principal intermediate filament protein (Eng, 1985; Eng et al., 2000; Hol and Pekny, 2015). However, due to its filament nature the immunolabeling of GFAP may not show the complete surface of the astrocyte in traditional microscopic preparations. Thus, other methodological approaches have been developed for staining astrocytes in microscopic preparations. In this sense, Green Fluorescence Protein, also known as GFP (Shimomura et al., 1962; Chalfie, 1995), is a cytoplasmically expressed diffusible protein. When inserted in astrocytes, this protein appears distributed throughout the whole cytoplasm, allowing its immunolabeling to be used for a better detection of the whole astrocytic surface (Nolte et al., 2001). More particularly, GFP protein from the sea pansy *Renilla reniformis* (Ward et al., 1978; Ward and Cormier, 1979) and its humanized isoform, known as hrGFP, has been used as a marker in a wide variety of studies (Navarro-Galve et al., 2005; De Francesco et al., 2015; Kerr et al., 2015). More specifically, it has been used as an astroglial marker (Hadaczek et al., 2009).

2.2 HIPPOCAMPUS

2.2.1 DEFINITION OF THE HIPPOCAMPUS

The hippocampus is a structure that includes distinct brain regions differentiated by its cytoarchitecture. The main reason to include these areas under the same anatomical structure is that they are largely connected via unidirectional projections (the perforant path, mossy fibers and Schaffer collaterals). However, as I will describe later, two-way connections may also exist in this region.

The hippocampus can be divided in two cortical structures i.e. the hippocampal formation and the parahippocampal region (Fig. 7). On the one hand, the hippocampal formation is a C-shaped structure localized in the caudal part of the brain. It is comprised by three areas cytoarchitecturally differentiated: the dentate gyrus, the hippocampus properly speaking which is subdivided into three fields (CA1, CA2 and CA3), and the subiculum. On the other hand, the parahippocampal region lies adjacent to the hippocampal formation and comprises five subregions; the perirhinal cortex, the entorhinal cortex, the postrhinal cortex, the presubiculum and the parasubiculum.

The three-dimensional location of the hippocampus in the mouse brain is complex. It appears as an elongated structure, with its septo-temporal C-shaped axis extending towards the temporal lobe in its caudo-ventral end running above and behind the diencephalon. It is delimited by the septal nuclei rostrally and by the lateral ventricles laterally. The longitudinal axis of the hippocampal formation is located in septum-temporal direction, while the transverse or orthogonal axis is extending between temporal lobes. The regions of the hippocampal formation are distributed along the septum-temporal axis so that the dentate gyrus and CA1 and CA3 regions of the hippocampus appear at more septal levels. The subiculum begins to appear in the first third of septotemporal axis, while the presubiculum and parasubiculum are appearing progressively towards the temporal region. Finally, ventrally in the caudal portion of the hippocampus the entorhinal cortex is confined.

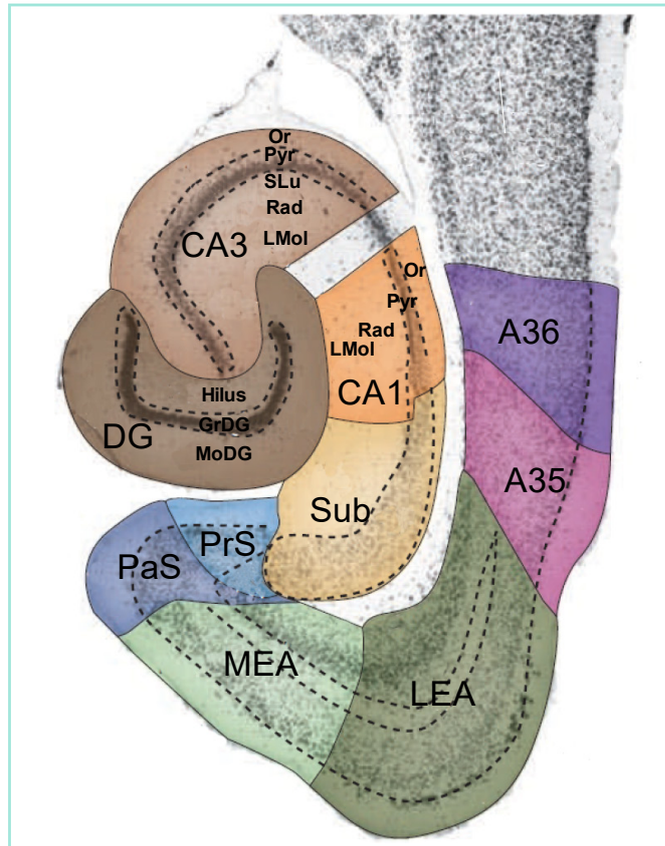


Figure 7. Representation of the hippocampal formation and the parahippocampal region in the rat brain. Dentate gyrus, DG (dark brown); Region 3 of Cornu Ammonis, CA3 (medium brown); Region 2 of Cornu Ammonis CA2 (not indicated); Region 1 of Cornu Ammonis, CA1 (orange). The subiculum, Sub (yellow); presubiculum, PrS (medium blue) and parasubiculum (PaS; dark blue). The entorhinal cortex, which has a lateral, LEA (dark green) and a medial, MEA (light green) areas. The perirhinal cortex (consisting of Brodmann areas: A35 (pink) and A36 (purple). Stratum oriens, Or; Pyramidal cell hippocampus, Py; Stratum radiatum, Rad; stratum lucidum, SLu; Lacunosum moleculare, LMol; Molecular dentate gyrus, MoDG and Granular dentate gyrus, GrDG. Modified from van Strien et al., 2009.

2.2.2 DENTATE GYRUS

The dentate gyrus comprises three layers, the molecular layer, the granular cell layer and the hilus.

2.2.2.1 Molecular layer.

The molecular layer is located edging the hippocampal fissure and contains the perforant path fibers connecting the entorhinal cortex with the dentate gyrus. This layer is mainly occupied by the dendrites of the granule, basket and various polymorphic cells, as well as axon terminals from several sources. Remarkably, synaptic terminals of the commissural/associational system are found in the inner third of the dentate molecular layer, containing a substantial input from the CA3 in addition to the excitatory terminals of the hilar mossy cells (Ribak et al., 1985; Li et al., 1994; Blasco-Ibáñez and Freund, 1997; Scharfman, 2007).

Furthermore, a few neuronal cell types are also present in the molecular layer. The first type of neurons is located in the innermost area, shows a triangular or

multipolar soma and present axons projecting to the granular layer basket plexus cells. They are considered as a type of basket cells (Hazlett and Farkas, 1978; Ribak and Seress, 1983; Witter and Amaral, 2004; Jinno and Kosaka, 2006; Witter, 2012).

The other kind of neurons that are localized in the molecular layer are immunoreactive for the neurotransmitter GABA (Celio, 1990), which indicates that they are modulating the activity of granule cells. They are similar to chandelier cells (Kosaka, 1983; Somogyi et al., 1985; Soriano and Frotscher, 1989; Halasy and Somogyi, 1993; Witter and Amaral, 2004; Witter, 2012). Their dendrites remain primarily in the molecular layer, while their axons project from the molecular layer into the granular cell layer and contact with initial axonal segments of granule cells.

2.2.2.2 Granular cell layer

The granule cell layer is situated deeply into the molecular layer. It is constituted mainly of densely packed excitatory glutamatergic neurons known as granule cells. Their dendritic extensions give the cell a characteristic conical appearance and are oriented towards the molecular layer. Furthermore, they have numerous dendritic spines which are able to contact with many axon terminals (Desmond and Levy, 1982, 1985) and increase in length at more distal positions along the dendrite (Stanfield and Cowan, 1979; Vuksic et al., 2008). On the other hand, their glutamatergic axons known as mossy fibers invade the hilar region where they contact with the mossy cells, CA3 pyramidal neurons and GABAergic interneurons (Frotscher et al., 1994). No remarkable morphological differences appeared between granule cells taken from the dorsal or ventral portions, except for the more extensive degree of arborisation found in ventral cells (Vuksic et al., 2008).

Another type of cells in this layer are the GABAergic basket cells (Aika et al., 1994) whose somata are located in the innermost side of the granular cell layer. They owe their name to their axons which form a pericellular plexus surrounding the granule cells bodies (Ribak and Seress, 1983; Ribak et al., 1985) modulating granule cells activity (Meyer et al., 2002; Jinno and Kosaka, 2006).

2.2.2.3 Hilus

Also known as the polymorphic layer, the hilus contains a wide variety of cell types. The most common are the glutamatergic mossy cells (Frotscher et al., 1994; Blasco-Ibáñez and Freund, 1997). Their large cell bodies are oval and their dendrites extend for long distances within the hilus, sometimes being able to reach out the granule cell layer and the molecular layer (Blasco-Ibáñez and Freund, 1997; Fujise and Kosaka, 1999). Remarkably, dendrites of the mossy cells do not leave the confines of the hilus to enter the adjacent CA3 field. The most proximal dendrites are covered by the very large and complex spines called “thorny excrescences”, which establish synaptic contacts with the granule mossy fibers (Ribak et al., 1985; Frotscher et al., 1991; Blasco-Ibáñez and Freund, 1997; Fujise and Kosaka, 1999). Importantly, the axon terminals of the hilar mossy cells contain abundant tightly packed synaptic vesicles and end into the inner third of dentate molecular layer forming asymmetric synapses with granule cell dendrites and dendritic spines (Ribak et al., 1985; Blasco-Ibáñez and Freund, 1997).

Other cell types located in the hilus are the fusiform cells whose dendrites run parallel to the granular cell layer (Bakst et al., 1986; Halasy and Somogyi, 1993), and the multipolar neurons whose axons and dendrites remain within the hilus (Amaral, 1978).

2.2.3 HIPPOCAMPUS

As Lorente de Nó proposed (1934) the hippocampus can be divided into three regions, CA1, CA2 and CA3. The mossy fibers inputs that arrive to CA3 from the dentate gyrus not reaching CA1 and CA2 represent the main anatomical difference among these three regions. In addition to the structural characteristics, different patterns of gene expression also exist among the different regions (Dong et al., 2009; Fanselow and Dong, 2010) that make a real physiological separation between the three different fields.

The hippocampus shows a trilaminar structure, folded on itself. Each of these layers can be defined depending on the cell types that host and their spatial distribution and density. Remarkably, the cellular distribution is correlated with the performance of specific functions in the hippocampal network (Klausberger and Somogyi, 2008). The ventricular surface of the hippocampus is covered by the alveus, which is a thin layer of myelinated fibers.

2.2.3.1 Stratum oriens

The basal dendritic tree of pyramidal neurons will be part of the stratum oriens, which is a relatively free layer of cells situated in the outer side of the pyramidal layer (Lorente De Nó, 1934).

2.2.3.2 Pyramidal cell layer

The pyramidal cell layer (Lorente De Nó, 1934), consists of glutamatergic cells (Liu et al., 1989) whose somata have a pyramidal look due to the presence of a prominent apical dendrite. Their basal dendritic trees distribute in the stratum oriens while the apical dendritic tree is oriented in ventromedial direction.

There is also an assorted population of basket cells with their cell bodies confined within the pyramidal cell layer (Frotscher et al., 1991; Jinno and Kosaka, 2006). Their axons expand transversely from the cell body forming a basket plexus that innervates the pyramidal cell bodies.

2.2.3.3 Stratum lucidum

This is an acellular CA3 layer just above the pyramidal cell layer that contains the dentate granule mossy fibers axons.

2.2.3.4 Stratum radiatum

The stratum radiatum is located superficial to stratum lucidum in CA3 and above the pyramidal cell layer in CA2. The majority of the CA3 to CA3 associational fibers and the ipsilateral CA3 to CA1 Schaffer collaterals are found in this layer.

2.2.3.5 Stratum lacunosum-moleculare

Finally, the stratum lacunosum-moleculare is located in the vicinity of the hippocampal fissure and the dentate gyrus. About 20% of the dendritic tree of the pyramidal cells reaches the stratum lacunosum-moleculare (Amaral et al., 1990). This layer receives synapses from the entorhinal cortex.

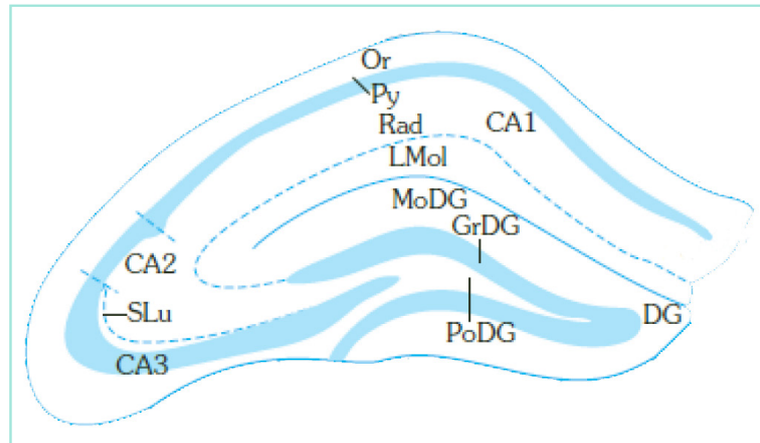


Figure 8. Representation of a sagittal section of the mouse hippocampal formation. Laminar structure of the hippocampus. DG, Dentate gyrus; CA1, Region 1 of Cornu Ammonis; CA2, Region 2 of Cornu Ammonis; CA3, Region 3 of Cornu Ammonis; Or, Stratum oriens; Py, Stratum pyramidale; Rad, Stratum radiatum; SLu, stratum lucidum; LMol, Stratum lacunosum moleculare; MoDG, Molecular dentate gyrus; GrDG, Granular dentate gyrus; PoDG, Polymorph dentate gyrus. Modified from Franklin and Paxinos, 2008.

2.2.3.6 GABAergic cells of the hippocampus

Although there are a wide variety of cells in the stratum oriens, stratum radiatum, and stratum lacunosum-moleculare of the hippocampus, most of them are GABAergic neurons (Ribak et al., 1978; Aika et al., 1994). Moreover, there are eight well defined GABAergic subclasses in the hippocampus based on their colocalization with parvalbumin, calretinin, calbindin, neuropeptide-Y, somatostatin, cholecystokinin, vasoactive intestinal polypeptide (VIP), or nitric oxide synthase (NOS) (Morrison et al., 1983; Somogyi et al., 1984; Sloviter and Nilaver, 1987; Sloviter, 1989; Gulyás et al., 1991, 1992, 1999; Miettinen et al., 1992; Tóth and Freund, 1992; Jinno and Kosaka, 2000). Some studies have shown the relation between chemical phenotypes and electrophysiological properties of the interneurons (Jinno and Kosaka, 2006; Klausberger and Somogyi, 2008) but a clear correlation between them have not been fully established yet and their function in the hippocampal network is still poorly understood (Witter and Amaral, 2004; Witter, 2012)

2.2.4 CONNECTIONS OF THE HIPPOCAMPUS

The hippocampal formation is one of the most important functional structures of the mammalian brain. There exists high connectivity with other brain regions to perform its functions properly. These connections allow playing a key role in memory formation, learning, neurogenesis and cognitive function. Also, they keep informed the hippocampal formation of the brain activity. The connections of the hippocampal formation can be divided into two groups: afferent (connections from other regions) and efferent (connections to other regions) projections.

2.2.4.1 Afferent connections

The hippocampal formation receives main connections from: the neocortex, the septal area, the contralateral hippocampus (commissural fibers), several nuclei of the diencephalon and the reticular formation of the brain stem. The neocortex of the temporal lobe which receives association fibers of different sensory association areas, sends information to the entorhinal cortex of the hippocampal formation. The entorhinal cortex is a transition structure between the neocortex and hippocampal cortex. Then, the entorhinal cortex sends fibers through the subiculum and intersects the hippocampal sulcus to end in the dentate gyrus forming the so-called perforant pathway. Fewer fibers from the entorhinal cortex form the alveolar via crossing the subcortical white matter as well as the alveus and ending in the hippocampus.

In addition, fibers from the septal nuclei reach the hippocampus through the fornix. Moreover, other afferent fibers also included in the fornix come from the thalamic nuclei, hypothalamus, diencephalon and brainstem regions such as the ventral tegmental area (dopaminergic), the locus coeruleus (noradrenergic) and nuclei raphe (serotonergic).

2.2.4.2 Efferent connections

The main efferent pathway from the hippocampal formation is the fornix, which is composed by the axons of the CA1 pyramidal neurons and of the subicular cells. The axons enter the alveus and together constitute the hippocampal fimbria followed by the fornix after leaving the fibers the hippocampus and going into the forebrain.

These glutamatergic fibers travel along the entire fornix which branches near the anterior commissure into two components: precommissural and postcommissural fibers. The precommissural fibers (formed by fibers originating mainly in the hippocampus) extend rostrally, innervating the septal nuclei, the nucleus accumbens and the substantia innominata. Meanwhile, the postcommissural fibers (composed mainly of fibers originating in the subiculum) extend caudally to the diencephalon to finally reach the hypothalamus, where they divide into the medial cortico-hypothalamic tract (that innervates the mammillary bodies, the posterior hypothalamus and other ventromedial hypothalamic nuclei) and the subiculum-thalamic tract (reaching the thalamic nuclei) (Swanson et al., 1987).

A lot of the fimbria fibers are crossed, before being included into the fornix, in the caudal region of the septal nuclei. Most of these fibers are directed to the contralateral hippocampal formation. However, a few of them are integrated into the fornix to innervate the same structures that receive precommissural and postcommissural projections from the ipsilateral fornix.

The fornix is not the only efferent pathway from the hippocampal formation, since there are significant projections from the subicular complex and entorhinal cortex to cortical areas, such as the perirhinal cortex and the parahippocampal gyrus.

2.2.4.3 Hippocampal synaptic circuit

It is well characterized that the hippocampus has a unidirectional pathway that connects all its regions sequentially. The start of the hippocampal circuit is the dentate gyrus which receives the perforant pathway from the entorhinal cortex. Then, the mossy axon fibers of the dentate granule cells project unidirectionally to the proximal dendrites of the CA3 pyramidal cells. In turn, the CA3 pyramidal cell Schaffer collaterals reach the CA1 stratum radiatum. The CA1 hippocampal region projects intensely into the subicular complex which, sends axons back to the entorhinal cortex. Although the circuit forms a loop, each of the hippocampal fields also sends axons out.

However, even though the classical view proposes that the connectivity is unidirectional, several types of retrograde connections also exist in the hippocampal formation (van Strien et al., 2009). For instance, CA3 pyramidal cells are connected with the hilus and the dentate molecular layer (Laurberg, 1979; Buckmaster et al., 1993; Witter, 2007). Furthermore, retrograde connections from CA1 to CA3 through interneurons (Laurberg, 1979; Amaral et al., 1991; Cenquizca and Swanson, 2007) and from the subiculum to CA1 (Finch et al., 1983) have been identified. Also, recurrent collaterals in CA3 (Laurberg, 1979; Amaral et al., 1990; Buckmaster et al., 1993; Sik et al., 1993; Siddiqui and Joseph, 2005; Wittner et al., 2007) and bidirectional connections between the entorhinal cortex and subiculum have been described.

2.3 WORKING HYPOTHESIS

As I have already described, the CB₁ receptor expression is very high in inhibitory GABAergic synaptic terminals and low in glutamatergic neurons as well as in brain astrocytes. Subcellularly, CB₁ receptors are localized in the plasma membranes and in intracellular organelles, such as brain mitochondria. Due to the hugely different levels of expression among different cell types, it is extremely difficult to identify low CB₁ receptor expression in cell types and/or in subcellular compartments of wild-type brains. This led in the past to consider low CB₁ receptor levels as background staining (see below).

Based on methodological “loss of function” approaches, the use of mutant mice lacking CB₁ receptors in specific cell populations helped identifying the necessary role of the CB₁ receptor for some brain functions. Moreover, neuroanatomical analyses of these mutants determined the presence of the CB₁ receptor at sites previously not known (e.g. cortical glutamatergic neurons or neuronal mitochondria). However, these approaches do not provide definitive information concerning the possibly sufficient role of the cell-type expression of CB₁ receptors for a given function. Indeed, distinguishing between low levels (wild-type) and no expression (conditional mutant) is very difficult.

Therefore, to establish precise causal relationships and to identify low levels of expression, “rescue” strategies are needed to analyze the function of endogenous CB₁ receptor expression exclusively in specific cell types. Indeed, these rescue mice have allowed already determining the sufficient causal role of the CB₁ receptor in the anterior olfactory cortex area as well as in dorsal telencephalic glutamatergic neurons. However, so far no studies have addressed the subcellular localization of the CB₁ receptor in these mutant brains in the electron microscope. This is particularly important because the huge expression of the CB₁ receptor in specific cell types (e.g. hippocampal GABAergic interneurons) does not appreciate the detailed pattern of expression of the receptor in other cell types (e.g. hippocampal glutamatergic pyramidal neurons) in wild-type mouse samples. Hence, the conditional rescue mutants emerge as key tools for the detailed anatomical characterization of the subcellular distribution of the receptor in specific cell types, independently of its level of expression.

In this study, we have examined the CB₁ receptor expression in the hippocampus of rescue mice modified to express the gene exclusively in specific brain cell types, such as dorsal telencephalic glutamatergic neurons (Glu-CB₁-RS), GABAergic neurons (GABA-CB₁-RS) and brain astrocytes (GFAP-CB₁-RS), for three reasons. First, to elucidate the real levels of CB₁ receptors in specific cell types expressing very little CB₁ protein eliminating the interference of other cells highly expressing CB₁ receptors. Second, to verify that the re-expression is limited to the specific cell types and maintains endogenous levels of CB₁ receptors where the regulatory sequences of the receptor are normally activated. And third, to know whether the rescue mutant mice are good and reliable tools for functional studies of the endocannabinoid system in the brain. Finally, we have also analyzed the CB₁ receptor expression in the hippocampus of transgenic mice expressing the hrGFP protein in astrocytes (GFAPhrGFP-CB₁-WT and GFAPhrGFP-CB₁-KO mice) in order to investigate the CB₁ receptor distribution in astrocytes relative to the tripartite synapses.

3 OBJECTIVES

The general goal of this Thesis was the anatomical characterization of new different mutant mice expressing CB₁ receptors in specific brain cells. In particular, the study was focused on the hippocampus processed for a preembedding immunogold method for electron microscopy.

The proposed objectives of the present Thesis work are:

- To study the CB₁ receptor expression in the hippocampus of mice with the CB₁ receptor rescued in dorsal glutamatergic neurons (Glu-CB₁-RS).
- To investigate the CB₁ receptor expression in the hippocampus of mice with the CB₁ receptor rescued in GABAergic neurons (GABA-CB₁-RS).
- To determine the CB₁ receptor expression in the hippocampus of mice with the CB₁ receptor rescued in astrocytes (GFAP-CB₁-RS).
- To compare the CB₁ receptor expression in hippocampal astrocytes of transgenic mice with GFAP or with hrGFP as the astrocytic marker.
- To establish the anatomical distribution of the astrocytic CB₁ receptors relative to the surrounding excitatory and inhibitory synapse forming the tripartite synapse.

4 MATERIAL AND METHODS

I have used specific antibodies in combination with an immunocytochemical technique (immunoperoxidase) for light microscopy; a preembedding silver intensified immunogold method and a double preembedding immunogold and immunoperoxidase method for electron microscopy. In addition, I have done the corresponding quantification and statistical analysis of the data obtained in the electron microscopic experiments.

4.1 ANTIBODIES

Polyclonal Anti-Cannabinoid Receptor Type-1 (anti-CB1 receptor), monoclonal Anti-Glial Fibrillary Acidic Protein (anti-GFAP) and polyclonal humanized *Renilla reniformis* Green Fluorescence Protein (anti-hrGFP) antibodies were used (Table 1).

Table 1. Details of the antibodies used in this Thesis.

ANTIBODY	MANUFACTURER	HOST	CONCENTRATION
Polyclonal Anti-Cannabinoid Receptor Type-1 (CB ₁)	Frontier Institute	Goat	1:100
Monoclonal Anti-Glial Fibrillary Acidic Protein (GFAP)	Sigma-Aldrich	Mouse	1:1000
Polyclonal humanized <i>Renilla reniformis</i> Green Fluorescence Protein (hrGFP)	Stratagene	Rabbit	1:500

The experiments were done under the same conditions. Moreover, a negative control by omitting the primary antibody was performed. Importantly, the CB₁ receptor antibodies used were tested in CB₁-KO brain tissue. Finally, the specificity of the other antibodies has been thoroughly confirmed in previous studies.

CB₁: The sequence of the immunizing peptide used to generate the antibody corresponds with the last 31 amino acids of the C-terminus of the mouse CB₁ receptor (NM007726), as provided by the manufacturer (NCBI Reference Sequence: NP_031752.1; 443-473 amino acid residues: MHRAAESCISKSTVKIAKVTMSVSTDTSAEAL). The specificity of the CB₁ receptor antibody (Frontier Institute; Goat polyclonal; #CB1-Go-Af450) was assessed in previous publications (Bellocchio et al., 2010; Reguero et al., 2011; Bénard et al., 2012; Hebert-Chatelain et al., 2014b; Soria-Gómez et al., 2014; Martín-García et al., 2015). Furthermore, the specificity was tested in this study by applying the antibody to CB₁-KO hippocampus. No immunostaining was observed in this tissue in the light microscope. Also, the specificity of the CB₁ receptor pattern was analyzed in hippocampi of CB₁-WT and CB₁-KO processed together for immunoelectron microscopy. Thus, CB₁ receptor immunoparticles were localized in the inhibitory and excitatory synaptic terminals of CB₁-WT mouse while the CB₁ receptor

pattern disappeared in CB_1 -KO tissue, indicating the high specificity of the antibody used. Moreover, the antibody detects a single protein band of 52 kDa on immunoblots (Manufacturer's provided information, Frontier Science).

- **GFAP:** The specificity of the GFAP antibody (Sigma-Aldrich; mouse monoclonal; #G3893) was established in previous publications (Han et al., 2012; Bosier et al., 2013). This antibody has been tested for the GFAP immunolocalization in human, pig and rat tissues, and specifically detects GFAP on immunoblots. In addition, it does not cross react with vimentin, which is frequently co-expressed in glioma cells and some astrocytes (Manufacturer's provided information Sigma-Aldrich).
- **hr-GFP:** The humanized *Renilla reniformis* green fluorescence protein (hr-GFP) antibody (Stratagene; rabbit polyclonal; #240142) has been tested in brain and spinal cord and other tissues previously (Sakata et al., 2009; Zhang et al., 2013; Kerr et al., 2015). Moreover, anti-hrGFP antibody was not observed to bind to any cell in wild type mice.

4.2 RESEARCH ANIMALS

4.2.1 CB_1 RECEPTOR MUTANT LINES

4.2.1.1 Generation of the CB_1 -KO mice

CB_1 -KO mice were generated and genotyped as described (Marsicano et al., 2002). Conditional CB_1 receptor mutant mice were obtained by crossing the respective Cre expressing mouse line with CB_1^{ff} mice (Marsicano et al., 2003), using a three-step breeding protocol (Monory et al., 2006). All mice were in a predominant C57BL/6N background.

4.2.1.2 Generation of conditional mutant mice bearing a selective deletion of the CB_1 receptor in cortical glutamatergic neurons (hereafter Glu- CB_1 -KO)

The helix-loop-helix transcription factor NEX is a marker of embryonic neuronal progenitors, which will develop into mature cortical glutamatergic neurons (Wu et al., 2005). In the adult brain, NEX is expressed in mature glutamatergic cortical neurons, but not in cortical GABAergic interneurons and to a much lesser extent in subcortical regions (Bartholomä and Nave, 1994). Cre expression under the control of the regulatory sequences of NEX in mutant mice as generated by knock-in into the NEX locus (NEX-Cre mice), leads to the specific deletion of "floxed" alleles in forebrain neurons (Kleppisch et al., 2003). Thereby, $CB_1^{ff}; NEX-Cre$ mice Glu- CB_1 -KO, (Monory et al., 2006) were obtained by crossing CB_1^{ff} with NEX-Cre mice (Schwab et al., 2000; Kleppisch et al., 2003). Mutants were obtained by crossing CB_1^{ff} females with $CB_1^{ff}; NEX-Cre$ males.

4.2.1.3 Generation of conditional mutant mice bearing a selective deletion of the CB₁ receptor in GABAergic neurons (hereafter GABA-CB₁-KO)

The *Dlx5/Dlx6* genes are homeobox genes that are expressed in differentiating and migrating forebrain GABAergic neurons during embryonic development (Stuhmer, 2002). *Dlx5/6-Cre* (Zerucha et al., 2000) were crossed with *CB₁^{ff}* to obtain *CB₁^{ff}; Dlx5/6-Cre* mice. Thus, the expression of Cre recombinase under the control of the regulatory sequences of *Dlx5/Dlx6* genes drives to a recombination of loxP sites in GABAergic neurons (GABA-CB₁-KO mice, (Monory et al., 2006). Mutants were obtained by crossing *CB₁^{ff}* females with *CB₁^{ff}; DLX5/6-Cre* males.

4.2.1.4 Generation of conditional mutant mice bearing a selective deletion of the CB₁ receptor in astrocytes (hereafter GFAP-CB₁-KO)

Transgenic mice *CB₁^{ff}; GFAP-CreERT2* were obtained from crossing mice carrying CB₁ receptor “floxed” sequence (Marsicano et al., 2003) with transgenic mice expressing the inducible version of the Cre recombinase CreERT2 under the control of the human glial fibrillary acid protein promoter, i.e. GFAP-CreERT2 mice (Hirrlinger et al., 2006). This animal model allows the on-demand control of astroglial CB₁ receptor recombination in adult mice (Han et al., 2012).

4.2.1.5 Generation of Glu-CB₁-RS and GABA-CB₁-RS

Rescue mice were produced as previously described (Ruehle et al., 2013). Briefly, Stop-CB₁ mice were generated by inserting a loxP-flanked stop cassette into the 5'UTR of the coding exon of the *CB₁* gene, 32 nucleotides upstream of the translational start codon to obtain *CB₁^{stop/stop}* mice, lacking expression of the CB₁ protein (Ruehle et al., 2013). Conditional rescue mice were obtained by crossing Stop-CB₁ with NEX-Cre or *Dlx5/6-Cre* mice to generate Glu-CB₁-RS (rescue) and GABA-CB₁-RS, respectively. Genotyping was performed as previously described (Ruehle et al., 2013), and rescue mutants were obtained by crossing *CB₁^{stop/stop}* females with *CB₁^{stop/stop}; Nex-Cre* or *CB₁^{stop/stop}; DLX5/6-Cre* males, respectively.

4.2.1.6 Generation of GFAP-CB₁-RS

To obtain specific CB₁ rescue in astrocytes STOP-CB₁ mice (Ruehle et al., 2013) were crossed with GFAP-CreERT2 mice (Han et al., 2012) as previously described (Ruehle et al., 2013). Three weeks before neuroanatomical characterization, deletion of stop cassette for CB₁ rescue was achieved by injection of 4-hydroxytamoxifen as previously described (Han et al., 2012).

4.2.1.7 Generation of GFAPhrGFP-CB₁-WT and GFAPhrGFP-CB₁-KO

CB₁-WT and *CB₁-KO* mice received an intrahippocampal injection of a recombinant adeno associated virus expressing humanized Renilla GFP protein under the control of human GFAP promoter (von Jonquieres et al., 2013). Virus production and purification, as well as injection procedure were performed as

previously described (Chiarlone et al., 2014). After surgery, mice were allowed to recover for at least 4 weeks before anatomical characterization.

4.3 ANIMAL TREATMENT

The experiments were approved by the Committee of Ethics for Animal Welfare of the University of the Basque Country (CEEA/408/2015/GRANDES MORENO, CEIAB/213/2015/GRANDES MORENO). Animals were treated according to the European Community Council Directive 2010/63/UE and the Spanish legislation (RD 53/2013 and Ley 6/2013). Finally, great efforts were made in order to minimize the number and the suffering of the animals used.

4.3.1 TRANSCARDIALLY PERFUSION OF THE ANIMALS

1. C57BL/6N adult mice (between 60 and 90 postnatal days) of either sex were housed under standard conditions (food and water ad libitum; 12h/12h light/dark cycle). CB_1 wild type (hereafter CB_1 -WT), Glu- CB_1 -KO, GABA- CB_1 -KO, GFAP- CB_1 -KO, CB_1 -STOP, Glu- CB_1 -RS, GABA- CB_1 -RS, GFAP- CB_1 -RS, CB_1 -KO, GFAPhrGFP- CB_1 -WT and GFAPhrGFP- CB_1 -KO mice (at least three animals of 3 each condition) were deeply anesthetized by intraperitoneal injection of ketamine/xylazine (80/10 mg/kg body weight).
2. Animals were transcardially perfused at room temperature (20-25 °C) with phosphate buffered saline (0.1 M, pH 7.4) for 20 seconds, followed by the fixative solution made up of 4% formaldehyde (freshly depolymerized from paraformaldehyde), 0.2% picric acid, and 0.1% glutaraldehyde in phosphate buffer (0.1 M Phosphate buffer, pH 7.4) for 10-15 minutes.
3. Brains were carefully removed from the skull and post-fixed in the fixative solution for approximately 1 week at 4 °C. Finally, brains were stored at 4 °C in 1:10 diluted fixative solution plus 0.025% sodium azide at 4°C until its processing.

4.3.2 AVIDIN-BIOTIN PEROXIDASE METHOD FOR LIGHT MICROSCOPY

This is the protocol applied:

1. Brain coronal sections were cut at 50 μ m in a vibratome and collected in 0.1 M Phosphate buffer at room temperature.
2. Preincubation of the hippocampal sections with blocking solution composed by 10% horse serum, 0.1% sodium azide and 0.5% triton X-100 prepared in Tris-Hydrogen Chloride buffered saline 1X, pH 7.4 for 30 minutes at room temperature.
3. Incubation with the primary anti- CB_1 receptor antibody (1:100) prepared in the blocking solution, with continuous gentle shaking for 2 days at 4 °C.
4. Several washes in 1% horse serum and 0.5% triton X-100 in Tris-Hydrogen Chloride buffered saline for 30 minutes to remove excess of the antibody.
5. Incubation with a secondary biotinylated horse anti-goat Inmunoglobulin-G (1:200, Vector Laboratories, Burlingame, CA, USA) prepared in the washing solution for 1 hour on a shaker at room temperature.

6. Washes with 1% horse serum and 0.5% triton X-100 in Tris-Hydrogen Chloride buffered saline.
7. Incubation in 1:50 avidin-biotin complex (Avidin-biotin peroxidase complex, Elite, Vector Laboratories, Burlingame, CA, USA) prepared in the washing solution for 1 hour at room temperature.
8. Several rinses with 1% horse serum and 0.5% triton X-100 in Tris-Hydrogen Chloride buffered saline. Last rinses were done with 0.1M phosphate buffer and 0.5% triton X-100.
9. Incubation with 0.05% diaminobenzidine as a chromogen in 0.1 M phosphate buffer containing 0.5% triton-X100 and 0.01% hydrogen peroxide, for 5 minutes at room temperature.
10. Several washes in 0.1 M phosphate buffer with 0.5% triton-X100.
11. Mounting on gelatinized slides.
12. Dehydration in graded alcohols (50°, 70°, 96°, 100°) for 5 minutes each.
13. Clearing in xylol (3 times of 5 min).
14. Coverslipped with DPX.
15. Tissue sections were observed and photographed with a Zeiss Axiocam light microscope coupled to a Zeiss AxioCam HRc Camera.

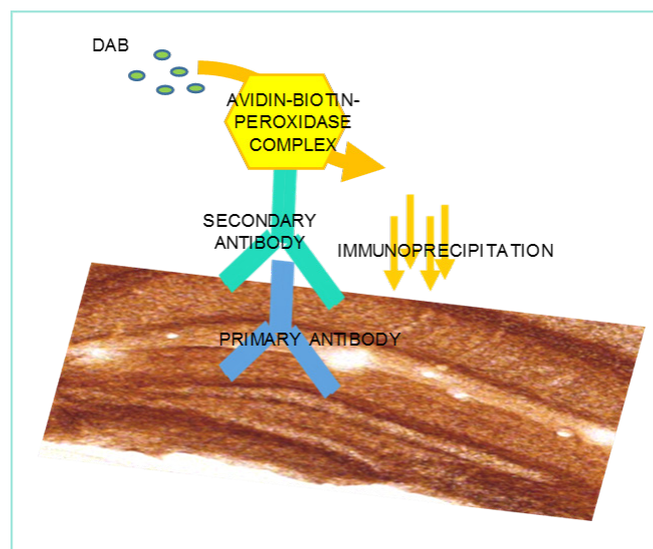


Figure 9. Graphical representation of the Immunoperoxidase method for light microscopy.

4.3.3 PREEMBEDDING SILVER-INTENSIFIED IMMUNOGOLD METHOD FOR ELECTRON MICROSCOPY

The following protocol was applied:

1. Coronal hippocampal vibrosections were cut at 50 μm and collected in 0.1 M phosphate buffer (pH 7.4) at room temperature.
2. Preincubation of the sections in a blocking solution of 10% bovine serum albumin, 0.1% sodium azide, and 0.02% saponin prepared in Tris-

- Hydrogen Chloride buffered saline 1X, pH 7.4 for 30 minutes at room temperature.
3. Incubation with the primary goat polyclonal anti-CB₁ receptor antibody (1:100) in 10% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline containing 0.1% sodium azide and 0.004% saponin on a shaker for 2 days at 4°C.
 4. Several washes in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline.
 5. Incubation in a secondary 1.4 nm gold-labeled rabbit anti-goat Immunoglobulin-G (Fab' fragment, 1:100, Nanoprobes Inc., Yaphank, NY, USA) in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline with 0.004% saponin on a shaker for 3 hours at room temperature.
 6. Several washes in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline overnight on a shaker at 4°C.
 7. Post-fixation with 1% glutaraldehyde prepared in Tris-Hydrogen Chloride buffered saline for 10 minutes at room temperature.
 8. Several washes in double distilled water for 30 min.
 9. Silver intensification of gold particles with a HQ Silver kit (Nanoprobes Inc., Yaphank, NY, USA) for about 12 minutes in the dark.
 10. Several washes in double distilled water for 10 minutes.
 11. Several washes in 0.1 M phosphate buffer (pH 7.4) for 30 minutes.
 12. Osmication (1% osmium tetroxide, in 0.1 M phosphate buffer pH 7.4) for 20 minutes.
 13. Several washes in 0.1 M phosphate buffer (pH 7.4) for 30 minutes.
 14. Dehydration in graded alcohols (50°, 70°, 96° and 100°) for 5 min each and 3 times of 5 min for 100°.
 15. Clearing in propylene oxide 3 times, 5 min each.
 16. Embedding in a mixture of 1:1 propylene oxide and Epon resin 812 overnight on a shaker at room temperature.
 17. Resin polymerization in a heater at 60°C for 2 days.
 18. Cutting 1µm semithin sections in the Reichert-Jung ultracut.
 19. Collection of 60nm ultrathin sections on mesh nickel grids.
 20. Staining with 2.5% lead citrate for 20 min.
 21. Examination under a Philips EM208S electron microscope. Tissue was photographed by using a digital Morada Camera from Olympus coupled to the electron microscope.

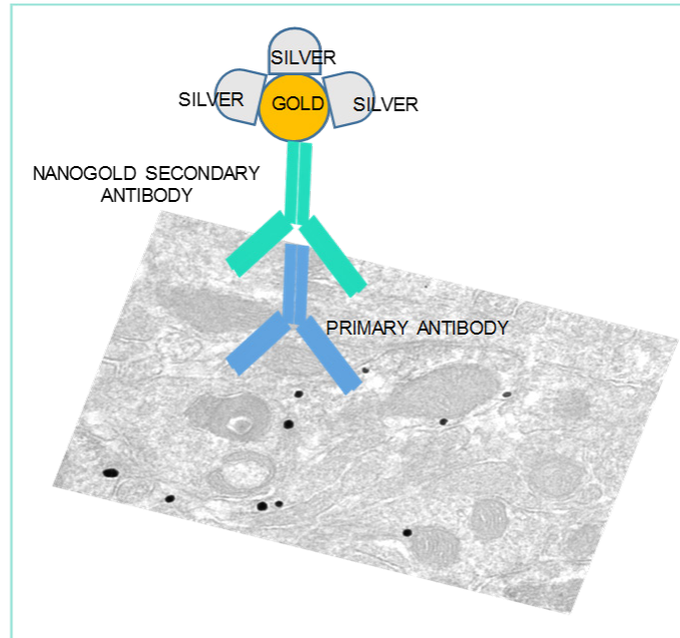


Figure 10. Graphical representation of the Immunoperoxidase method for light microscopy.

4.3.4 PREEMBEDDING DOUBLE LABELING OF SILVER-INTENSIFIED IMMUNOGOLD AND IMMUNOPEROXIDASE METHOD FOR ELECTRON MICROSCOPY

This is the protocol followed:

1. Coronal hippocampal vibrosections were cut at 50 μm and collected in 0.1 M phosphate buffer (pH 7.4) at room temperature.
2. Preincubation in a blocking solution of 10% bovine serum albumin, 0.1% sodium azide, and 0.02% saponin prepared in Tris-Hydrogen Chloride buffered saline 1X, pH 7.4 for 30 minutes at room temperature.
3. Incubation with the primary goat polyclonal anti- CB_1 receptor antibody (1:100) and the primary mouse monoclonal anti-GFAP (1:1000) or the primary rabbit polyclonal hrGFP (1:500) prepared in the blocking solution in 10% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline containing 0.1% sodium azide and 0.004% saponin on a shaker for 2 days at 4°C.
4. Several washes in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline.
5. Incubation with the corresponding biotinylated secondary antibody (1:200) in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline with 0.004% saponin on a shaker for 3 hours at room temperature.
6. Several washes in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline overnight on a shaker at 4°C.
7. Incubation in a secondary 1.4 nm gold-labeled rabbit anti-goat Immunoglobulin-G (Fab' fragment, 1:100, Nanoprobes Inc., Yaphank, NY, USA) in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline with 0.004% saponin on a shaker for 3 hours at room temperature.

8. Several washes in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline for 30 minutes on a shaker at 4°C.
9. Incubation in avidin-biotin complex (1:50) prepared in the washing solution for 1.5 hour at room temperature.
10. Several washes in 1% Bovine Serum Albumin / Tris-Hydrogen Chloride buffered saline overnight on a shaker at 4°C.
11. Post-fixation with 1% glutaraldehyde in Tris-Hydrogen Chloride buffered saline for 10 minutes at room temperature.
12. Several washes in double distilled water for 30 min.
13. Silver intensification of the gold particles was with a HQ Silver kit (Nanoprobes Inc., Yaphank, NY, USA) for about 12 minutes in the dark.
14. Several washes in double distilled water for 10 minutes.
15. Several washes in 0.1 M phosphate buffer (pH 7.4) for 30 minutes.
16. Incubation in 0.05% DAB and 0.01% hydrogen peroxide prepared in 0.1 M phosphate buffer for 5 minutes at room temperature.
17. Osmication (1% osmium tetroxide, in 0.1 M phosphate buffer pH 7.4) for 20 minutes).
18. Several washes in 0.1 M phosphate buffer (pH 7.4) for 30 minutes.
19. Dehydration in graded alcohols (50°, 70°, 96° and 100°) for 5 min each and 3 times of 5 min for 100°.
20. Clearing in propylene oxide 3 times, 5 min each.
21. Embedding in a mixture of 1:1 propylene oxide and Epon resin 812 overnight on a shaker at room temperature.
22. Resin polymerization in a heater at 60°C for 2 days.
23. Cutting 1µm semithin sections in the Reichert-Jung ultracut.
24. Collection of 60nm ultrathin sections on mesh nickel grids.
25. Staining with 2.5% lead citrate for 20 min.
26. Examination under a Philips EM208S electron microscope. Tissue preparations were photographed by using a digital Morada Camera from Olympus coupled to the electron microscope.

4.4 SEMI-QUANTIFICATION OF THE CB1 RECEPTOR IMMUNOGOLD STAINING

With the aim of maximizing the standard conditions, the pre-embedding immunogold method was systematically applied simultaneously to all the sections collected from the all different animals (n=3 of each condition) in each of the three replicated experiments done for each mouse performed.

Immunogold-labeled hippocampal sections were then visualized in the light microscope in order to select portions of the CA1 and the dentate molecular layer with good and reproducible CB₁ receptor immunolabeling. Then the semithin sections were cut with an ultra-microtome and just the first 5 ultrathin sections (60 nm thick) were collected onto two grids. Moreover, to further standardize the conditions between the immunolabeled sections obtained from the different mice, only the first 1.5 µm from the section surface was photographed. All electron micrographs were taken at 18,000x using a Digital Morada Camera from Olympus. Sampling was always performed accurately in the same way for all the animals studied.

The analyzed excitatory and inhibitory synapses were identified by their characteristic ultrastructural features. Hence, excitatory synapses were distinguished by their asymmetric synapses with thick postsynaptic densities and presynaptic terminals containing abundant, clear and spherical synaptic vesicles. Inhibitory synapses were identified by their symmetric synaptic contacts and terminal boutons with pleomorphic synaptic vesicles.

To determine the proportion of the labeled CB₁ receptor terminals, positive labeling was considered if at least one immunoparticle was within approximately 30 nm from the plasmalemma of the synaptic terminal. Metal particles at these synaptic membranes were visualized and counted. Immunopositive astrocytic processes were considered positive if the precipitates of DAB immunoreaction product for hrGFP or GFAP, were inside the astrocytic elements. Image-J software was used to measure the membrane length and distance to synapses. Percentages of CB₁ receptor positive profiles, density (particles/ μm membrane) and proportion of CB₁ immunoparticles in terminals versus total CB₁ receptor expression in cellular membranes were analyzed and displayed as mean \pm Standard Error of the Mean using a statistical software package (GraphPad Prism, GraphPad Software Inc, San Diego, USA). The normality test (Kolmogorov-Smirnov normality test) was applied before running statistical tests and subsequently data were analyzed using nonparametric tests (Man-Whitney U test when $k=2$ or Kruskal-Wallis test when $k>2$). A potential variability between mice of the same mutant line was analyzed statistically. Since there were no differences between them, all data from each line were pooled.

4.5 SEMI-QUANTIFICATION OF THE DISTANCE FROM ASTROCYTIC CB₁ RECEPTORS TO THE NEAREST SYNAPSE.

Immunogold and immunoperoxidase hippocampal sections were visualized in the light microscope in order to select portions of the CA1 and the dentate molecular layer with good and reproducible immunolabeling. The first 5 ultrathin sections (60 nm thick) were collected onto two grids. Moreover, to further standardize the conditions between the immunolabeled sections obtained from the different mice, only the first 1.5 μm from the section surface was photographed. Once again, all electron micrographs were taken at 18,000x using a Digital Morada Camera from Olympus. Sampling was always accurately done in the same way for all the animals studied.

Positive astrocytic processes were considered by the presence of immunoprecipitation of DAB in the cytoplasm (see above). Moreover, to determine the proportion of labeled CB₁ receptor astrocytic processes, positive labeling was considered if at least one immunoparticle was within approximately 30 nm from the plasmalemma.

Afterward, the nearby synapses surrounding the immunopositive astrocytic process were identified, the distances were measured using the Image-J software, and the nearest synapse to the astrocytic CB₁ receptor immunogold particle was selected. The data from all the nearest synapses to the astrocytic CB₁ receptor particles were tabulated, analyzed and displayed as mean \pm S.E.M. using a

statistical software package (GraphPad Prism, GraphPad Software Inc, San Diego, USA).

5 RESULTS

5.1 CB₁ RECEPTOR DISTRIBUTION IN THE BRAIN OF GLU-CB₁-RS AND GABA-CB₁-RS MICE. LIGHT MICROSCOPY

The pattern of CB₁ receptor immunostaining was first analyzed in the Glu-CB₁-RS and GABA-CB₁-RS brains by light microscopy (Figs. 11 and 12). As expected, a much weaker and diffuse CB₁ receptor pattern was observed in Glu-CB₁-RS than in GABA-CB₁-RS, in accordance with the distribution of CB₁ receptor in glutamatergic and GABAergic cells, respectively (Figs. 11 and 12). Thus, noticeable CB₁ receptor staining was observed in the striatum, cortex, olfactory tubercle and amygdala of Glu-CB₁-RS (Fig. 11). In GABA-CB₁-RS, strong CB₁ receptor immunoreactivity was detected in the cortex, anterior olfactory nucleus, piriform cortex, globus pallidus, entopeduncular nucleus, amygdala and the substantia nigra (Fig. 12). The striatum exhibited moderate to strong immunostaining (Fig. 12). Altogether, the overall CB₁ receptor pattern in Glu-CB₁-RS and GABA-CB₁-RS matches the known receptor distribution in the brain.

5.2 CB₁ RECEPTOR DISTRIBUTION IN HIPPOCAMPUS OF GLU-CB₁-RS AND GABA-CB₁-RS MICE. LIGHT MICROSCOPY

We particularly focused on the pattern of the CB₁ receptor immunostaining in the hippocampus (Fig.13). In CB₁-WT, a distinct intensity pattern was observed throughout the layers (Fig. 13a). Thus, the strongest CB₁ receptor immunostaining was revealed in the stratum pyramidale and at the limit between the strata radiatum and lacunosum-moleculare. Also, a dense neuropil labeling was seen throughout the strata radiatum and oriens. In the dentate gyrus, the most prominent CB₁ receptor staining localized in the innermost third of the stratum moleculare whereas a remarkable but weaker staining was found in the outer 2/3 of the layer. Finally, immunoreactive fibrous processes surrounding the granule cell bodies were visible in the stratum granulosum (Fig. 13a).

In the Glu-CB₁-RS hippocampus, a faint CB₁ receptor immunoreactivity was observed throughout the strata oriens and radiatum of the hippocampal Ammon's horn (Cornu ammonis, CA) (Fig. 13b). Remarkably, a strong band of CB₁ receptor immunostaining appeared in the innermost third of the dentate molecular layer which corresponds to the zone of the synapses of the commissural/associational system (Fig. 13b) (Monory et al., 2006; Ruehle et al., 2013). In GABA-CB₁-RS, heavy CB₁ receptor immunoreactivity was distributed throughout the CA stratum radiatum and the dentate molecular layer (Fig. 13c). However, a stronger CB₁ receptor staining was found in the pyramidal cell layer and at the limit between strata radiatum and lacunosum-moleculare of the CA1, as well as in the molecular innermost third and the infragranular zone of the dentate gyrus (Fig.13c). Importantly, the CB₁ receptor immunostaining was absent in the CB₁-KO hippocampus, indicating the specificity of the CB₁ receptor antibody used (Fig. 13d).

5.3 SUBCELLULAR LOCALIZATION OF THE CB₁ RECEPTOR IN HIPPOCAMPUS OF GLU-CB₁-RS AND GABA-CB₁-RS. HIGH RESOLUTION ELECTRON MICROSCOPY

To study the localization of CB₁ receptor in detail, a preembedding immunogold method for electron microscopy was used. Hence, hippocampal sections of CB₁-WT, CB₁-KO, Glu-CB₁-RS, GABA-CB₁-RS, STOP-CB₁, Glu-CB₁-KO and GABA-CB₁-KO mice were used to investigate the cellular and subcellular localization of the receptor in the CA1 stratum radiatum (Fig.14) and in the inner third of the dentate molecular layer (Fig.15). As expected in CB₁-WT, presynaptic inhibitory terminal membranes forming symmetric synapses were decorated with a high density of CB₁ receptor immunoparticles, whereas a much lower labeling was observed in excitatory terminals making asymmetric synapses. CB₁ receptor immunoparticles were located at a distance from the active zones of the symmetric and asymmetric synapses formed with postsynaptic dendrites and dendritic spines, respectively (Figs.14a and 15a). Noticeably, the CB₁ receptor pattern virtually disappeared in CB₁-KO meaning that the CB₁ receptor antibody used was highly specific (Figs. 14b and 15b).

In Glu-CB₁-RS, the CB₁ receptor was re-expressed in excitatory synapses identified by their typical ultrastructural features, namely, axon boutons with abundant, clear and spherical synaptic vesicles forming asymmetric synapses with postsynaptic dendritic spines, whereas CB₁ receptor immunolabeling was absent in symmetric synapses (Figs. 14 d and 15d). CB₁ receptor was also re-expressed in synaptic terminals of the commissural/associational system in the inner third of the dentate molecular layer that contained abundant tightly packed synaptic vesicles and forming asymmetric synapses with granule cell dendrites and dendritic spines (Fig. 15d) (Ribak et al., 1985; Li et al., 1994; Blasco-Ibáñez and Freund, 1997; Scharfman, 2007). By contrast, in Glu-CB₁-KO CB₁ receptor immunonegative excitatory synapses but numerous CB₁ receptor immunoparticles accumulated in terminals with pleomorphic synaptic vesicles and symmetric synapses with dendrites were observed (Figs.14e and 15e).

In the case of GABA-CB₁-RS, rich CB₁ receptor immunolabeling was restricted to presynaptic terminal plasma membranes making symmetric synapses with dendrites; no immunoparticles were found at excitatory synapses (Figs.14f and 15 f). On the other hand, only presynaptic boutons forming asymmetric synapses with spines were CB₁ receptor immunopositive in GABA-CB₁-KO (Figs.14g and 15g). No immunolabeling was observed in excitatory or inhibitory synapses of STOP-CB₁ hippocampus (Figs.14c and 15c). In addition, CB₁ receptor immunoparticles were also observed in mitochondrial membranes of CB₁-WT and the conditional mutants, but not in CB₁-KO (Figs.14b and 15b)

5.4 SUBCELLULAR LOCALIZATION OF THE CB₁ RECEPTOR IN HIPPOCAMPUS OF GFAP-CB₁-RS AND GFAPhrGFP-CB₁-WT. HIGH RESOLUTION ELECTRON MICROSCOPY

For the study in detail of the localization of CB₁ receptors in astrocytes of the hippocampus, a preembedding immunogold and immunoperoxidase method for electron microscopy were applied. In order to investigate the cellular and subcellular localization of the receptor in the CA1 stratum radiatum (Fig.16) and in the dentate molecular layer (Fig.17), hippocampal sections of *CB₁-WT*, *CB₁-KO*, *GFAP-CB₁-RS*, *STOP-CB₁* and *GFAP-CB₁-KO* mice were used. Besides, *CB₁-WT* and *CB₁-KO* mice expressing hrGFP in astrocytes (*GFAPhrGFP-CB₁-WT* and *GFAPhrGFP-CB₁-KO* respectively) were also analyzed.

As expected, CB₁ receptor immunoparticles were distributed in astrocytic processes membranes and terminal plasmalemma of *CB₁-WT* (Figs.16a and 17a) and of *GFAPhrGFP-CB₁-WT* (Figs 16f, 16g, 17f and 17g). However, in *GFAP-CB₁-RS*, CB₁ receptor immunoparticles appeared only in astrocytic processes but not in synaptic terminals (Figs. 16c and 17c). Conversely in *GFAP-CB₁-KO* (Figs. 16d and 17d), CB₁ receptor labeling disappeared from astrocytes membranes whereas the pattern of the CB₁ receptor in synaptic terminals remained unchangeable relative to *CB₁-WT* (Figs. 16a and 17a). Importantly, there was not CB₁ receptor immunolabeling in synaptic terminals nor in astrocytic processes of *STOP-CB₁* hippocampus (Figs.16e and 17e). Furthermore, the CB₁ receptor staining pattern disappeared in *CB₁-KO* (Figs. 16b and 17b) and in *GFAPhrGFP-CB₁-KO* (Figs. 16h and 17h) hence demonstrating the great specificity of the anti-CB₁ receptor antibody used.

5.5 STATISTICS OF THE CB₁ RECEPTOR IN EXCITATORY TERMINALS OF THE GLU-CB₁-RS AND GABA-CB₁-RS HIPPOCAMPUS

The proportion of CB₁ receptor immunopositive excitatory synaptic terminals in the CA1 stratum radiatum of *CB₁-WT* (24.29% ± 1.09), *Glu-CB₁-RS* (21.89% ± 1.21) and *GABA-CB₁-KO* (21.37% ± 1.13) was not significantly different (Fig. 18a). However, the percentage of positive terminals greatly decreased in *Glu-CB₁-KO* (2.50% ± 0.76) and virtually disappeared in *STOP-CB₁*, *GABA-CB₁-RS* and *CB₁-KO* (Fig. 18a). Furthermore, no statistical differences were found in CB₁ receptor density (particles/μm) between excitatory synaptic terminals of *CB₁-WT* (0.45 ± 0.01), *Glu-CB₁-RS* (0.45 ± 0.02) and *GABA-CB₁-KO* (0.46 ± 0.02) (Fig. 18b). Finally, the proportion of CB₁ receptor gold particles in asymmetric terminals versus total CB₁ receptor expression in plasmalemmal structures was analyzed in CA1 stratum radiatum. 88.78% ± 1.96 of the total CB₁ receptor immunoparticles were located in excitatory terminals of *Glu-CB₁-RS*, while 12.35% ± 1.04 and 27.30% ± 3.34 were found in *CB₁-WT* and *GABA-CB₁-KO* respectively. Only residual CB₁ receptor immunoparticles were observed in *STOP-CB₁*, *GABA-CB₁-RS*, *Glu-CB₁-KO* and *CB₁-KO* (Fig. 18c).

The analysis of the inner third of the dentate molecular layer showed comparable percentages of CB₁ receptor immunopositive excitatory synaptic terminals in *CB₁-WT* (55.98% ± 2.51), *Glu-CB₁-RS* (53.19% ± 2.89) and *GABA-CB₁-KO* (52.12% ± 3.00) (Fig. 19a). The proportion decreased in *Glu-CB₁-KO* (5.48% ± 1.70) and practically disappeared in *STOP-CB₁*, *GABA-CB₁-RS* and *CB₁-KO* (Fig. 19a). CB₁ receptor immunoparticle density, expressed as the number of particles per μm of

cell perimeter, was statistically similar in *CB₁*-WT (0.39 ± 0.02), *Glu-CB₁*-RS (0.39 ± 0.02) and *GABA-CB₁*-KO (0.39 ± 0.02) (Fig. 19b). Furthermore, the proportion of *CB₁* receptor immunogold particles in glutamatergic terminals versus total *CB₁* receptor expression in plasmalemma was $12.80\% \pm 1.39$ in *CB₁*-WT, $88.64\% \pm 1.71$ in *Glu-CB₁*-RS, $29.22\% \pm 3.65$ in *GABA-CB₁*-KO and only residual *CB₁* receptor immunoparticles were found in excitatory terminals of *GABA-CB₁*-RS, *Glu-CB₁*-KO, *STOP-CB₁* and *CB₁*-KO mice (Fig. 19c). Altogether, these results indicate that the subcellular *CB₁* receptor distribution and expression levels at excitatory synapses were preserved in the *Glu-CB₁*-RS and *GABA-CB₁*-KO mutants relative to the wild type, remaining residual at the same synapses of *Glu-CB₁*-KO and *GABA-CB₁*-RS mice.

5.6 STATISTICS OF THE *CB₁* RECEPTOR IN INHIBITORY TERMINALS OF THE *GLU-CB₁*-RS AND *GABA-CB₁*-RS HIPPOCAMPUS

As to the inhibitory synapses, the amount of *CB₁* receptor immunopositive terminals in *CB₁*-WT ($78.50\% \pm 2.44$), *GABA-CB₁*-RS ($77.92\% \pm 2.63$) and *Glu-CB₁*-KO ($79.63\% \pm 2.35$) was maintained in the CA1 stratum radiatum (Fig. 20a). However, this proportion drastically decreased in *STOP-CB₁*, *GABA-CB₁*-KO and *Glu-CB₁*-RS and virtually disappeared in *CB₁*-KO (Fig. 20a). There were not statistically significant differences in *CB₁* receptor immunoparticle density between inhibitory synaptic terminals of *CB₁*-WT (4.57 ± 0.13), *GABA-CB₁*-RS (4.33 ± 0.11) and *Glu-CB₁*-KO (4.50 ± 0.15) (Fig. 20b). Furthermore, $44.06\% \pm 2.67$ of the total immunoparticles in *CB₁*-WT, $97.10\% \pm 0.46$ in *GABA-CB₁*-RS and $54.87\% \pm 3.94$ in *Glu-CB₁*-KO were located in inhibitory terminals. Just residual particles were observed in *STOP-CB₁*, *Glu-CB₁*-RS, *GABA-CB₁*-KO and *CB₁*-KO mice (Fig. 20c).

The proportion of *CB₁* receptor immunopositive inhibitory synaptic terminals in the inner third of the dentate molecular layer was found to be similar between *CB₁*-WT ($84.27\% \pm 2.38$), *GABA-CB₁*-RS ($85.07\% \pm 1.76$) and *Glu-CB₁*-KO ($88.02\% \pm 1.93$) (Fig. 21a). These values almost disappeared in *STOP-CB₁*, *GABA-CB₁*-KO, *Glu-CB₁*-RS and in *CB₁*-KO (Fig. 21a). No statistical differences in labeling density were found between inhibitory terminals of *CB₁*-WT (7.54 ± 0.16), *GABA-CB₁*-RS (7.47 ± 0.14) and *Glu-CB₁*-KO (7.69 ± 0.21) (Fig. 21b). Moreover, $45.38\% \pm 3.18$ of the total immunoparticles in the inner third of the dentate molecular layer were located in GABAergic terminals of *CB₁*-WT, $93.87\% \pm 0.93$ in *GABA-CB₁*-RS and $59.45\% \pm 4.00$ in *Glu-CB₁*-KO, whereas only residual particles were found in inhibitory terminals of *STOP-CB₁*, *Glu-CB₁*-RS, *GABA-CB₁*-KO and *CB₁*-KO mice (Fig. 21c). Hence, the subcellular distribution and expression levels of *CB₁* receptors at inhibitory synapses were kept in the *GABA-CB₁*-RS and *Glu-CB₁*-KO mutants relative to the wild type, remaining residual at the same synapses of *STOP-CB₁*, *GABA-CB₁*-KO and *Glu-CB₁*-RS mice.

5.7 STATISTICS OF THE CB₁ RECEPTOR IN ASTROCYTIC PROCESSES OF THE GFAP-CB₁-RS AND GFAPhrGFP-CB₁-WT HIPPOCAMPUS

The analysis of the percentage of CB₁ receptor immunopositive astrocytic processes (Fig. 22a) did not show statistical differences between CB₁-WT (42.06% ± 3.56) and GFAP-CB₁-RS (37.12% ± 3.79) in the CA1 stratum radiatum. In GFAPhrGFP-CB₁-WT, the proportion of astrocytic elements immunolabeled for CB₁ receptors increased to 59.91% ± 3.29 being the difference statistically significant when compared with CB₁-WT. Conversely, only scarce CB₁ receptor immunolabeling was found in astrocytes of GFAP-CB₁-KO, STOP-CB₁, CB₁-KO and in GFAPhrGFP-CB₁-KO mice.

Additionally, CB₁ receptor immunoparticle density on astrocytic membranes (particles/μm) was analyzed (Fig. 22b). Our data showed statistically similar densities in CB₁-WT (0.135 ± 0.019) and GFAP-CB₁-RS (0.128 ± 0.020), whereas a great statistically significant increase was found in GFAPhrGFP-CB₁-WT (0.384 ± 0.039). Nevertheless, only residual unspecific particles were found in GFAP-CB₁-KO (0.005 ± 0.003), in STOP-CB₁ (0.005 ± 0.003), in CB₁-KO (0.001 ± 0.001) and in GFAPhrGFP-CB₁-KO (0.004 ± 0.002).

Finally, the proportion of CB₁ immunogold particles in astrocytic processes versus total CB₁ expression in plasmalemma was analyzed in CA1 stratum radiatum (Fig. 22c). 95.45% ± 1.82 of the total CB₁ immunoparticles were located in astrocytic processes of GFAP-CB₁-RS, while 23.08% ± 1.99 of the total immunogold particles were in the astrocytic elements of CB₁-WT. This proportion reached 33.71% ± 2.75 in GFAPhrGFP-CB₁-WT. As expected, only residual CB₁ receptor immunoparticles were located in astrocytes of GFAP-CB₁-KO, STOP-CB₁, GFAPhrGFP-CB₁-KO and CB₁-KO mice.

Furthermore, 75.13% ± 4.06 of the total CB₁ immunoparticles in plasmalemma were in terminal membranes of GFAP-CB₁-KO, while 65.52% ± 2.44 were in CB₁-WT, and 56.32% ± 2.73 in GFAPhrGFP-CB₁-WT (Fig. 22d). As expected, only residual CB₁ receptor immunoparticles were located in astrocytic processes of GFAP-CB₁-RS, STOP-CB₁, GFAPhrGFP-CB₁-KO and CB₁-KO mice (Fig. 22d).

The analysis of the hippocampal dentate molecular layer showed comparable percentages of CB₁ receptor immunopositive astrocytic processes between CB₁-WT (44.67% ± 3.85) and GFAP-CB₁-RS (39.84% ± 3.50), whereas the proportion significantly increased to 59.99% ± 3.37 in GFAPhrGFP-CB₁-WT. The proportion decreased in GFAP-CB₁-KO, STOP-CB₁, GFAPhrGFP-CB₁-KO and CB₁-KO mice (Fig. 23a).

Besides, no statistical differences were found between the CB₁ receptor density (particles/μm) on astrocytic processes of CB₁-WT (0.112 ± 0.011) and GFAP-CB₁-RS (0.138 ± 0.016) mice but there was when compared with GFAPhrGFP-CB₁-WT (0.334 ± 0.033) (Fig. 23b). However, only residual particles were observed in GFAP-CB₁-KO (0.006 ± 0.003), STOP-CB₁ (0.006 ± 0.003), GFAPhrGFP-CB₁-KO (0.002 ± 0.002) and CB₁-KO mice (0.004 ± 0.002) (Fig. 23b).

Furthermore, we examined the proportion of CB₁ receptor labeling in astrocytic processes versus total CB₁ receptor expression (Fig. 23c). The values were

22.40% \pm 2.28 in *CB₁*-WT, 95.61% \pm 1.56 in GFAP-*CB₁*-RS and 32.57% \pm 2.09 in GFAPhrGFP-*CB₁*-WT. Only non-specific *CB₁* receptor immunoparticles were found in astrocytic processes of GFAP-*CB₁*-KO, STOP-*CB₁*, and GFAPhrGFP-*CB₁*-KO and *CB₁*-KO mice (Fig. 23c).

Finally, *CB₁* gold labeling in terminal membranes versus total *CB₁* expression was also studied. 76.17% \pm 4.70 of the total *CB₁* immunoparticles were located in terminals of GFAP-*CB₁*-KO, 64.27% \pm 2.88 in *CB₁*-WT and 57.17% \pm 2.19 in GFAPhrGFP-*CB₁*-WT (Fig. 23d). Just residual particles were in astrocytic processes of GFAP-*CB₁*-RS, STOP-*CB₁*, GFAPhrGFP-*CB₁*-KO and *CB₁*-KO mice (Fig. 23d).

5.8 STATISTICAL ANALYSES OF THE DISTANCE FROM ASTROCYTIC *CB₁* RECEPTORS TO THE NEAREST SYNAPSE

Taking advantage of the better detection of astrocytes achieved with the hr-GFP protein in the electron microscope, the distance between the astrocytic *CB₁* receptor immunoparticles and the midpoint of the nearest synapse surrounded by the immunopositive astrocytic element was measured in GFAPhrGFP-*CB₁*-WT.

In CA1 (Fig 24a), 50.76% \pm 4.10 of the total synapses surrounded by astrocytic processes were at a distance between 400-800 nm away from the nearest astrocytic immunoparticle. Of those, 40.89% \pm 2.57 were excitatory synapses whereas 9.86% \pm 1.97 were inhibitory synapses. 20.60% \pm 6.1 of the synapses were found to be located in a range of 0 nm to 400 nm from the astrocytic *CB₁* receptor to the nearest synapse. Of these, 17.52% \pm 5.63 were excitatory synapses and only 3.08% \pm 0.49 were inhibitory synapses. 18.46% \pm 2.53 of the synapses were found to be located in a range of 800 nm to 1200 nm of which 13.96% \pm 1.60 were excitatory and 4.50% \pm 0.95 were inhibitory synapses. Only 10.18% \pm 7.81 synapses were found at more than 1200 nm from the astrocytic *CB₁* receptor metal particle, being 9.62% \pm 8.01 of them excitatory and only 0.56% \pm 0.56 were inhibitory synapses.

In the dentate molecular layer (Fig 24b), 57.25% \pm 3.19 of the synapses surrounded by astrocytic processes were at 400-800 nm from the astrocytic *CB₁* receptor particle, 46.67% \pm 2.17 of them were excitatory synapses and the other 10.59% \pm 2.18 were inhibitory synapses. 19.83% \pm 0.58 of the synapses were in a range of 0 nm to 400 nm from the astrocytic *CB₁* receptor metal particle of which 16.67% \pm 1.31 were excitatory and 3.16% \pm 0.82 inhibitory synapses. Finally, 16.42% \pm 4.37 of the total synapses were found between 800 nm and 1200 nm; 13.22% \pm 3.19 being excitatory and 3.21% \pm 2.25 inhibitory synapses. Only 6.48% \pm 2.31 synapses were found at more than 1200 nm from the astrocytic *CB₁* receptor. 5.16% \pm 2.72 of them were excitatory and 1.32% \pm 0.66 inhibitory synapses.

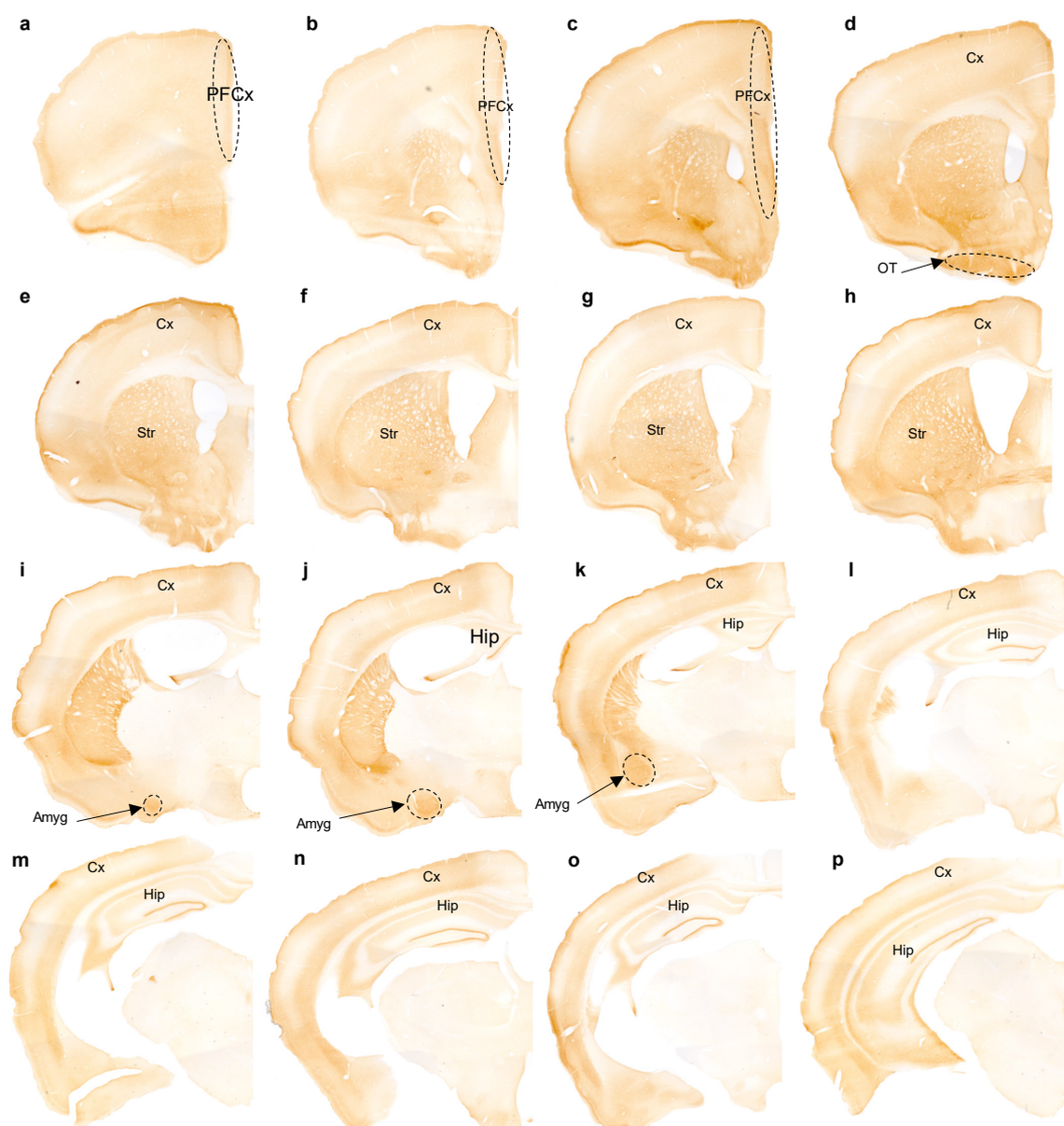


Figure 11. CB₁ receptor immunostaining in the Glu-CB₁-RS mouse brain.

An immunoperoxidase method for light microscopy was used for CB₁ receptor immunostaining detection in the Glu-CB₁-RS mouse brain. Representative rostro-caudal coronal brain sections stained with the CB₁ antibody (left to right and top to bottom). Observe a diffuse CB₁ receptor staining throughout but somehow more detectable in OT, Cx, Str, Amyg and hippocampus.

PFCx, Prefrontal cortex; OT, Olfactory tubercle; Cx, Cortex; Str, Striatum; Amyg, Amygdala; Hip, Hippocampus; a to d: Bregma 2.34 to 1.34 mm; e to h: Bregma 0.98 to 0.14 mm; i to l: Bregma -0.46 to -1.70 mm; m to p: Bregma -2.46 to -2.80 mm.

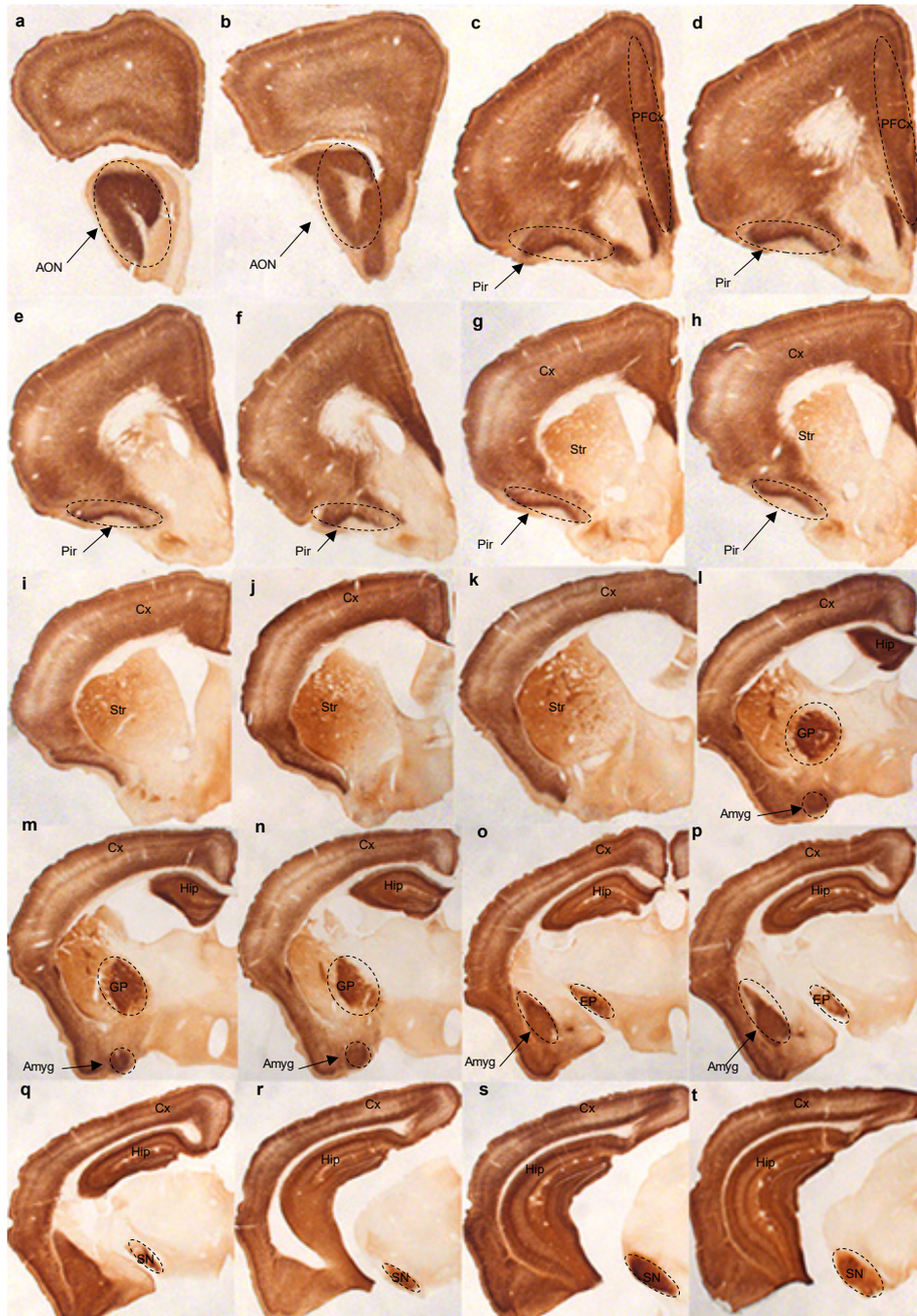


Figure 12. CB₁ receptor immunostaining in the GABA-CB₁-RS mouse brain.

An immunoperoxidase method for light microscopy was used for CB₁ receptor immunostaining detection in the GABA-CB₁-RS mouse brain. Representative rostro-caudal coronal brain sections (left to right and top to bottom). Strong CB₁ receptor immunostaining is found in AON, Pir, PFCx, Cx, Str, GP, EP, Amyg, SN and hippocampus.

AON, Anterior olfactory nucleus; Pir, Piriform cortex; PFCx, Prefrontal cortex; Cx, Cortex; Str, Striatum; GP, Globus pallidus; Amyg, Amygdala; Hip, Hippocampus; EP, Entopeduncular nucleus; SN, Substantia nigra; a to d: Bregma 2.80 to 1.70 mm; e to h: Bregma 1.54 to 1.18 mm; i to l: Bregma 0.74 to -0.94 mm; m to p: Bregma -1.22 to -1.58 mm; q to t: Bregma -2.46 to -2.92.

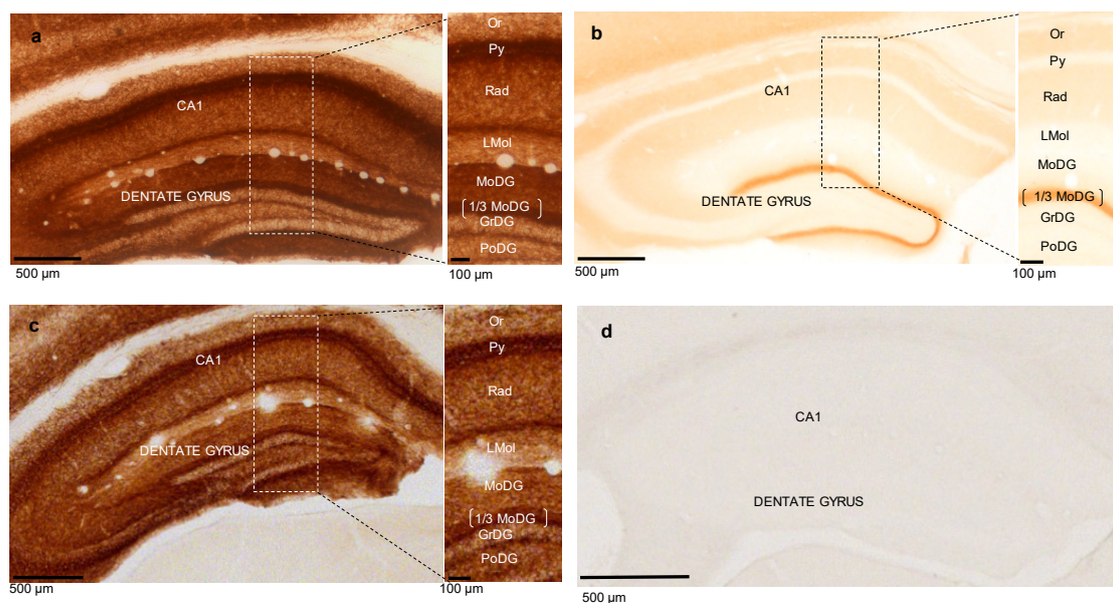


Figure 13. CB₁ receptor immunostaining in the hippocampus of CB₁-WT, Glu-CB₁-RS and GABA-CB₁-RS and CB₁-KO mice.

An immunoperoxidase method for light microscopy was used. In the CB₁-WT hippocampus a stronger immunolabeling is observed in Py and at the limit Rad and LMol than throughout the Or and Rad. In the dentate gyrus, the densest immunolabeling is found in the inner third of MoDG whereas less staining, although intense, was found in the outer parts of this layer. Fibrous processes surrounding the granule cell bodies are observed in GrDG (a). Detailed magnification of the Glu-CB₁-RS hippocampus: noticeable CB₁ receptor neuropil labeling is detected in Or and Rad. A dense band of immunostaining is observed in the inner third of MoDG (b). Detailed magnification of the GABA-CB₁-RS hippocampus: a denser labeling area is seen in Py and also at the limit between Rad and LMol. MoDG is heavily CB₁ receptor immunoreactive exhibiting a conspicuous band at its inner third (c). Detailed magnification of the CB₁-KO hippocampus: immunostaining completely disappears in all the CB₁-KO layers demonstrating that the CB₁ antibody is highly specific (d).

CA1, Region 1 of Cornu Ammonis; Or, Stratum oriens; Py, Pyramidal cell hippocampus; Rad, Stratum radiatum; LMol, Lacunosum moleculare; MoDG, Molecular dentate gyrus; 1/3 MoDG, inner third of molecular dentate gyrus; GrDG, Granular dentate gyrus; PoDG, Polymorph dentate gyrus.

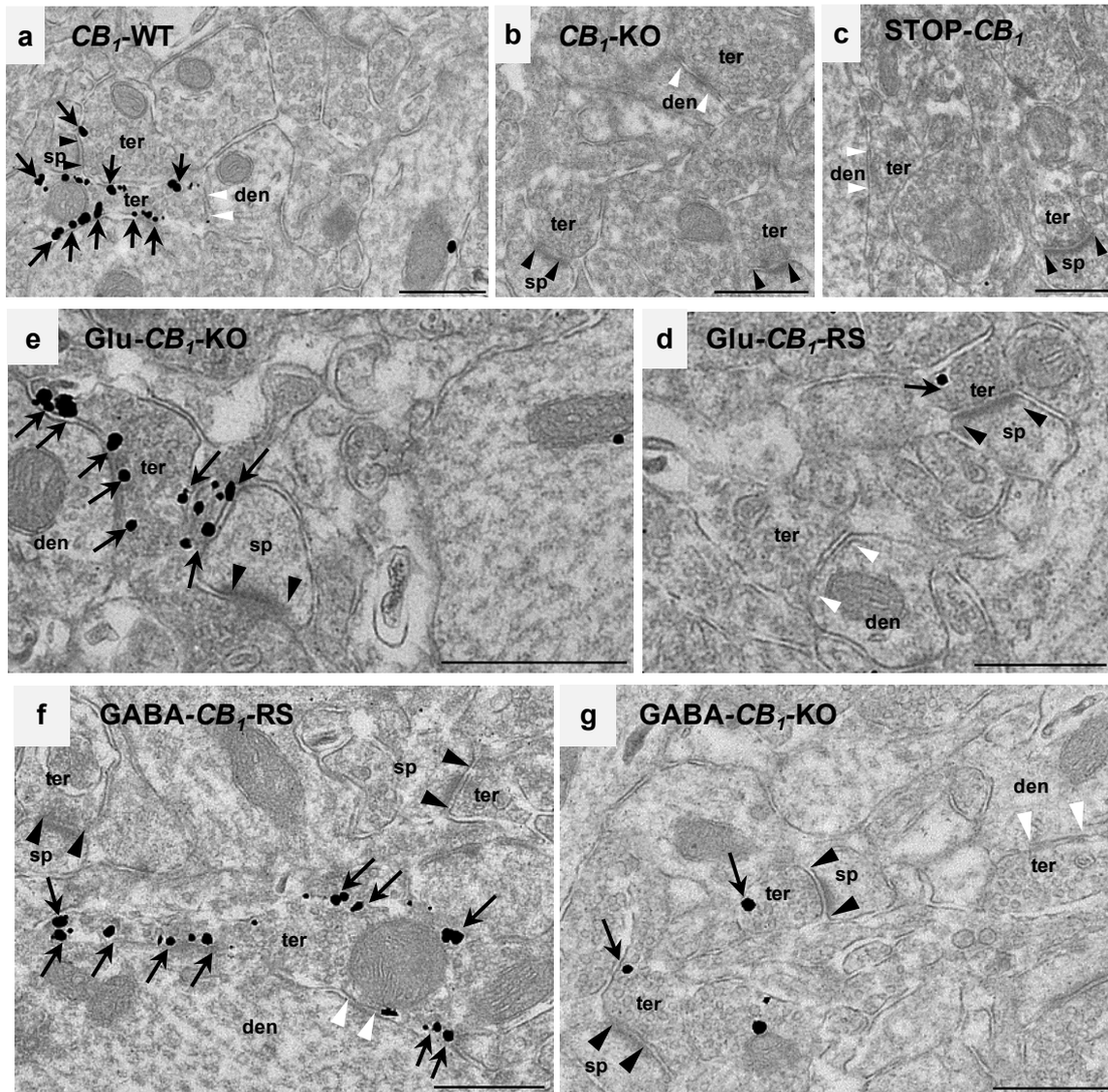


Figure 14. Subcellular CB₁ receptor localization in synaptic terminals of the CA1 stratum radiatum. Preembedding immunogold method for electron microscopy.

In *CB₁-WT*, immunoparticles are localized on excitatory (asymmetric synapses) and inhibitory (symmetric synapses) terminal membranes making synaptic contacts with dendritic spines or dendrites, respectively (**a**). No CB₁ receptor immunolabeling is detected in *CB₁-KO*, indicating that the CB₁ receptor antibody used is specific (**b**). In *STOP-CB₁*, no CB₁ receptor labeling is observed in excitatory or inhibitory terminals (**c**). In *Glu-CB₁-RS*, CB₁ receptor immunoparticles are at asymmetric but not symmetric synapses (**d**). On the contrary, CB₁ receptor immunolabeling is distributed on inhibitory presynaptic terminal membranes and not on membranes of excitatory boutons in *Glu-CB₁-KO* (**e**). Similarly, many CB₁ receptor immunoparticles are confined to symmetric synapses in *GABA-CB₁-RS* (**f**). However, synaptic CB₁ receptor immunoparticles are observed on excitatory presynaptic terminal membranes and not in inhibitory terminals of *GABA-CB₁-KO* (**g**). Note the presence of CB₁ receptor labeling in mitochondria (a, d-g).

Black arrowheads: excitatory synapses; white arrowheads: inhibitory synapses; black arrows: CB₁ receptor immunoparticles. ter, terminal; den, dendrite; sp, dendritic spine. Scale bars: 0.5 μm.

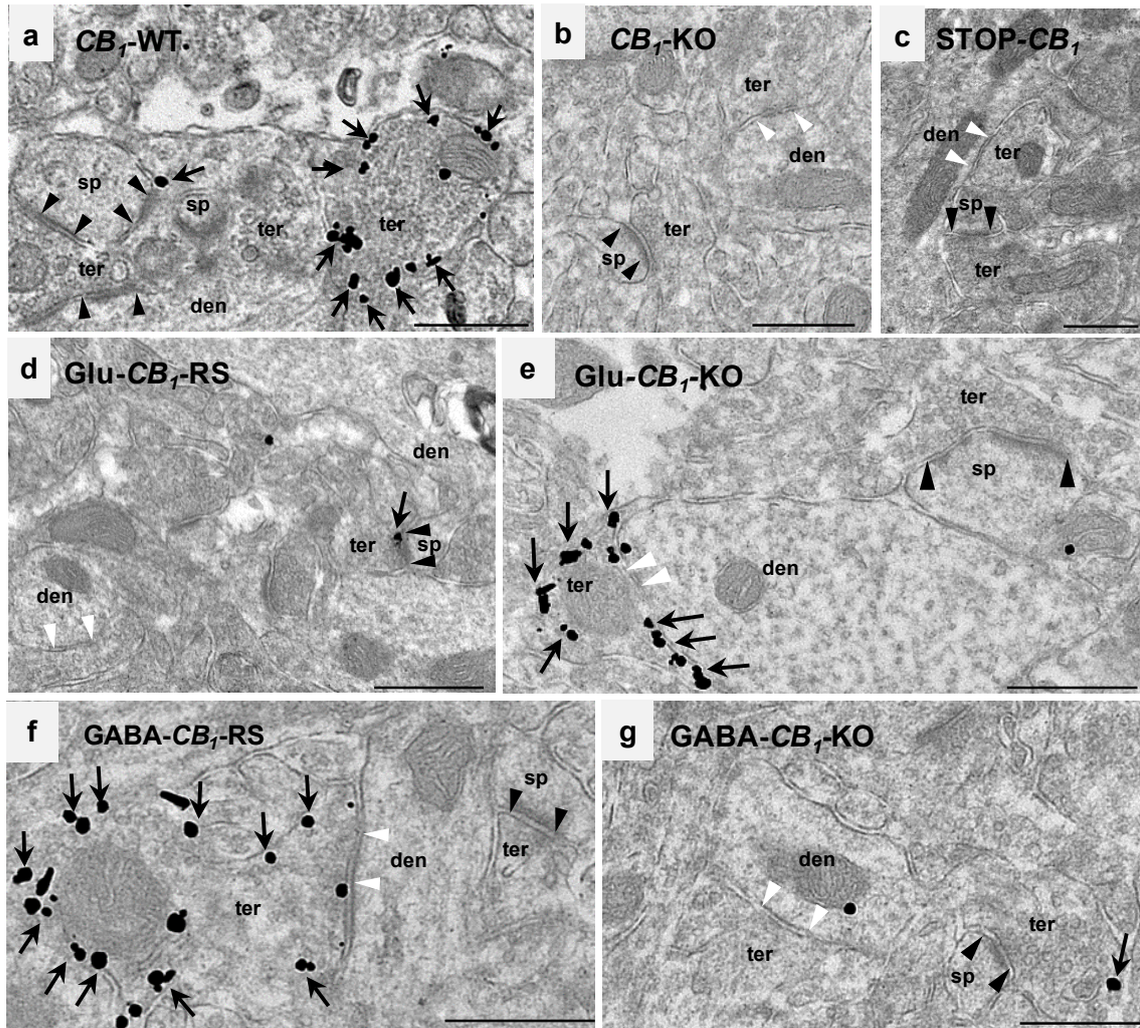


Figure 15. Subcellular CB₁ receptor localization in synaptic terminals of the dentate inner 1/3 molecular layer. Preembedding immunogold method for electron microscopy.

In *CB₁-WT*, scarce CB₁ receptor immunoparticles are localized on presynaptic terminal membranes making excitatory synapses (asymmetric) with dendritic spines while numerous metal particles are at inhibitory terminals (a). The CB₁ receptor labeling pattern is neither observed in *CB₁-KO* (b) nor in *STOP-CB₁* mice (c). In *Glu-CB₁-RS*, CB₁ receptor gold particles are only seen on presynaptic mossy cell axon boutons and not on presynaptic inhibitory terminal membranes (d). However, in *Glu-CB₁-KO*, excitatory mossy cell terminals are immunonegative but inhibitory boutons contain abundant CB₁ receptor labeling (e). In *GABA-CB₁-RS*, numerous CB₁ receptor immunoparticles are only on presynaptic inhibitory axon terminals but not at excitatory synapses (f). In contrast, immunolabeling is not observed in GABAergic terminals but it is present in excitatory boutons of *GABA-CB₁-KO* (g). Observe CB₁ receptor immunoparticles on mitochondria (a, d-g).

Black arrowheads: excitatory synapses; white arrowheads: inhibitory synapses; black arrows: CB₁ receptor immunoparticles. ter, terminal; den, dendrite; sp, dendritic spine. Scale bars: 0.5 μm.

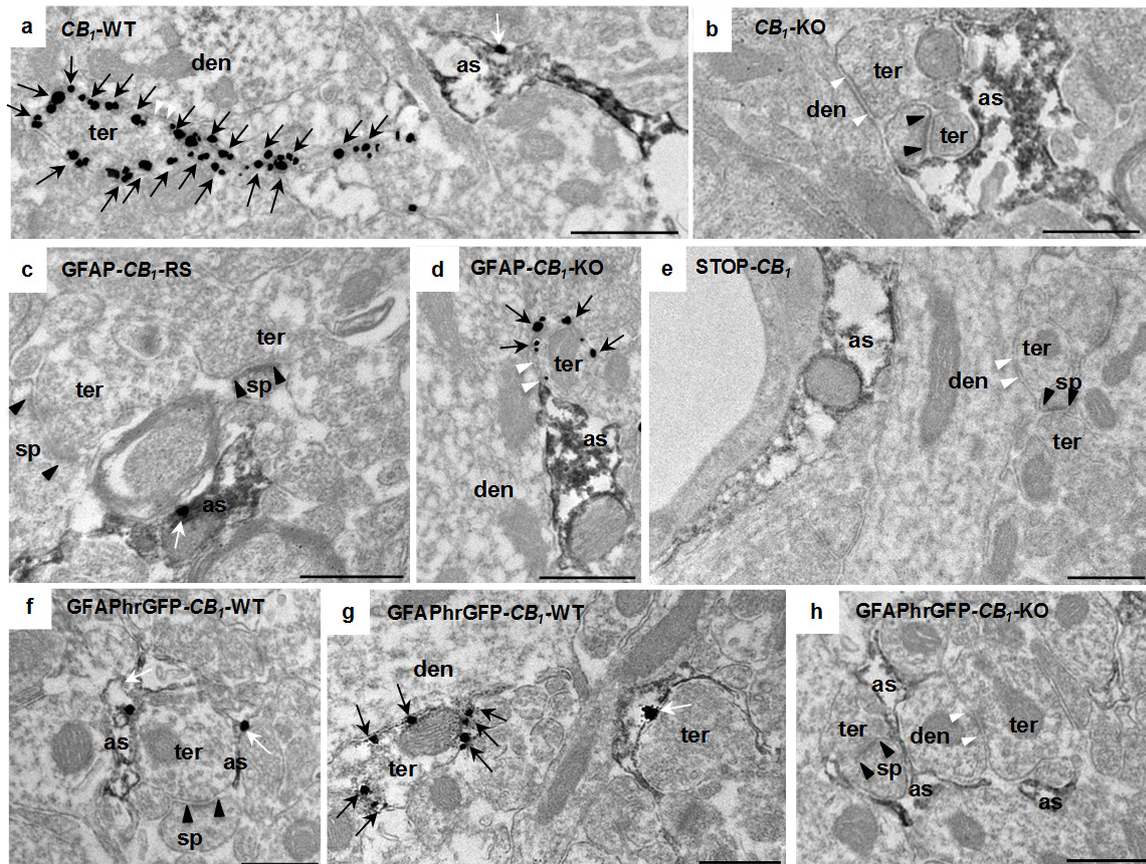


Figure 16. Subcellular CB₁ receptor localization in astrocytic processes of the hippocampal CA1 stratum radiatum. Combined preembedding immunogold and immunoperoxidase methods for electron microscopy.

In *CB₁-WT* and in *GFAPhrGFP-CB₁-WT* mice, CB₁ receptor immunoparticles are localized in membranes of astrocytic processes. CB₁ receptor immunoparticles are also observed on terminal membranes (**a, f, g**). CB₁ receptor immunolabeling is neither detected in *CB₁-KO* nor in *GFAPhrGFP-CB₁-KO*, indicating that the CB₁ receptor antibody used is highly specific (**b, h**). CB₁ receptor immunoparticles appear in astrocytic processes but not in terminals of *GFAP-CB₁-RS* (**c**). In *GFAP-CB₁-KO*, CB₁ receptors disappear from membranes of astrocytic processes whereas they remain in synaptic terminals (**d**). There is not CB₁ receptor immunoparticles in *STOP-CB₁* mice (**e**).

Black arrowheads: excitatory synapses; white arrowheads: inhibitory synapses; black arrows: CB₁ receptor immunoparticles; white arrows: astrocytic CB₁ receptor immunoparticles. as: astrocytic processes; ter: terminal; den: dendrite; sp: dendritic spine. Scale bars: 0.5 μ m.

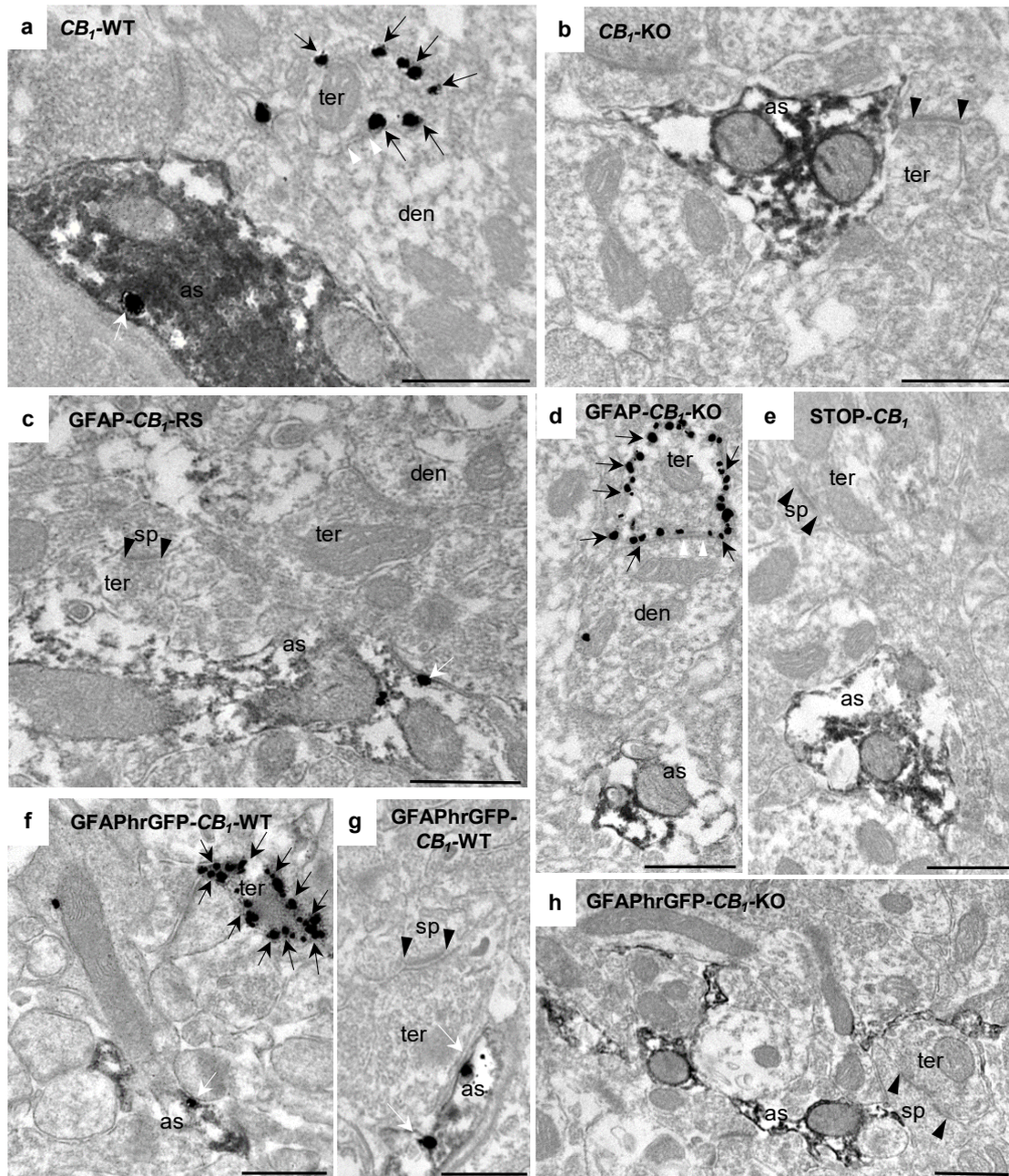


Figure 17. Subcellular CB₁ receptor localization in astrocytic processes of the dentate molecular layer. Combined preembedding immunogold and immunoperoxidase methods for electron microscopy.

In *CB₁-WT* and *GFAPhrGFP-CB₁-WT* mice, CB₁ receptor is localized on presynaptic terminal membranes and on membranes of astrocytic processes (**a**, **f**, **g**). The CB₁ receptor labeling pattern disappears in *CB₁-KO* and in *GFAPhrGFP-CB₁-KO* showing the specificity of the antibody used (**b**, **h**). In *GFAP-CB₁-RS*, CB₁ receptor immunogold particles are on astrocytic processes and not on neuronal terminal membranes (**c**). However, in *GFAP-CB₁-KO*, astrocytic processes are CB₁ receptor immunonegative but inhibitory and excitatory terminals contain the receptor (**d**). There is no CB₁ receptor immunolabeling in *STOP-CB₁* (**e**).

Black arrowheads: excitatory synapses; white arrowheads: inhibitory synapses; black arrows: CB₁ receptor immunoparticles; white arrows: astrocytic CB₁ receptor immunoparticles. as: astrocytic processes; ter: terminal; den: dendrite; sp: dendritic spine. Scale bars: 0.5 μm.

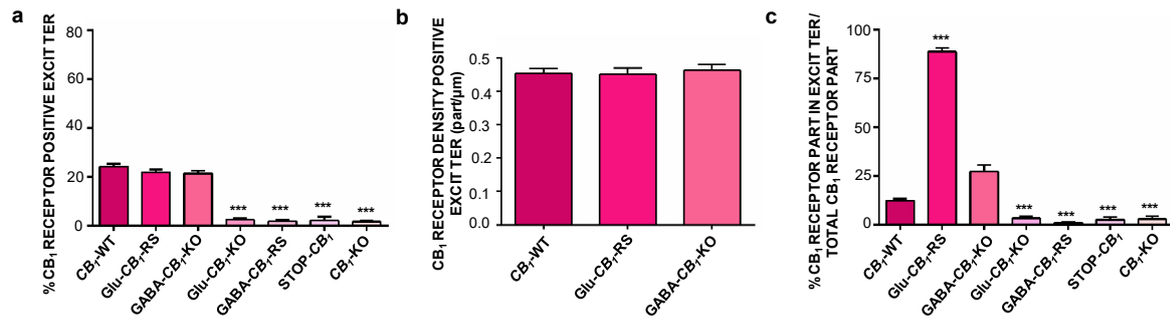


Figure 18. Statistics of the CB₁ receptor immunopositive glutamatergic terminals in the CA1 stratum radiatum.

No significant differences are found between *CB₁-WT* (24.29% ± 1.09), *Glu-CB₁-RS* (21.89% ± 1.21) and *GABA-CB₁-KO* (21.37% ± 1.13). Just residual excitatory terminals with metal particles were found in *Glu-CB₁-KO* (2.50% ± 0.76), *GABA-CB₁-RS* (1.92% ± 0.66) *STOP-CB₁* (2.20% ± 1.54) and *CB₁-KO* (1.68% ± 0.62) (a). Graphical representation of the CB₁ receptor density (particles/μm) in CB₁ receptor positive excitatory terminals. No statistical differences are obtained between *CB₁-WT* (0.45 ± 0.01), *Glu-CB₁-RS* (0.45 ± 0.02) and *GABA-CB₁-KO* (0.46 ± 0.02) (b). Proportion of CB₁ receptor in glutamatergic terminals normalized to the total CB₁ receptor signal in plasmalemma of cellular structures: 12.35% ± 1.04 of the total immunoparticles are located in glutamatergic terminals of *CB₁-WT* and 27.30% ± 3.34 of *GABA-CB₁-KO*. In *Glu-CB₁-RS* the proportion increases to 88.78% ± 1.96 and only residual CB₁ receptor immunoparticles are found in excitatory terminals of *GABA-CB₁-RS* (1.10% ± 0.30), *Glu-CB₁-KO* (3.39% ± 0.96), *STOP-CB₁* (3.03% ± 1.52) and *CB₁-KO* (2.65% ± 1.45) (c). Data are expressed as mean ± Standard Error of the Mean of three different animals. Data were analyzed by means of Kruskal-Wallis Test and the Dunn's Multiple Comparison Post-hoc test. *** indicate statistically significant differences with $p < 0.001$. EXCIT, excitatory; PART, particles; TER, terminals.

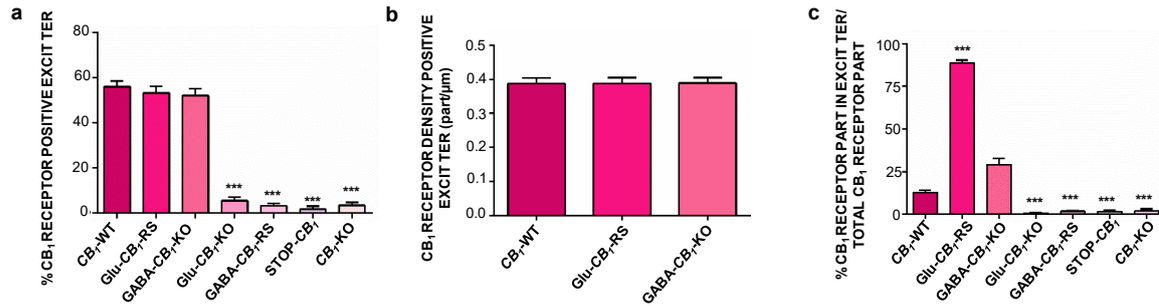


Figure 19. Statistics of the CB₁ receptor distribution in glutamatergic terminals of the dentate inner 1/3 molecular layer.

The percentage of CB₁ receptor immunopositive excitatory synaptic terminals is statistically similar between CB₁-WT (55.98% ± 2.51), Glu-CB₁-RS (53.19% ± 2.89) and GABA-CB₁-KO (52.12% ± 3.00). The terminal labeling drastically decreases in Glu-CB₁-KO (5.48% ± 1.70) and practically disappears in GABA-CB₁-RS (3.19% ± 1.08), STOP-CB₁ (1.69% ± 1.36) and CB₁-KO (3.41% ± 1.36) (a). Graphical representation of the CB₁ receptor density (particles/μm) in CB₁ receptor positive excitatory terminals. Density in CB₁-WT (0.39 ± 0.02), Glu-CB₁-RS (0.39 ± 0.02) and GABA-CB₁-KO (0.39 ± 0.02) is not statistically different (b). Proportion of CB₁ receptors in glutamatergic terminals normalized to the total CB₁ receptor signal in plasmalemma of cellular structures: 12.80% ± 1.39 of the total CB₁ receptor immunoparticles in CB₁-WT, 29.22% ± 3.65 in GABA-CB₁-KO and 88.64% ± 1.71 in Glu-CB₁-RS are localized in excitatory terminals. Only scarce CB₁ receptor immunoparticles are in excitatory terminals of GABA-CB₁-RS (1.78% ± 0.45), Glu-CB₁-KO (0.65% ± 0.27), STOP-CB₁ (1.54% ± 1.10) and CB₁-KO (1.87% ± 1.14) (c). Data are expressed as mean ± Standard Error of the Mean of three different animals. Data were analyzed by means of Kruskal-Wallis Test and the Dunn's Multiple Comparison Post-hoc test. *** indicate statistically significant differences with $p < 0.001$. EXCIT, excitatory; PART, particles; TER, terminals.

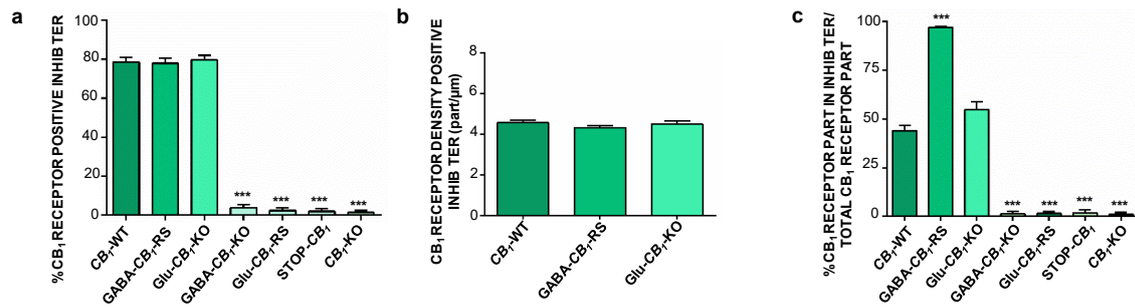


Figure 20. Statistics of the CB₁ receptor distribution in GABAergic terminals of the CA1 stratum radiatum.

Statistical analysis indicates that the percentages of CB₁ receptor immunopositive inhibitory synaptic terminals in CB₁-WT (78.50% ± 2.44), GABA-CB₁-RS (77.92% ± 2.63) and Glu-CB₁-KO (79.63% ± 2.35) are not different. The values virtually disappear in GABA-CB₁-KO (2.71% ± 1.20), Glu-CB₁-RS (2.38% ± 1.37), STOP-CB₁ (1.95% ± 1.44) and CB₁-KO (1.53% ± 0.90) (a). Graphical representation of the CB₁ receptor density (particles/μm) in CB₁ receptor positive inhibitory terminals. There are no statistical differences between CB₁-WT (4.57 ± 0.13), GABA-CB₁-RS (4.33 ± 0.11) and Glu-CB₁-KO (4.50 ± 0.15) (b). Proportion of CB₁ receptor in GABAergic terminals normalized to the total CB₁ receptor signal in plasmalemma of cellular structures: 44.06% ± 2.67 of CB₁ receptor immunoparticles in CB₁-WT, 54.87% ± 3.94 in Glu-CB₁-KO and 97.10% ± 0.46 in GABA-CB₁-RS are in inhibitory terminals. Only sparse CB₁ receptor immunoparticles are in inhibitory terminals of Glu-CB₁-RS (1.60% ± 0.92), GABA-CB₁-KO (1.35% ± 1.35), STOP-CB₁ (1.97% ± 1.46) and CB₁-KO (1.18% ± 0.83) (c). Data are expressed as mean ± Standard Error of the Mean of three different animals. Data were analyzed by means of Kruskal-Wallis Test and the Dunn's Multiple Comparison Post-hoc test. *** indicate statistically significant differences with $p < 0.001$. INHIB, inhibitory; PART, particles; TER, terminals.

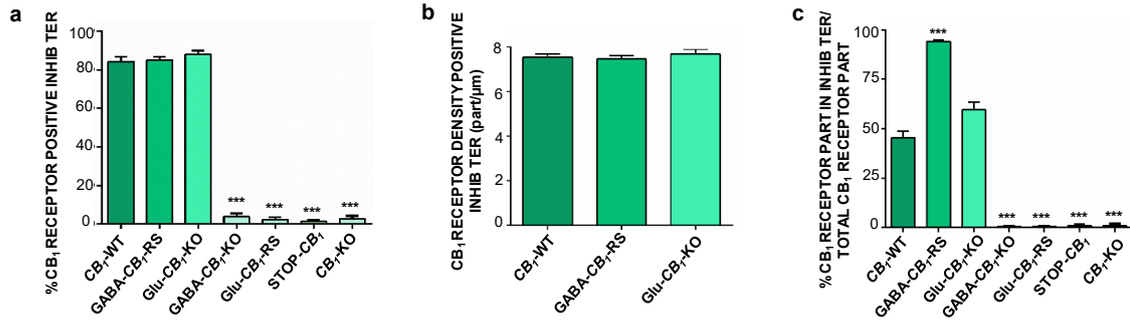


Figure 21. Statistics of the CB₁ receptor distribution in GABAergic terminals of the dentate inner 1/3 molecular layer.

Percentages of CB₁ receptor immunopositive inhibitory synaptic terminals do not defer statistically between CB₁-WT (84.27% ± 2.38), GABA-CB₁-RS (85.07% ± 1.76) and Glu-CB₁-KO (88.02% ± 1.93). Only residual particles are detected in scarce terminals of GABA-CB₁-KO (3.82% ± 1.68), Glu-CB₁-RS (2.30% ± 1.32), STOP-CB₁ (1.28% ± 0.90) and CB₁-KO (2.84% ± 1.48) (a). Graphical representation of the CB₁ receptor density (particles/μm) in CB₁ receptor positive inhibitory terminals. Density in CB₁-WT (7.54 ± 0.16), GABA-CB₁-RS (7.47 ± 0.14) and Glu-CB₁-KO (7.69 ± 0.21) is not statistically different (b). Proportion of CB₁ receptor immunoparticles in GABAergic terminals normalized to the total CB₁ signal in plasmalemma of cellular structures: 45.38% ± 3.18 in CB₁-WT, 59.45% ± 4.00 in Glu-CB₁-KO and 93.87% ± 0.93 in GABA-CB₁-RS are distributed in GABAergic terminals. Rare CB₁ receptor immunoparticles are in inhibitory terminals of Glu-CB₁-RS (0.60% ± 0.43), GABA-CB₁-KO (0.53% ± 0.53), STOP-CB₁ (0.86% ± 0.86) and CB₁-KO (0.93% ± 0.93) (c). Data are expressed as mean ± Standard Error of the Mean of three different animals. Data were analyzed by means of Kruskal-Wallis Test and the Dunn's Multiple Comparison Post-hoc test. *** indicate statistically significant differences with $p < 0.001$. INHIB, inhibitory; PART, particles; TER, terminals.

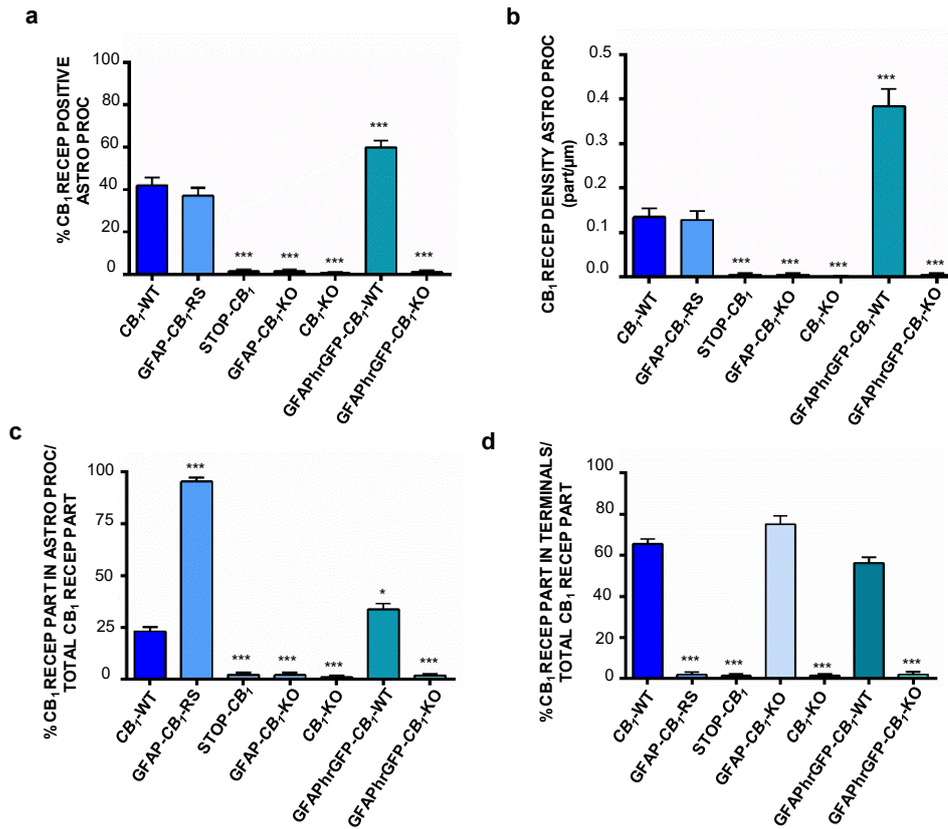


Figure 22. Statistics of the CB₁ receptor distribution in astrocytic processes of the CA1 stratum radiatum.

Percentages of CB₁ receptor immunopositive astrocytes do not show statistical differences between *CB₁-WT* mice ($42.06\% \pm 3.56$) and *GFAP-CB₁-RS* mice ($37.12\% \pm 3.79$). In *GFAPhrGFP-CB₁-WT*, proportion increases to $59.91\% \pm 3.29$. Residual immunolabeling is found in *GFAP-CB₁-KO* ($1.46\% \pm 0.78$), *STOP-CB₁* ($1.15\% \pm 0.67$), *CB₁-KO* ($0.54\% \pm 0.39$) and *GFAPhrGFP-CB₁-KO* ($1.60\% \pm 0.66$) (a). CB₁ immunoparticle density on membranes of astrocytic processes (particles / μm) are analyzed. *CB₁-WT* (0.135 ± 0.019) and *GFAP-CB₁-RS* (0.128 ± 0.020) are statistically similar, whereas a significant increase is found in *GFAPhrGFP-CB₁-WT* (0.384 ± 0.039). Just unspecific particles are observed in *GFAP-CB₁-KO* (0.005 ± 0.003), *STOP-CB₁* (0.005 ± 0.003), *CB₁-KO* (0.001 ± 0.001) and *GFAPhrGFP-CB₁-KO* (0.004 ± 0.002) (b). Proportion of CB₁ gold particles in astrocytic processes versus total CB₁ expression in plasmalemma of cellular structures: $95.45\% \pm 1.82$ of the total CB₁ receptor immunoparticles are located in astrocytic processes of *GFAP-CB₁-RS* and $23.08\% \pm 1.99$ are in *CB₁-WT* mice. In *GFAPhrGFP-CB₁-WT*, the proportion increases to $33.71\% \pm 2.75$. Only residual CB₁ immunoparticles are located in astrocytic processes of *GFAP-CB₁-KO* ($1.94\% \pm 1.27$), *STOP-CB₁* ($1.96\% \pm 1.28$), *GFAPhrGFP-CB₁-KO* ($1.62\% \pm 0.94$) and *CB₁-KO* mice ($1.01\% \pm 0.71$) (c). Proportion of immunogold particles localized in terminals versus total CB₁ receptor expression in plasmalemma: $75.13\% \pm 4.06$ of the total CB₁ immunoparticles are in terminals of *GFAP-CB₁-KO* and $65.52\% \pm 2.44$ in *CB₁-WT*. In *GFAPhrGFP-CB₁-WT*, the proportion is $56.32\% \pm 2.73$. Residual CB₁ receptor immunoparticles are located in astrocytic processes of *GFAP-CB₁-RS* ($2.02\% \pm 1.17$), *STOP-CB₁* ($1.47\% \pm 0.84$), *GFAPhrGFP-CB₁-KO* ($2.08\% \pm 1.19$) and *CB₁-KO* ($1.51\% \pm 0.86$) mice (d). Data are expressed as mean \pm Standard Error of the Mean of three different animals. Data were analyzed by means of Kruskal-Wallis Test and the Dunn's Multiple Comparison Post-hoc test. *** indicate statistically significant differences with $p < 0.001$; ** indicate statistically significant differences with $p < 0.01$; * indicate statistically significant differences with $p < 0.05$.

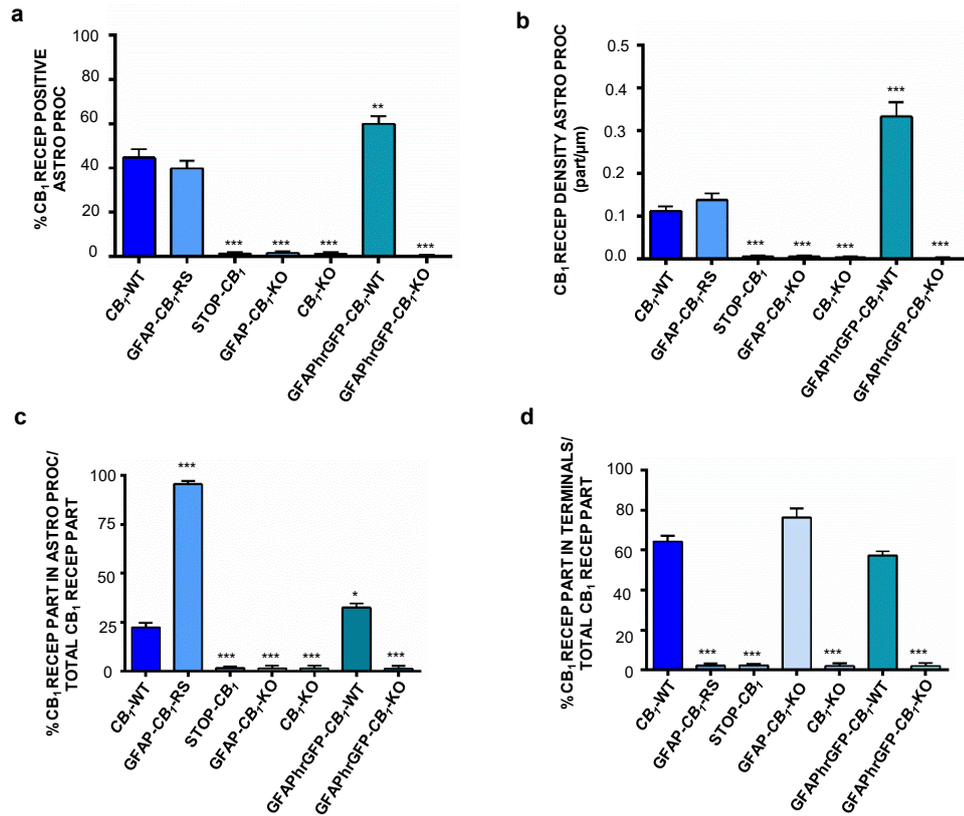


Figure 23. Statistics of the CB₁ receptor distribution in astrocytic processes of the dentate molecular layer.

Similar percentages of CB₁ receptor immunopositive astrocytic processes in CB₁-WT (44.67% ± 3.85) and GFAP-CB₁-RS (39.84% ± 3.50) are found. Statistical differences are obtained with GFAPhrGFP-CB₁-WT (59.99% ± 3.37). Proportion decreases to reach just residual level in: GFAP-CB₁-KO (1.56% ± 0.65), STOP-CB₁ (1.33% ± 0.64), GFAPhrGFP-CB₁-KO (0.47% ± 0.36) and CB₁-KO mice (1.19% ± 0.70) (a). Analysis of CB₁ receptor density (particles/μm) on astrocytic processes shows no statistical differences comparing: CB₁-WT (0.112 ± 0.011) and GFAP-CB₁-RS (0.138 ± 0.016) mice. However, statistical differences are found with GFAPhrGFP-CB₁-WT (0.334 ± 0.033). Just residual particles are observed in: GFAP-CB₁-KO (0.006 ± 0.003), STOP-CB₁ (0.006 ± 0.003), GFAPhrGFP-CB₁-KO (0.002 ± 0.002) and CB₁-KO (0.004 ± 0.002) (b). Proportion of CB₁ gold particles in astrocytic processes versus total CB₁ expression in plasmalemma is 22.40% ± 2.28 in CB₁-WT, 95.61% ± 1.56 in GFAP-CB₁-RS and 32.57% ± 2.09 in GFAPhrGFP-CB₁-WT. Only non-specific CB₁ receptor immunoparticles are found in astrocytic processes of GFAP-CB₁-KO (1.43% ± 1.43), STOP-CB₁ (1.65% ± 0.66) GFAPhrGFP-CB₁-KO (1.37% ± 1.37) and CB₁-KO (1.43% ± 1.43) mice (c). Proportion of immunogold particles localized in terminals versus total CB₁ receptor expression in plasmalemma: 76.17% ± 4.70 of the total CB₁ immunoparticles are in terminals of GFAP-CB₁-KO, 64.27% ± 2.88 in CB₁-WT and 57.17% ± 2.19 in GFAPhrGFP-CB₁-WT. Residual CB₁ receptor immunoparticles are located in astrocytic processes of GFAP-CB₁-RS (2.19% ± 1.10), STOP-CB₁ (2.36% ± 0.85), GFAPhrGFP-CB₁-KO (2.05% ± 1.52) and CB₁-KO (2.14% ± 1.52) mice (d). Data are expressed as mean ± Standard Error of the Mean of three different animals. Data were analyzed by means of Kruskal-Wallis Test and the Dunn's Multiple Comparison Post-hoc test. *** indicate statistically significant differences with $p < 0.001$; ** indicate statistically significant differences with $p < 0.01$; * indicate statistically significant differences with $p < 0.05$.

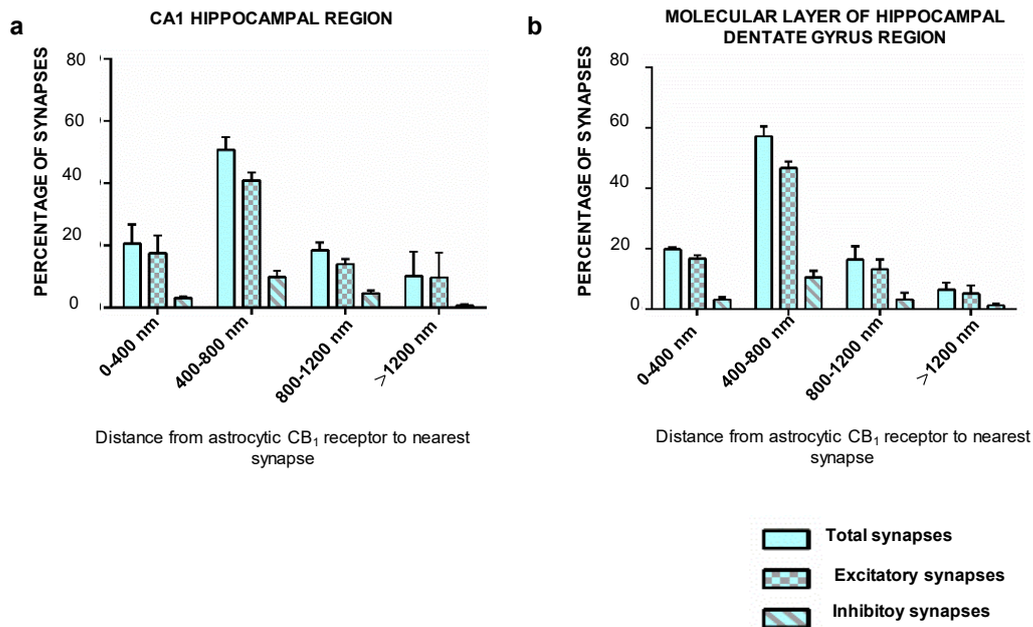


Figure 24. Statistics of distances from the astrocytic CB₁ receptors to the nearest synapse in the hippocampus.

Analysis of the distance from the astrocytic CB₁ receptors and the midpoint of the nearest synapse surrounded by the astrocytic processes in the GFAPhrGFP-CB₁-WT mice. In CA1, of the total analyzed synapses (n= 159) 50.76% ± 4.10 are localized at a distance between 400-800 nm from the astrocytic CB₁ receptor immunoparticle. 40.89% ± 2.57 of them are excitatory and 9.86% ± 1.97 are inhibitory synapses. 20.60% ± 6.12 of the synapses are in a range of 0-400 nm from the astrocytic CB₁ receptor immunoparticle. Of those, 17.52% ± 5.63 are excitatory synapses and 3.08% ± 0.49 are inhibitory synapses. 18.46% ± 2.53 of the total synapses are between 800 nm and 1200 nm; 13.96% ± 1.60 are excitatory and 4.50% ± 0.95 are inhibitory synapses. Only 10.18% ± 7.81 of the synapses are found at more than 1200 nm from the astrocytic CB₁ receptor. Of those, 9.62% ± 8.01 are excitatory and only 0.56% ± 0.56 are inhibitory synapses (**a**). In the dentate molecular layer, 57.25% ± 3.19 of the total analyzed synapses (n=166) are placed at a distance between 400-800 nm from the astrocytic CB₁ receptor particle; 46.67% ± 2.17 of them are excitatory and 10.59% ± 2.18 are inhibitory synapses. 19.83% ± 0.58 of the synapses are in a range of 0-400 nm; 16.67% ± 1.31 of them are excitatory and 3.16% ± 0.82 are inhibitory synapses. 16.42% ± 4.37 of the total synapses are located between 800-1200 nm. Of these synapses, 13.22% ± 3.19 are excitatory and 3.21% ± 2.52 are inhibitory. Only 6.48% ± 2.31 of the synapses are observed at more than 1200 nm from the astrocytic CB₁ receptor metal particle; 5.16% ± 2.72 being excitatory and 1.32% ± 0.66 inhibitory synapses (**b**). Data are expressed as mean ± Standard Error of the Mean of three different animals.

6 DISCUSSION

6.1 THE IMPORTANCE OF CB₁ RECEPTOR MUTANTS IN THE STUDY OF CB₁ RECEPTOR

The CB₁ receptor is expressed in different brain cell types populations (Marsicano and Lutz, 1999; Tsou et al., 1999; Nyíri et al., 2005b; Monory et al., 2006; Häring et al., 2007; Scavone et al., 2010; Han et al., 2012; Metna-Laurent and Marsicano, 2015). Moreover, a wide variety of intracellular effects such as modulation of kinases, ion channels and transcription factors are triggered by the activation of CB₁ receptors (Bosier et al., 2010; Pertwee, 2015). One of the most known and important result of these intracellular events is the retrograde inhibition of transmitter release (Kano et al., 2009). Consequently, the activation of CB₁ receptors modulates the release of several neurotransmitters, such as: glutamate, GABA, glycine, acetylcholine, norepinephrine, dopamine, serotonin and cholecystokinin (Kano et al., 2009).

Anatomically, the CB₁ receptor is also widely distributed in the brain with a preferential localization in motor, limbic, reward and cortical regions (Matsuda et al., 1993; Tsou et al., 1998). Its high concentration in certain brain areas is an advantage for studying the functional role of the receptor in the neural circuits where it is abundantly localized (Katona et al., 1999; Kawamura et al., 2006; Ludányi et al., 2008; Marsicano and Kuner, 2008; Katona and Freund, 2012; De-May and Ali, 2013; Steindel et al., 2013; Hu and Mackie, 2015).

However, CB₁ receptor density is not uniform through the regions expressing the receptor which makes extremely difficult to identify low CB₁ receptor expression in cell types and/or in subcellular compartments of wild-type brains (Busquets-Garcia et al., 2015).

The hippocampus is one of the brain structures with the highest CB₁ receptor immunoreactivity (Herkenham et al., 1990; Mailloux and Vanderhaeghen, 1992; Matsuda et al., 1993; Tsou et al., 1998; Marsicano and Lutz, 1999; Egertová and Elphick, 2000; Katona et al., 2006; Kawamura et al., 2006; Ludányi et al., 2008; Katona and Freund, 2012; Steindel et al., 2013; Hu and Mackie, 2015) and where CB₁ receptors have shown to play key functional roles (Campbell et al., 1986; Heyser et al., 1993; Stella et al., 1997; Hampson and Deadwyler, 1999; Katona et al., 2000; Carlson et al., 2002; Chevaleyre and Castillo, 2003, 2004; Vanderbyl et al., 2005; Robbe et al., 2006; Akirav, 2011; Puighermanal et al., 2012; Han et al., 2012; Basavarajappa and Subbanna, 2014).

The development and use of cell type specific CB₁-receptor-knockout mice, lacking the CB₁ receptor in specific brain cell populations served to identify low CB₁ receptor expression in cellular, subcellular or intracellular compartments as well as to understand its physiological functions in those compartments (Marsicano et al., 2003; Monory et al., 2006, 2007; Puighermanal et al., 2009; Bellocchio et al., 2010; Bénard et al., 2012; Han et al., 2012; Ruehle et al., 2013; Steindel et al., 2013; Soria-Gómez et al., 2014; Busquets-Garcia et al., 2015; Martín-García et al., 2015; Oliveira da Cruz et al., 2016). For instance, in the GABA-CB₁-KO brain, a drastic decrease of the CB₁ receptor immunoreactivity was observed throughout the hippocampus but remaining a noticeable immunoreactive band in the inner third of the dentate molecular layer (Martín-García et al., 2015). On the other hand, just a very faint decrease of the CB₁ receptor immunostaining was noticed in the Glu-

CB₁-KO hippocampus (Martín-García et al., 2015). Yet, there was not a full disappearance of the *CB₁* receptor immunoreactivity in either condition as it occurred in *CB₁*-KO. The reason is because the *CB₁* receptor localization is mostly, but not exclusively, restricted to neuronal membrane compartments, as *CB₁* receptors are also distributed in astrocytic processes (Bosier et al., 2013) and mitochondrial membranes of the hippocampus (Bénard et al., 2012; Hebert-Chatelain et al., 2014a; b). The light microscopic observations previously described by our group (Martín-García et al., 2015) correlate well with the present findings in the electron microscope of the subcellular *CB₁* receptor localization in hippocampal glutamatergic and GABAergic axon terminals of GABA-*CB₁*-KO and Glu-*CB₁*-KO, respectively. This work has also shown that *CB₁* receptor immunolabeling was abolished in excitatory terminals of Glu-*CB₁*-KO while the proportion and density of *CB₁* receptors in GABA-*CB₁*-KO were maintained in the glutamatergic excitatory terminals of CA1 stratum radiatum (Takahashi and Castillo, 2006; Takács et al., 2012; Witter, 2012). On the contrary, *CB₁* receptor metal particles were not observed in the hippocampal inhibitory synaptic terminals of GABA-*CB₁*-KO though *CB₁* receptor expression and distribution in CA1 and dentate inhibitory boutons were undistinguishable from *CB₁*-WT.

Moreover, the using of GFAP-*CB₁*-KO mice have already demonstrated that the activation of *CB₁* receptors in CA1 astrocytes mediates the impairment of working memory elicited by acute cannabinoids through the modulation of hippocampal long term depression (Han et al., 2012). The results obtained on the proportion of *CB₁* receptor immunopositive CA1 hippocampal astrocytes in in this thesis is similar to the value reported in our previous work (Han et al. in 2012). Furthermore, the expression levels of *CB₁* receptor immunoparticles in terminals of GFAP-*CB₁*-KO mice was comparable to *CB₁*-WT indicating that GFAP-*CB₁*-KO are good tools for studies based on the loss of function of astrocytic *CB₁* (Han et al., 2012).

However, conditional mutant mice have limitations as a biological compensation derived from the *CB₁* receptor deletion could occur. In addition, they do not allow establishing a link between the anatomical *CB₁* receptor localization in some nerve cells and a certain physiological task (Ruehle et al., 2013).

6.2 *CB₁* RECEPTOR EXPRESSION IN SPECIFIC BRAIN CELL TYPES OF RESCUE MUTANTS

The *CB₁* receptor expression protein restoration in phenotype-specific brain cell populations can provide great insights into the sufficiency of the *CB₁* receptor for specific brain functions and behaviors. Thus, it is crucial to ensure that the rescue of *CB₁* receptors in particular cell types of *CB₁* receptor-null mutants correlates with the endogenous *CB₁* receptor expression and localization in the wild-type mouse brain. This thesis work has analyzed in detail the anatomical distribution pattern of the *CB₁* receptor in the mutant mice.

6.2.1 CB₁ RECEPTOR EXPRESSION IN SPECIFIC NEURONAL CELL TYPES OF THE RESCUE MUTANT MICE

The cellular CB₁ receptor staining described here for the Glu-CB₁-RS and GABA-CB₁-RS mutants by a preembedding immunoperoxidase method for light microscopy reflects the distribution of CB₁ receptors throughout the brain (Herkenham et al., 1990; Tsou et al., 1998; Marsicano and Lutz, 1999; Egertová and Elphick, 2000; Mackie, 2005; Monory et al., 2006, 2007; Kano et al., 2009; Ruehle et al., 2013; Martín-García et al., 2015). CB₁ receptor immunoreactivity was much fainter in Glu-CB₁-RS than in GABA-CB₁-RS, as expected for the much lower CB₁ receptor expression in glutamatergic (Marsicano and Lutz, 1999; Katona et al., 2006; Kawamura et al., 2006; Monory et al., 2006, 2007; Martín-García et al., 2015; Lu and Mackie, 2016) than in GABAergic neurons (Katona et al., 1999; Hájos et al., 2000; Nyíri et al., 2005a; Kano et al., 2009; Takács et al., 2015; Lu and Mackie, 2016). Subcellularly, CB₁ receptors were localized in glutamatergic terminals of the CA1 stratum radiatum and the inner third of the dentate molecular layer of the Glu-CB₁-RS, the CB₁-WT and the GABA-CB₁-KO hippocampus. Furthermore, no significant differences in the proportion of immunopositive excitatory synaptic terminals and density of CB₁ receptors were detected between Glu-CB₁-RS, CB₁-WT or GABA-CB₁-KO (Gutiérrez-Rodríguez et al., 2016). This pattern corresponds well with the distribution and expression of CB₁ receptors in the intrahippocampal excitatory pathways (Marsicano and Lutz, 1999; Katona et al., 2006; Kawamura et al., 2006; Monory et al., 2006; Uchigashima et al., 2011; Katona and Freund, 2012). Very importantly, the inhibitory presynaptic profiles in the Glu-CB₁-RS hippocampus were virtually devoid of CB₁ immunoparticles. Also, the labeling was almost nil in the CB₁-KO, indicating that the CB₁ receptor antibody used in this study was highly specific. Previous electron microscopy studies described that more than 80% of excitatory synapses are CB₁ receptor immunopositive in the inner third of the dentate molecular layer, compared to only 30–50% in other hippocampal layers (Katona et al., 2006; Uchigashima et al., 2011). Although these values were somehow higher than those obtained in CB₁-WT and Glu-CB₁-RS in our study, it is plausible to assume that this is probably due to the different CB₁ receptor antibodies used and/or the immunocytochemical protocols applied (Katona et al., 2006; Uchigashima et al., 2011). Besides, in these studies only anatomical features of the synapses were used to distinguish between excitatory and inhibitory synapses. In our study, the CB₁ receptor localization was determined not only based on anatomical features but also using genetic tools (Glu-CB₁-RS mice and GABA-CB₁-RS mice) probably making a difference.

CB₁ receptors in GABA-CB₁-RS were densely localized in axon terminals of presumably cholecystokinin-containing interneurons as well as of calbindin D28k positive interneurons innervating the proximal portions of the pyramidal cell dendrites in the CA1 stratum radiatum and the dentate molecular layer (Katona et al., 1999; Marsicano and Lutz, 1999; Hájos et al., 2000; Nyíri et al., 2005; Kano et al., 2009; Katona and Freund, 2012; Lu and Mackie, 2016; Takács et al., 2015). Significantly, the excitatory presynaptic profiles in the GABA-CB₁-RS hippocampus were virtually lacking CB₁ immunoparticles. Furthermore, the proportion of the CB₁ receptor immunopositive inhibitory synaptic terminals was similar in GABA-CB₁-RS, Glu-CB₁-KO and CB₁-WT (Gutiérrez-Rodríguez et al., 2016), and the

localization coincides with the CB₁ receptor distribution pattern described in rodent interneurons (Katona et al., 1999; Hájos et al., 2000; Nyíri et al., 2005a) and human hippocampus (Katona et al., 2000).

Another parameter studied here was the density of immunogold particles in the CB₁ receptor positive excitatory and inhibitory terminals. The molecular and microscopic analysis of the hippocampal CB₁ receptors in Glu-CB₁-RS and GABA-CB₁-KO, on the one hand, and GABA-CB₁-RS and Glu-CB₁-KO, on the other, confirmed that the CB₁ receptor is present at much higher density in hippocampal GABAergic than glutamatergic cells (Katona et al., 2006; Kawamura et al., 2006; Gutiérrez-Rodríguez et al., 2016). The study by Kawamura et al. (2006) revealed a CB₁ receptor density of 5.75 particles/μm in inhibitory terminals and only 0.29 particles/μm in excitatory terminals of the CA1 stratum radiatum. In the same study, inhibitory terminals in the inner third of the dentate molecular layer had 7.14 particles/μm in contrast to 0.62 particles/μm found in excitatory terminals (Kawamura et al., 2006). The CB₁ receptor densities observed in the CB₁ receptor positive excitatory and inhibitory terminals of the mutant mice are in line with these previous hippocampal data obtained in CB₁-WT mice, though the differences observed could be attributed to the diverse antibodies and methods used, as mentioned before. To mention that higher receptor abundance in inhibitory synapses does not directly correlate with downstream signaling activation, as the glutamatergic CB₁ receptor is more efficiently coupled to G protein signaling than the GABAergic CB₁ receptor (Steindel et al., 2013).

6.2.2 CB₁ RECEPTOR EXPRESSION IN ASTROCYTES OF RESCUE MUTANT MICE

In this study, the subcellular CB₁ receptor labeling was also described in rescue mice where the CB₁ receptor expression has been specifically restored in astrocytes of GFAP-CB₁-RS mice. The low levels of expression of CB₁ receptors in astrocytes can only be accurately detected using high resolution immunocytochemical techniques for electron microscopy. Hence, a preembedding immunogold and immunoperoxidase methods applied to hippocampal sections had previously shown in our laboratory to be an excellent methodological strategy for the localization of CB₁ in astrocytes (Han et al., 2012; Bosier et al., 2013). Our present results have shown the proportion of the CB₁ receptor positive astrocytic processes and the density of the receptor in the astrocytic elements (particles /μm) were not statistically significant different in CB₁-WT and GFAP-CB₁-RS. Moreover, the percentage of the CB₁ receptor immunopositive CA1 astrocytes obtained here is similar to the previous values reported by our group (Han et al., 2012). The analysis of the CB₁ particles localized in astrocytic processes or terminals versus total CB₁ receptor particles in plasmalemmal structures of the GFAP-CB₁-RS hippocampus showed that almost all the labeling was in astrocytic processes and just only residual particles were in terminals confirming the high specificity of the astrocytic CB₁ receptor genetic rescue approach carried in the mutant mice. Consequently, these rescue mice emerge as excellent models to study the contribution to brain functions of the CB₁ receptors in astrocytes. Specifically, a more comprehensive characterization of the functional consequences of the eCB signaling through the astrocytic CB₁ receptor could be achieved in the tripartite synapses using these rescue mice.

Importantly, the CB₁ receptor expression virtually disappeared in the STOP-CB₁ mice corroborating previous findings published with these mutants (Ruehle et al., 2013; de Salas-Quiroga et al., 2015).

To sum up, the genetic rescue mouse models of the Glu-CB₁-RS expressed CB₁ receptors only in dorsal telencephalic glutamatergic neurons (Ruehle et al., 2013; Soria-Gómez et al., 2014; de Salas-Quiroga et al., 2015; Gutiérrez-Rodríguez et al., 2016), the GABA-CB₁-RS only in GABAergic neurons (de Salas-Quiroga et al., 2015; Gutiérrez-Rodríguez et al., 2016) and the GFAP-CB₁-RS exclusively in astrocytic processes. Furthermore, these results may suggest that the regulation of the CB₁ receptor expression in glutamatergic and GABAergic cells may be independent one from another. Moreover, astrocytic CB₁ receptor expression may be also independent from the CB₁ receptor in neurons. However, the absence of a crosstalk in the expression of CB₁ receptor between these type of cells cannot be proved with the present experimental design, since the analysis of only one point in the mouse's life time is insufficient to assess the regulation pathway. Therefore, further analysis would be required in order to corroborate this assumption.

The demonstration in this thesis work that the hippocampus of the rescue Glu-CB₁-RS, GABA-CB₁-RS and GFAP-CB₁-RS mutant mice maintain the normal anatomical distribution and expression levels of CB₁ receptors in the cell types with restored receptors, proves the great potential of these mutants for the study of the CB₁ receptor function in specific brain cell populations. In fact, the rescue strategies have the advantage of the reestablishment and visualization of existing CB₁ receptors levels in glutamatergic neurons, GABAergic neurons and astrocytes more accurately, without the interference of additional cells expressing CB₁ receptors. This is especially important for the anatomical and functional investigation of the endocannabinoid system in cell types or brain areas with sparse CB₁ receptors. Finally, the comprehension of the exact expression and distribution of the CB₁ receptor will not only help to better understand the receptor architecture of the brain, but will also improve the conceptual framework for a more specific pharmacological intervention against complex brain diseases in regions where balanced CB₁ receptors are crucial for brain function.

6.3 CB₁ RECEPTOR EXPRESSION IN GFAPhrGFP-CB₁-WT MUTANT MICE

Additionally, mutant mice target to express hrGFP into astroglial cells were used in this thesis. In this animal model, the percentage and density (particles/μm) of astrocytic elements expressing CB₁ receptor were higher and more statistically significant when compared with the GFAP-CB₁-RS mutants using GFAP as a marker for astrocytes. These results suggest that the expression of CB₁ receptors on astrocytes could actually be higher than in previous observations reported using GFAP as a marker for the identification of the astrocytes (Han et al., 2012; Bosier et al., 2013). A plausible explanation could be based on the different molecular nature of the GFAP and hr-GFP proteins. GFAP is a cytoskeletal protein which is assembled in intermediate filament packet (Inagaki et al., 1994; Eng et al., 2000; Hol and Pekny, 2015), then GFAP immunostaining only shows the main radial processes of the astrocyte. However, hr-GFP is a diffusible protein which fills all the cytoplasmic regions, including the fine processes of astrocytes that are

normally lacking GFAP (Nolte et al., 2001), hence, a better detection of astrocytic processes can be accomplished.

6.3.1 TRIPARTITE SYNAPSE: ANATOMICAL INTERPLAY BETWEEN CB₁ RECEPTORS IN ASTROCYTES AND THE NEARBY SYNAPSES

Taking advantage of the better detection of CB₁ receptors in astrocytes of the GFAP^{hr}GFP-CB₁-WT mouse, we also examined the distance between the astrocytic CB₁ receptor and the nearest synapse in order to put the CB₁ receptors in astrocytes in the anatomical context of the functional tripartite synapse (Navarrete and Araque, 2008, 2010; Navarrete et al., 2013, 2014; Gómez-Gonzalo et al., 2014). It was demonstrated that the endocannabinoids coming from a postsynaptic neuron activate astrocytic CB₁ receptors leading to an increase in the intracellular calcium that triggers a release of astrocytic glutamate, which stimulates a neurotransmitter release in heteroneuronal synapses (Navarrete and Araque, 2008, 2010; Navarrete et al., 2013, 2014; Gómez-Gonzalo et al., 2014).

Glial cells constitute the most abundant cellular population in the central nervous system. Amongst glial cell types, astrocytes are excellent players in brain information processing (Volterra and Meldolesi, 2005), due to the bidirectional communication established with neurons (Araque et al., 2001; Bezzi and Volterra, 2011) through intricate morphological and biochemical interactions. The morphology of astrocytes is particularly complex, suggesting that their structures have important functional roles in brain functions. In rodents, astrocytes are distributed covering non-overlapping domains in the cerebral cortex, where they contact tens of thousands of synapses (Halassa et al., 2007). Astrocytes are in close apposition to the synaptic structures, forming tripartite synapses, and play important roles in maintaining and regulating synaptic physiology (Perez-Alvarez et al., 2014). Functionally, endocannabinoids, through CB₁, promote astroglial differentiation and mediate neuron-astrocyte communication regulating synaptic transmission (Aguado et al., 2006; Navarrete and Araque, 2008, 2010). Furthermore, our laboratories (Han et al., 2012) showed that the impairment of spatial working memory and *in vivo* long-term depression at hippocampal CA3-CA1 synapses, induced by an acute exposure of exogenous cannabinoids, is fully abolished in conditional mutant mice lacking CB₁ receptors in brain astroglial cells but is conserved in mice lacking CB₁ receptors in glutamatergic or GABAergic neurons. Blockade of neuronal glutamate N-methyl-D-aspartate receptors (NMDAR) and of synaptic trafficking of glutamate α -amino-3-hydroxy-5-methylisoxazole propionic acid receptors (AMPA) also abolished cannabinoid effects on spatial working memory and long-term depression induction and expression (Han et al., 2012).

In addition, CB₁ activation in astrocytes is involved in energy supply to the brain. Our findings have revealed that genetic and pharmacological manipulations of the CB₁ receptor expression and activity in cultured cortical and hypothalamic astrocytes demonstrated that cannabinoid signaling controls the levels of leptin receptors expression (Bosier et al., 2013). Lack of CB₁ receptors also markedly impaired leptin-mediated activation of signal transducers and activators of transcription 3 and 5 (STAT3 and STAT5) in astrocytes. In particular, CB₁ deletion

determined a basal over activation of STAT5, thereby leading to the downregulation of leptin receptors expression, and leptin failed to regulate STAT5-dependent glycogen storage in the absence of CB₁ receptors in astrocytes (Bosier et al., 2013).

Our present results showed that the most common distance between astrocytic CB₁ receptors and the nearest synapses is in a rank of 400-800 nm. Moreover, most of those synapses were found to be excitatory. Thus, it is tempting to suggest that this distance would represent the distance travelled by the endocannabinoid molecules from the postsynaptic neuron to the CB₁ receptor localized in the astrocyte. According to this interpretation, once inside the astrocyte, the endocannabinoid would activate the astrocytic CB₁ receptor, hence triggering the increase of calcium and stimulate the release of neurotransmitter that modulates synaptic transmission and plasticity. Finally, the fixed anatomical distribution pattern of the astrocytic CB₁ receptors relative to the nearest synapse both in the CA1 stratum radiatum and in the dentate molecular layer suggests the existence of a molecular architecture underlying the functional activity of the astrocytic CB₁ receptors at the tripartite synapse.

7 CONCLUSIONS

The conclusions of the Thesis work are the following:

1. The CB₁ receptor in glutamatergic neurons of the Glu-CB₁-RS mouse hippocampus maintains the same levels of expression and localization of the CB₁-WT mouse hippocampus.
2. The CB₁ receptor in GABAergic neurons of the GABA-CB₁-RS mouse hippocampus maintains the same levels of expression and localization of the CB₁-WT mouse hippocampus.
3. The CB₁ receptor in astrocytes of the GFAP-CB₁-RS mouse hippocampus maintains the same levels of expression and localization of the CB₁-WT mouse hippocampus.
4. The detection of CB₁ receptors in astrocytes of the GFAPhrGFP-CB₁-WT mouse hippocampus is significantly higher than in CB₁-WT mouse hippocampus with GFAP as an astrocytic marker.
5. The most common distance between the astrocytic CB₁ receptors and the nearest synapses in the hippocampus, is between 400-800 nm.
6. The majority of the CA1 and dentate molecular layer synapses surrounded by CB₁ receptor immunopositive astrocytes in the 400-800 nm range are of excitatory nature.
7. The CB₁ receptor rescue mutant mice characterized in this Doctoral Thesis have proven: 1) to express CB₁ receptors in specific brain cell types; 2) the re-expression is limited to the particular neuronal populations or to astrocytes; 3) the endogenous levels of CB₁ receptors are maintained in the brain cell types re-expressing the receptor.
8. Altogether, the studied CB₁ receptor rescue mutant mice are excellent tools for functional and translational investigations on the role of the CB₁ receptors in the normal and diseased brain.

8 ABBREVIATIONS

- 2-AG: 2-arachidonoylglycerol.
- AA: Arachidonic acid.
- ABHD12: α/β -hydrolase domain containing 12.
- ABHD6: α/β -hydrolase domain containing 6.
- AC: Adenylate cyclase.
- AEA: N-arachidonylethanolamine or anandamide.
- AMPAR: α -amino-3-hydroxy-5-methyl-isoxazole propionic acid receptor.
- Amyg: Amygdala.
- AON: Anterior olfactory nucleus.
- as: Astrocyte.
- Astro: Astrocytic.
- CA: Cornu ammonis or Ammon's horn.
- CA1: Region 1 of Cornu Ammonis.
- CA2: Region 2 of Cornu Ammonis.
- CA3: Region 3 of Cornu Ammonis.
- Cb: Cerebellar cortex.
- CB₁: Cannabinoid type I receptor
- CB₁-KO: Cannabinoid type-1 receptor knock-out mouse.
- CB₁-WT: Cannabinoid type-1 receptor wild type mouse.
- CB₂: Cannabinoid type 2 receptor.
- CNS: Central Nervous System.
- CPu: Caudate putamen.
- Cx: Cortex.
- DAB: Diaminobezidine.
- den: Dendrite.
- DG: Dentate gyrus.
- DG: Diacylglycerol.
- DGL: Diacylglycerol lipase.
- DSE: Depolarization-induced suppression of excitation.
- DSI: Depolarization-induced suppression of inhibition.
- eCB-LTD: Endocannabinoid-mediated long-term depression.
- eCBs: Endocannabinoids.
- eCB-STD: Endocannabinoid-mediated short-term depression.
- EP: Entopeduncular nucleus.
- EXCIT: Excitatory.
- FAAH: Fatty acid amide hydrolase.
- GABA: Gamma-Aminobutyric acid.
- GABA-CB₁-KO: GABAergic neurons cannabinoid type-1 receptor knock-out mouse.
- GABA-CB₁-RS: GABAergic neurons cannabinoid type-1 receptor rescue mouse
- GFAP: Glial Fibrillary Acidic Protein.
- GFAP-CB₁-KO: Astrocyte cannabinoid type-1 receptor knock-out mouse
- GFAP-CB₁-RS: Astrocyte cannabinoid type-1 receptor rescue mouse
- GFAPhrGFP-CB₁-KO: CB₁-KO mouse that express hrGFP in astrocytes.
- GFAPhrGFP-CB₁-WT: CB₁-WT mouse that express hrGFP in astrocytes.
- GFP: Green Fluorescence Protein.
- Glu-CB₁-KO: Dorsal telencephalic glutamatergic neurons cannabinoid type-1 receptor knock-out mouse.
- Glu-CB₁-RS: Dorsal telencephalic neurons cannabinoid type-1 receptor rescue mouse.

- GP: Globus pallidus.
- GPCRs: G protein coupled receptors.
- GrDG: Granular dentate gyrus.
- Hi: Hippocampus.
- INHIB: Inhibitory.
- LEA: Lateral entorhinal cortex.
- LMol: Lacunosum moleculare,
- LTD: Long-term depression.
- LTP: Long-term potentiation.
- M1: Primary motor cortex.
- MAGL: Monoacylglycerol lipase.
- MEA: Medial Entorhinal Cortex.
- mGluR: Metabotropic glutamate receptors.
- Mid: Midbrain.
- MO: Medulla oblongata.
- MoDG: Molecular dentate gyrus.
- NAc: Nucleus accumbens.
- NAPE: N-arachidonoyl phosphatidylethanolamine.
- NAPE-PLD: N-acyl phosphatidylethanolamine specific phospholipase D.
- NAT: N-acyltransferase.
- NMDAR: N-Methyl-D-aspartate receptor.
- NOS: Nitric oxide synthase.
- OEA: Oleoylethanolamine.
- Or: Stratum oriens.
- OT: Olfactory tubercle.
- PART: Particles.
- PaS: Parasubiculum.
- PEA: Palmitoylethanolamine.
- PFCx: Prefrontal cortex.
- Pir: Piriform cortex.
- PLC: Phospholipase C.
- Po: Pons.
- PoDG: Polymorph dentate gyrus.
- PPAR: Peroxisome Proliferator-Activated Receptor.
- PROC: Processes.
- PrS: Presubiculum.
- Py: Pyramidal cell hippocampus.
- Rad: Stratum radiatum.
- RECEP: Receptor.
- S1: Primary somatosensory cortex.
- SLu: Stratum lucidum.
- SNR: Substantia nigra pars reticulata.
- sp: Dendritic spine.
- STAT: Signal transducer and activator of transcription.
- STOP-CB₁: mouse carrying a loxP-flanked stop cassette inserted in the 5'UTR upstream of the CB₁ receptor translational start codon.
- Str: Striatum.
- Sub: The subiculum.
- TER: Terminals.
- Th: Thalamus.

- TRPA1: Transient Receptor Potential Ankyrin 1.
- TRPV1: Transient Receptor Potential Vanilloid 1.
- V1: Primary visual cortex.
- VIP: Vasoactive intestinal polypeptide.
- VP: Ventral pallidum.
- WT: Wild type.
- Δ^9 -THC or THC: Delta (9) - tetrahydrocannabinol.

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