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Review



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#### Author for correspondence:

Katelyn Cuttler e-mail: kcuttler@sun.ac.za

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# Emerging evidence implicating a role for neurexins in neurodegenerative and neuropsychiatric disorders

Katelyn Cuttler<sup>1</sup>, Maryam Hassan<sup>3</sup>, Jonathan Carr<sup>2,4</sup>, Ruben Cloete<sup>3</sup> and Sorava Bardien<sup>1,4</sup>

<sup>1</sup>Division of Molecular Biology and Human Genetics, Department of Biomedical Sciences, and <sup>2</sup>Division of Neurology, Department of Medicine, Faculty of Medicine and Health Sciences, Stellenbosch University, Cape Town, South Africa

<sup>3</sup>South African Medical Research Council Bioinformatics Unit, South African National Bioinformatics Institute, University of the Western Cape, Cape Town, South Africa

<sup>4</sup>South African Medical Research Council/Stellenbosch University Genomics of Brain Disorders Research Unit, Cape Town, South Africa

(D) KC, 0000-0003-1833-5426

Synaptopathies are brain disorders characterized by dysfunctional synapses, which are specialized junctions between neurons that are essential for the transmission of information. Synaptic dysfunction can occur due to mutations that alter the structure and function of synaptic components or abnormal expression levels of a synaptic protein. One class of synaptic proteins that are essential to their biology are cell adhesion proteins that connect the pre- and post-synaptic compartments. Neurexins are one type of synaptic cell adhesion molecule that have, recently, gained more pathological interest. Variants in both neurexins and their common binding partners, neuroligins, have been associated with several neuropsychiatric disorders. In this review, we summarize some of the key physiological functions of the neurexin protein family and the protein networks they are involved in. Furthermore, examination of published literature has implicated neurexins in both neuropsychiatric and neurodegenerative disorders. There is a clear link between neurexins and neuropsychiatric disorders, such as autism spectrum disorder and schizophrenia. However, multiple expression studies have also shown changes in neurexin expression in several neurodegenerative disorders, including Alzheimer's disease and Parkinson's disease. Therefore, this review highlights the potential importance of neurexins in brain disorders and the importance of doing more targeted studies on these genes and proteins.

# 1. Introduction

There is accumulating evidence to suggest that synaptic dysfunction is present in both neuropsychiatric disorders, such as autism spectrum disorders (ASDs), schizophrenia and bipolar disorder (BD), and neurodegenerative disorders, such as Parkinson's disease (PD), Alzheimer's disease (AD) and Huntington's disease (HD) [1]. In fact, involvement of the synapse is such a prominent feature of the pathogenesis of various brain disorders that it has led to the coining of a specific term, 'synaptopathies'. Indeed, in the case of PD, the involvement of synaptopathy as an initial and central event in the disease pathogenesis, which precedes neuronal damage, has been postulated [2]. Synaptic dysfunction can occur due to mutations that alter the structure and function of synaptic components or abnormal expression levels of a synaptic protein.

Synapses are specialized junctions between neurons that transmit information and they connect neurons into millions of 'neural circuits' that underlie all brain functions [3]. The information transmitted allows the nervous system to respond

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to external stimuli and controls bodily functions, behaviour, emotions and memories [4]. This system is tightly controlled and regulated, and even slight perturbations can lead to synaptic dysfunction. An important aspect of synapse biology is the cell adhesion molecules that connect pre- and post-synaptic compartments [5]. These interactions in the synaptic cleft help to maintain synapse structure by delineating mutual boundaries [6]. These proteins are also important in synapse plasticity as synaptic cell adhesion is able to regulate the remodelling of synapses [7]. Interestingly, they are also involved in trans-synaptic signalling [5]. Thus, these proteins are highly important in the organization of synaptic junctions and overall brain function.

Neurexins are one type of synaptic cell adhesion molecule. They are pre-synaptically localized and bind to neuroligins and other proteins in the post-synapse (figure 1). Neurexins and their common binding partners, neuroligins, have recently gained more pathological interest as variants in both have been associated with several neuropsychiatric disorders, including autism and schizophrenia [8]. This further suggests that synaptic dysfunction plays a role in the development of these disorders. Synaptic dysfunction is also known to occur in neurodegenerative disorders [9]; however, it was considered an endpoint of these disorders, due to the considerably later onset of clinical symptoms and progressive appearance of cognitive deficits. This dichotomy has, recently, been challenged by the creation of 'disease-in-a-dish' models for multiple central nervous system (CNS) pathologies [9]. This research has identified commonalities between developmental and degenerative disorders, at both the cellular and molecular level, with most of these common mechanisms meeting at the synapse level [9]. Indeed, our laboratory has, recently, found a novel variant (p.G849D) in the NRXN2 gene which may be implicated in PD [10]. Therefore, we believe it is important to investigate the potential role of neurexins in various neuropsychiatric and neurodegenerative disorders.

In this review, we summarize some of the key physiological functions of the neurexin protein family and the protein networks they are involved in. We also examine the available published literature to determine what research has been done on neurexins in neuropsychiatric and neurodegenerative disorders. This analysis provides an overview on what progress has been made in understanding the roles of synaptic functioning in these disorders and reveals the gaps in knowledge in this field.

# 2. Structure and biological functions of neurexins

Neurexins were first identified using affinity chromatography when neurexin 1 $\alpha$  was found in rat brain extract on a column of  $\alpha$ -latrotoxin [11].  $\alpha$ -latrotoxin is a potent neurotoxin from black widow spider venom that stimulates synaptic vesicle exocytosis and induces massive neurotransmitter release [12]. This work has been continued by Südhof and coworkers [13] who have characterized the neurexin proteins and their binding partners, the neuroligins [14].

In mammals, the neurexins are encoded by three *NRXN* genes (*NRXN1-3*), each of which has both an upstream promoter that is used to generate the  $\alpha$ -neurexins, and a downstream promoter that is used to generate the shorter  $\beta$ -neurexins [13,15]. Neurexins also undergo extensive alternate splicing



**Figure 1.** Location of neurexins and their binding partners, neuroligins, in the synapse. Several neurexin–neuroligin pathway proteins are shown as well as synaptic vesicle-binding proteins. NMDAR, *N*-methyl-D-aspartate receptor; mGluR5, metabolic glutamate receptor 5; PSD-95, post-synaptic density protein 95; Shank, SH3 and multiple ankyrin repeat domains protein.



**Figure 2.** Structural domain organization of the  $\alpha$  and  $\beta$  forms of neurexin. (*a*)  $\alpha$ -neurexin. (*b*)  $\beta$ -neurexin. EGF, epidermal growth factor-like region; LNS, laminin/neurexin/sex hormone-binding domain; PDZ, PSD-95, DLG1, ZO-1 domain; SP, signal peptide; TM, transmembrane domain.

at five splice sites, generating significant diversity of more than 2000 potential variants [13,16]. The fact that neurexin splice insert sequences and their positions are well conserved among neurexin genes and between species supports the idea that alternative splicing has important functional roles.

The neurexins are transmembrane proteins that consist of an extracellular region responsible for trans-synaptic interactions, a transmembrane domain and a smaller cytoplasmic domain named PSD-95, DLG1, ZO-1 binding domain (PDZ) that is involved in intracellular protein interactions and signalling (figure 2) [13].  $\alpha$ -neurexins are composed of six large extracellular laminin/neurexin/sex



**Figure 3.** A summary of GO terms associated with neurexins 1–3. (*a*) Biological processes. (*b*) Cellular components. (*c*) Molecular functions. All: total number of proteins analysed. The number above each bar indicates the number of proteins assigned to that category. Figure generated by WebGestalt (http://www.webgestalt. org) [18].

hormone-binding (LNS) globulin domains with three interspersed epidermal growth factor (EGF)-like regions (figure 2).  $\beta$ -neurexins are shorter and only have the sixth extracellular LNS domain and no EGF-like regions (figure 2). Only neurexin 1 protein structures (both  $\alpha$  and  $\beta$  forms) have been solved experimentally in *Mus musculus, Rattus norvegicus* and *Bos taurus*. However, these structures have not yet been solved in humans.

Neurexins are localized pre-synaptically and are distributed to both excitatory and inhibitory synapses [8]. Their functions are mediated by their binding to neuroligins (figure 1). Neuroligins have five known isoforms and are expressed post-synaptically [17]. Consequently, neurexins and neuroligins form synaptic complexes in the synaptic cleft and have been found to control synapse formation, maturation, validation and function [17]. Various combinations of the different neurexins and neuroligin binding partners at synapses may also help determine synapse specificity through differential interactions between multiple splice variants and isoforms of these proteins [8].

Primarily, neurexins function to maintain synaptic organization. Gene ontology (GO) analysis by WebGestalt (http:// www.webgestalt.org) [18] of the three neurexins indicates that they all function in protein binding, ion binding and possess molecular transducer activity (figure 3). They are also involved in cellular component organization, developmental processes, response to stimuli, cell communication and biological regulation. These processes thus demonstrate how neurexins are able to maintain synaptic organization but also show their multi-functional nature. As such, it is conceivable that disruptions in neurexins could be detrimental to their various functions and affect overall neuronal function and integrity.

# 3. Biological pathways and interacting partners of neurexins

To understand the broader biological pathways that the neurexins are involved in, protein-protein interaction network analysis and co-expression analysis was performed using the tools, STRING (https://string-db.org) [19] and GeneMania (https://genemania.org) [20]. STRING finds related genes by accessing the STRING database which contains experimental data and computational predictions. Data in STRING are weighted and integrated and a confidence score of 0–1 is calculated for all interacting protein partners. GeneMANIA finds proteins related to neurexins by leveraging functional association data, such as interactions, pathways, co-expression, colocalization and protein domain similarity. All functional data for the proteins observed in these networks were obtained from UniProt (https://www.uniprot.org) [21], unless otherwise stated, while pathway data were obtained from KEGG (https://www.kegg.jp) [22].

#### 3.1. String analysis

Weighted string analysis was conducted on neurexin 1, 2 and 3 individually to determine their binding partners (figure 4*a*–*c*). Based on this analysis, there is strong evidence that neurexin 1 interacts with 10 proteins including calcium/ calmodulin-dependent serine protein kinase (CASK), leucinerich repeat transmembrane neuronal protein 1 (LRRTM1), LRRTM2, LRRTM3, neuroligin 1, neuroligin 2, neuroligin 3, neuroligin 4X, SH3 and multiple ankyrin repeat domains protein 2 (SHANK2) and synaptotagmin-1 with scores of 0.987, 0.983, 0.985, 0.975, 0.998, 0.997, 0.998, 0.997, 0.975, 0.974, respectively. Similarly, neurexins 2 and 3 also have 10 interactors each. There is strong evidence that neurexin 2 interacts with CASK, discs large homologue 4 (DLG4), LRRTM1, LRRTM2, LRRTM3, neuroligin 1, neuroligin 2, neuroligin 3, neuroligin 4X and SHANK2 with scores of 0.979, 0.977, 0.984, 0.983, 0.972, 0.998, 0.998, 0.998, 0.997 and 0.969, respectively. There is strong evidence that neurexin 3 interacts with CASK, DLG4, LRRTM1, LRRTM2, LRRTM3, neuroligin 1, neuroligin 2, neuroligin 3, neuroligin 4X and SHANK2 with scores of 0.978, 0.971, 0.979, 0.983, 0.971, 0.998, 0.997, 0.997, 0.997 and 0.969, respectively. The STRING analyses performed on the three neurexins identified



**Figure 4.** Protein interacting partners of neurexins 1-3. (a-c) STRING network of neurexins 1-3. Nodes represent gene-encoded proteins. Edges represent proteinprotein associations. Connections between nodes represent the relationship between proteins. A bold line implies a higher confidence level. (d) GeneMANIA network of all three neurexins. Nodes represent gene-encoded proteins. Larger nodes indicate higher protein scores. Edges represent protein-protein associations. Connections between nodes represent the relationship between proteins. A bold line implies a higher confidence level.

interacting proteins with very high confidence scores since the lowest score across the analyses was 0.969. This means that there is strong experimental evidence that these proteins interact with one or more of the neurexins.

Analysis of the identified neurexin binding partners revealed many proteins important in the maintenance and functioning of synapses. Notably, variants in several of these proteins are implicated in neuropsychiatric and developmental disorders. Variants in neuroligin 1 and SHANK2 have been implicated in susceptibility to autism [23,24], while variants in neuroligin 3 have been implicated in Asperger syndrome and autism [25]. Variants in neuroligin 4X have been implicated in X-linked forms of Asperger syndrome, autism susceptibility and mental retardation [25–27]. Variants in CASK have been implicated in FG syndrome 4, an X-linked genetic disorder and mental retardation [28–32], variants in DLG4 have been implicated in intellectual developmental disorder 62 [33,34] and variants in synaptotagmin-1 have been implicated in Baker–Gordon syndrome [35].

Furthermore, binding partners of these proteins as well as the pathways they occur in could also give insight into the development of disease. CASK binds to amyloid precursor protein and neuroligin 1 binds to amyloid- $\beta$ , both of which are important in AD. LRRTM3 is also a known positive regulator of amyloid- $\beta$  formation. Notably, *LRRTM3* may be considered a candidate gene for late-onset AD as it promotes the processing of amyloid precursor protein which leads to toxic amyloid- $\beta$  accumulation [36]. DLG4 is involved in dopamine receptor binding and synaptotagmin-1 regulates dopamine secretion. The loss of dopamine functioning is crucial in PD. Indeed, DLG4 is involved in several pathways of neurodegeneration (in multiple diseases), the HD pathway as well as cocaine addiction.

### 3.2. GeneMANIA

GeneMANIA analysis was performed on the neurexins to reveal further potential protein–protein interactions (figure 4*d*; electronic supplementary material, table S1). We performed the analysis by selecting only proteins with stronger evidence of neurexin interactions, such as interactions with physical evidence, and evidence from co-expression and co-localization studies.

All of the binding partners observed by STRING analysis were still present; however, more interacting proteins were also identified. These proteins have more diverse functions but still function in overall synapse maintenance.

This analysis further identified afadin (AFDN), Rho GTPase activating protein 10 (ARHGAP10), cerebellin 1 (CBLN1), dystroglycan (DAG1), microtubule actin cross-linking factor 1 (MACF1), neurexophilin-2, neurexophilin-3, PDZ domain-containing protein 2 (PDZD2), proteolipid protein 1 (PLP1), syndecan binding protein 1 (SDCBP), SDCBP2, SH3 domain-containing GRB2-like protein 2 (SH3GL2), signalinduced proliferation-associated 1-like protein 1 (SIPA1L1), synaptotagmin-13 (SYT13) TAFA chemokine-like family member 1 (TAFA1), TUBB-like protein 1 (TULP1) and XKrelated protein 4 (XKR4) as interactors of one or more neurexin proteins. AFDN, ARHGAP10, MACF1 and SIPA1L1 are all involved in actin filament binding/ organization, while PDZD2, SDCBP and SDCBP2 are involved in cell binding and cytoskeletal organization. Dysregulation of any of these proteins could thus affect cell adhesion and binding at the synapse. In addition, variants in MACF1 have been implicated in lissencephaly 9 with complex brainstem malformation [37]. CBLN1 is essential for synapse integrity and plasticity and its disruption could lead to synapse dysfunction. DAG1 has multiple functions, such as laminin and basement membrane assembly, cell survival, peripheral nerve myelination, nodal structure and cell migration. Variants in DAG1 have been implicated in both type A and type C muscular dystrophy-dystroglycanopathy [38-41]. Muscular dystrophies are genetic disorders characterized by the degeneration of skeletal muscle. Type C muscular dystrophy-dystroglycanopathy affects the limbgirdle area [40], while type C is congenital with brain and eye anomalies [39]. Neurexophilin-2 and neurexophilin-3 are both ligands for  $\alpha$ -neurexins and are involved in the neuropeptide signalling pathway. Disruption of these proteins could, therefore, affect neurotransmitter release and the subsequent signalling. PLP1 is the major myelin protein in the CNS and is important for maintaining the structure of myelin. Disruption of this protein could, therefore, negatively affect the downstream myelination of neurons, as is seen in multiple sclerosis (MS). Interestingly, PLP1 is also involved in the development of the substantia nigra, the main brain region affected by PD. Therefore, PLP1 alterations could also lead to disruptions in this brain region. SH3GL2 has been implicated in synaptic vesicle endocytosis, while synaptotagmin-13 may be involved in transport vesicle docking to the plasma membrane. Dysregulation of these proteins could thus affect neurotransmitter functioning. TAFA1 is involved in the modulation of neural stem cell proliferation and differentiation; therefore, dysregulation of this protein could result in developmental disorders. TULP1 is required for normal development of photoreceptor synapses. Variants in TULP1 are associated with Leber congenital amaurosis [42,43] and retinitis pigmentosa [42,44-47]. However, this protein is also involved in actin filament binding, therefore, its dysregulation could also affect cell adhesion and binding at the synapse. Not much is known about XKR4 except that it is involved in apoptosis during development. Therefore, its dysregulation could also possibly result in developmental disorders.

GO terms and physiological/pathway information on all binding partners identified by STRING and GeneMANIA are available in electronic supplementary material, figure S1 and table S1.

# 4. Role of neurexins in neuropsychiatric disorders

Literature-based searches using neurexin as a search term identified several studies that reported an association of neurexins in various neuropsychiatric disorders. The main findings of these studies are reported in table 1 and are summarized below.

### 4.1. Human studies

*NRXN1* has been well documented for its association with ASDs [54]. Several genetic analyses of families and populations of people with ASD have shown that copy number variations (CNVs) and de novo mutational events at the *NRXN1* locus are enriched in ASD [48,49,51,52,54]. In one

study, *NRXN1* was sequenced in cases of ASD with mental retardation [50]. Mutations (c.–3G > T in the Kozak region, c.3G > T at the initiation codon (p.M1), p.R375Q and p.G378S) were found in the *NRXN1* $\beta$  coding region thereby potentially implicate synapse dysfunction an important determinant in ASD [50].

The first evidence for a potential role of *NRXN2* in ASD was provided by a report of a frameshift mutation within *NRXN2* exon 12 (c.2733delT) in a boy with ASD and his father who had severe language delay [57]. This mutation results in a truncated neurexin  $2\alpha$  protein that lacks the binding sites for the established post-synaptic binding partners LRRTM2 and neuroligin 2 [57]. Subsequently, a 21-year-old man with a clinical phenotype including autistic traits, such as speech and language deficits and pathological insistence on routine, was reported to have a 570 kb de novo deletion of 24 genes at chromosome 11q13.1, including *NRXN2* [58].

Using microarray analyses on RNA extracted from brain tissue, Mirnics et al. [69] did not observe a difference in neurexin 1 expression between schizophrenia (SCZ) and control samples. However, since then, a link between neurexin 1 and SCZ has been reported in other studies. One study reported that NRXN1 deletions are more common in those with SCZ; however, it also found that there was incomplete penetrance of these deletions in families with SCZ [72]. Kirov et al. [70] observed a deletion in an SCZ patient at 2p16.3 that disrupts NRXN1 and predicted that it was highly likely to be pathogenic. Also, NRXN1 deletions were shown to segregate with several neuropsychiatric disorders in a study of a complex family [78]. The proband had SCZ and other members of his family had mental retardation, schizophreniform disorder and affective disorder [78]. After genotyping the proband and eight family members, they found two rare deletions upstream of the NRXN1 gene (2p16.3) that co-segregate with these disorders [78]. Notably, this shows that deletions in NRXN1 may manifest as multiple neuropsychiatric phenotypes.

Angione *et al.* [64] implicated *NRXN1* in epilepsy. They identified a 2p16.3 deletion in an 8-year-old male patient diagnosed with epilepsy showing symptoms of myoclonicatonic seizures (EMAS) [64]. This deletion included the first five exons of the *NRXN1* gene [64].

*NRXN* genes may also be involved in treatment response. In one study, it was found that variants in *NRXN1* may affect the long-term treatment outcome of patients with BD by modulating the effects of antipsychotics [61]. In a study of Levetiracetam resistance, an antiepileptic drug, Grimminger *et al.* [63] found that neurexin 1 is differentially expressed in non-responder and responder patients with mesial temporal lobe epilepsy (mTLE), whereby lower levels of neurexin 1 were observed in responder patients.

#### 4.1.1. Association studies

A genome-wide association study (GWAS) by Liu *et al.* [51] specifically examined *NRXN1* in an autism cohort of the Chinese Han population and discovered 22 variants that were associated with ASD. In this cohort, one SNP (rs2303298) was also significantly associated with a risk of developing ASD [51]. Furthermore, a GWAS of SCZ in Spain showed that a *NRXN1* single nucleotide polymorphism (SNP) (rs3850333) was close to the significance threshold [71], while another GWAS in American patients of European or African ancestry showed that *NRXN1* is associated with antipsychotic

Disorders Branch; CIBERSAM, Centro de Investigación Biomédica en Red de Salud Mental; CNV, copy number variation; EMAS, epilepsy with myoclonic-atonic seizures; GWAS, genome-wide association study; hESC, human embryonic stem Table 1. List of studies that have implicated neurexin genes in neuropsychiatric disorders. AAV, adeno-associated virus; AGRE, Autism Genetic Resource Exchange; Array-CGH, array comparative genomic hybridization; CBDB, Clinical Brain cell; iN, induced neuron; iPSC, induced pluripotent stem cell; KO, knockout; LC-MS/MS, liquid chromatography mass spectrometry/mass spectrometry; mESC, mouse embryonic stem cell; mTLE, mesial temporal lobe epilepsy; NGS, nextgeneration sequencing; NIMH, National Institute of Mental Health; RT–PCR, reverse transcriptase–polymerase chain reaction; SNP, single nucleotide polymorphism; SSC, Simons Simplex Collection; STEP-BD, Systematic Treatment Enhancement Program for Bipolar Disorder; WT, wild-type.

disorder/disease	neurexin				
process	gene	type of study	methods	main finding	reference
autism spectrum disorder (ASD)	NRXN1	genetic analysis	used SSC samples and the SSC database to extract 'trios' consisting of a mother, father and an ASD-affected child performed genetic analyses to identify CNVs	a de novo CNV in <i>NRXV1</i> was discovered in a large cohort of families with a single ASD-affected child and at least one unaffected sibling	[48]
ASD	NRXN1	GWAS	1174 families from SSC were genotyped identified CNVs and de novo events	rare de novo events/CNVs at <i>NRXN1</i> are strongly associated with autism	[49]
ASD	NRXN1ß	genetic analysis	86 patients with both ASD and mental retardation The coding sequence of the $NRXN1\beta$ gene was analysed by PCR	four novel mutations in <i>NRXN1</i> $\beta$ were identified by sequencing the exon of the gene in cases with autism and mental retardation	[50]
ASD	NRXN1	genetic analysis	313 ASD patients and 500 healthy controls from a Chinese autism cohort were recruited performed genomic DNA sequencing	22 variants in the <i>NRXN1</i> gene were discovered in the Chinese Han population; one SNP (rs2303298) was significantly associated with a risk of autism in this cohort	[51]
ASD	NRXN1	genetic analysis	2478 ASD individuals from SSC and 719 ASD individuals from AGRE 580 controls from ClinSeq and NIMH used a custom microarray to analyse CNVs	recurrent CNVs in <i>MRXN1</i> are enriched in autism.	[52]
ASD	NRXN1, 2 and 3	cell culture	iPSCs were produced from probands and unaffected family members iPSCs underwent neuronal differentiation to organoids RNA sequencing was performed on both iPSCs and differentiated organoids	neurexin 1, 2 and 3 mRNA is overexpressed in patient-derived iPSCs and differentiated organoids	[23]
ASD	NRXN1	genetic analysis	2591 families from SSC were genotyped identified CNVs, de novo deletions and ASD risk genes	<i>NRXV1</i> is an ASD risk gene	[54]
ASD	NRXN1 and 2	animal study	RNA was isolated from the whole brain of age-matched monoamine oxidase A KO mice and wild-type mice Microarrays were used to determine gene expression levels	neurexin 1 and 2 are downregulated in monoamine oxidase A KO mice	[55]
ASD	α-NRXNs	animal and cell culture study	transfected C. <i>elegans</i> strains and HEK-293 cells with plasmids expressing NRXN1 $\alpha$ and different $\alpha$ 2 $\delta$ subunits performed co-immunoprecipitation and pull-down assays	changes in $\alpha$ -neurexin binding to $\alpha$ 2 $\delta$ -3 subunits of N-type calcium channels could be important in some forms of autism spectrum disorders	[56]
ASD	NRXN2	genetic analysis	142 ASD patients and 94 non-syndromic controls sequenced NRXN 1,2 and 3 genes	observed a frameshift mutation in <i>NRXN2</i> exon 12 in a patient with ASD inherited from a father with severe language delay	[57]
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Table	

disorder/disease	neurexin				
process	gene	type of study	methods	main finding	reference
ASD	NRXN2	genetic analysis	recruited a patient with speech problems, autistic traits and pancreatic gastrinoma performed array-CGH	a de novo 0.57 Mb microdeletion was observed in chromosome 11q13.1, including NRXN2	[58]
ASD	NRXN2	animal study	used previously collected human faecal samples from typically developing children and children with ASD C57BL/6 J weanlings were colonized with human faecal samples brain tissue RNA was extracted and sequenced	mice colonized by microbiota from ASD patients showed differential splicing of NRXN2	[59]
bipolar disorder (BD)	NRXN3	GWAS	obtained participants from a family study of mood disorders in Taiwan (2008–2012) performed a multi-stage GVMS	NRXN3 shows a significant association with bipolar disorder	[60]
BD	NRXN1	genetic analysis	obtained patient genotyping and clinical data from STEP-BD analysed data to determine the effect of individual markers on phenotypes	NRXN1 may affect the long-term treatment outcome of bipolar disorder	[61]
borderline personality disorder (BPD)	NRXN3	association study	1439 heroin-dependent BPD cases and 507 neighbourhood controls genotyped NRXN3 SNPs and performed an association analysis	several NRXN3 SNPS were nominally associated with BPD phenotype in heroin- dependent cases	[62]
epilepsy	NRXN1	microarray analysis	obtained 53 biopsy specimens from mTLE patients performed microarray analysis	neurexin 1 is differentially expressed in non-responder and responder mTLE patients to the antiepileptic drug Levetiracetam lower levels of neurexin 1 are observed in responder patients	[03]
epilepsy	NRXN1	genetic testing	77 patients were identified at Children's Hospital Colorado various genetic tests were conducted	a 2p16.3 deletion, which includes the first five exons of the <i>NRXN1</i> gene, was identified in an 8-year-old male EMAS patient	[64]
epile psy/seizures	NRXN2α	animal study	treated adult Wistar rats with kainite or pentylenetetrazole to induce seizures isolated total RNA from whole-rat brains and hippocampi performed RT–PCR to determine the levels of different NRXNs	following kainate- and pentylenetetrazole-induced seizures in rats, neurexin $2lpha$ expression increased in the dentate gyrus of the hippocampus	[65]
fragile X syndrome	NRXN3	animal study	used male and female WT and FMR1 K0 mice (4–6 per experiment) analysed brain sections using riboprobes for NRXN1, 2 and 3 and NLGN 1, 2 and 3	there is increased neurexin 3 mRNA in female fragile X mice, but decreased neurexin 3 mRNA in male fragile X mice	[99]
major depressive disorder (MDD)	NRXN1, 2 and 3	animal study	81 healthy Sprague–Dawley rats were subjected to various mild stress factors extracted proteins from hippocampal post-synaptic density fractions analysis by LC-MS/MS	neurexin 1, 2 and 3 were not differentially expressed in a rat chronic mild stress model of depression	[67]
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disorder/disease	neurexin				
process	gene	type of study	methods	main finding	reference
neuropsychiatric disorders	NRXN1	cell culture	cultured cortical neurons from NRXN1 $\alpha$ KO mice introduced conditional NRXN1 mutations into hESCs using AAV recombination, and differentiated them into human iNs analysed neuronal development, synapse formation and	heterozygous NRXN1 mutations are able to selectively impair neurotransmitter release and increase the levels of the synaptic scaffolding protein, CASK in human iNs but not in the cortical neurons of NRXN1 $\alpha$ KO mice	[68]
schizophrenia (SCZ)	NRXN1	microarray analysis	obtained brain tissue from 12 SCZ patients and 10 controls extracted total RNA performed a microarray analysis	the expression of neurexin 1 was not significantly different between the schizophrenic and control subjects	[69]
SCZ	NRXN1	genetic analysis	selected 45 male and 48 female proband-parent trios from a sample of 600 Bulgarian SCZ trios performed Array-CGH	observed a 0.25 Mb deletion on 2p16.3 in both the proband and affected sibling which disrupts <i>NRXN1</i>	[70]
SCZ	NRXN1	GWAS	3063 SCZ patients and 2847 controls from CIBERSAM performed a GWAS at 95 SNPs	the rs3850333 SNP in the <i>NRXN1</i> gene was close to the significant threshold in a GWAS of schizophrenia in Spain	[17]
22	NRXN1	genetic analysis	obtained DNA of 635 S/Z patients and 635 controls from the CBDB Sibling Study genotyped samples and analysed them for CNVs and deletions	NRXN1 deletions are more frequent in schizophrenia patients there is incomplete penetrance of NRXN1 deletions in families with schizophrenia	[72]
SCZ	NRXN1	genetic analysis	data from 572 SCZ patients and 551 controls were used to select genes for sequencing 153 SCZ patients and 153 controls were sequenced for 21 chosen genes using NGS	missense variants at <i>NRXN1</i> may be protective against schizophrenia susceptibility	[73]
225	INXXN	cell culture	isolated primary rat neurons from hippocampi differentiated human neurons derived from human iPSGs overexpressed Caveolin-1 in both cell types western blotting was used to measure the expression of proteins involved in synaptic plasticity as well as DISC1, an SCZ-associated protein	overexpressing Caveolin-1, a potential therapeutic for schizophrenia, in neurons increased expression of proteins involved in synaptic plasticity (PSD95, synaptobrevin, synaptophysin, neurexin 1 and syntaxin 1) as well as DISC1	[74]
					(Continued.)

disorder/diseaseneurexinprocesgenetype of studymethodsmain findingprocesgenetype of studygenerated iPSCs from 5 childhood-onset SCZ patients and 4 controlsmain findingSCZNRN1animal studygenerated iPSCs into glial cellsmeurexin 1 was downregulated in chimetic miceSCZNRN1animal studygenerated iPSCs into glial cellsmeurexin 1 was downregulated in chimetic miceSCZNRN1animal studygenerated iPSCs into glial cellspatients with childhood-onset schizophreniaransplanted glial cellscollosumcollosumcollosumSCZNRN1GWSobtained genetic data and treatment response data of 302 SCZMRN11 is associated with antipsychotic response tSCZNRN1GWSobtained genetic data and treatment response data of 302 SCZMRN11 is associated with antipsychotic response data of 302 SCZSCZNRN1cell culturea placebo from two chincial SCZ trialsMRN11 dietion SCZ patients treated withSCZNRN1cell culturegenerated PSCs from 3 NRN11 dietion SCZ patients treated withmetrosponse data is controlsSCZNRN1cell culturegenerated PSCs from 3 NRN11 dietion SCZ patients treated withmetrosponse data is controlsSCZNRN1cell culturegenerated PSCs from 3 NRN11 dietion SCZ patients and 3 controlsmetrosponse data is controlsSCZNRN1cell culturegenerated PSCs from 3 NRN11 dietion SCZ patients and 3 controlsmetrosponse data of from sociated with scalebored of the synaptic scaleb	main finding         SCZ patients and 4 controls       neurexin 1 was downregulated in chimeric mice propatients with childhood-onset schizophrenia         cction into the corpus       cction into the corpus	main finding neurexin 1 was downregulated in chimeric mice prod patients with childhood-onset schizophrenia	uced from iPSCs derived from	reference
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and differentiated them into human iNs and increase the levels of the synaptic scaffold	SCZ patients and 3 controls heterozygous NRXN1 deletions impair neurotransmit	heterozygous NRXN1 deletions impair neurotransmitt	r release and synaptic function,	[77]
	ils and increase the levels of the synaptic scaffoldin	and increase the levels of the synaptic scaffolding	protein, CASK in human iNs but	
genetated medica medic	not mESCs generated from NRXN1 K0 mice	not mESCs generated from NRXN1 K0 mice		
analysed neuronal development, synapse formation and	formation and			
neurotransmitter release				
SCZ and other NRXN1 genetic analysis recruited a family with multiple neuropsychiatric disorders two rare deletions upstream of the NRXN1 gene	ychiatric disorders two rare deletions upstream of the NRXN1 gene (2)	two rare deletions upstream of the NRXN1 gene (2p'	6.3) segregate with schizophrenia,	[78]
neuropsychiatric schizophreniform disorder, and affective disord	' members have mental schizophreniform disorder, and affective disorder	schizophreniform disorder, and affective disorder i	ו a family	
disorders disorder and affective disorder and affective disorder	and affective disorder			
genotyped the proband and eight family members	wombove			

response to lurasidone in SCZ patients [76]. Additionally, an association study on Spanish SCZ patients showed that missense mutations in *NRXN1* may actually protect against susceptibility to SCZ [73].

In a Taiwanese GWAS study, a significant association between *NRXN3* and BD was found [60]. And finally, an association study on Australian borderline personality disorder (BPD) patients showed that several *NRXN3* SNPS were nominally associated with BPD in heroin-dependent cases [62].

### 4.2. In vitro and in vivo models of disease

Functional *in vitro* and *in vivo* studies have also found evidence for the roles of neurexins in ASD. Monoamine oxidase A knockout (KO) mice, which are an animal model for autism, exhibited downregulated levels of both neurexin 1 and neurexin 2 [55]. Furthermore, mice colonized with the microbiota of ASD patients showed differential splicing of *NRXN2* [59]. Another animal study showed that changes in the binding of  $\alpha$ -neurexins to N-type calcium channels could be important for some forms of ASD patient-derived induced pluripotent stem cells (iPSCs) and differentiated organoids showed that neurexin 1, 2 and 3 mRNA is overexpressed in these samples [53].

One study examined neurexins in Fragile X syndrome, a genetic disorder with features similar to ASD, and characterized by the silencing of the *FMR1* gene [79]. Individuals with Fragile X experience a range of neurodevelopmental problems, such as learning disabilities and cognitive impairment, and males are usually more severely affected. Using *FMR1* KO mice, researchers probed brain sections to determine the levels of neurexin 1, 2 and 3 [66]. Interestingly, they found that neurexin 3 mRNA levels are increased in female mice but decreased in male mice and predicted that this may help explain the sex difference observed in this disorder [66].

In an animal study of SCZ, neurexin 1 was found to be downregulated [75]. This study generated iPSCs from patients with childhood-onset SCZ, differentiated them into glial cells and injected the glial cells into mice to form chimeric mice as a model organism [75]. Interestingly, an *in vitro* study of SCZ showed that overexpressing Calveolin-1, a potential therapeutic for SCZ, actually increased the levels of neurexin 1 as well as other proteins involved in synaptic plasticity [74].

Neurexin  $2\alpha$  has been implicated in epilepsy and, more specifically, in seizures. Researchers observed an increase in neurexin  $2\alpha$  expression in the dentate gyrus of the hippocampus in an induced-seizure mouse model [65]. Finally, in one study, a rat chronic mild stress model of depression was used to determine if neurexin expression was altered in major depressive disorder; however, no change in neurexin 1, 2 or 3 levels was observed [67].

So far, there have been two studies validating the effect of NRXNs *in vitro*, both by Pak *et al.* [68,77]. These studies cultured human stem cells as well as mice cells generated from *NRXN1* KO mice. The first study introduced two conditional NRXN1 mutations previously seen in a range of neuropsychiatric disorders, including ASD and SCZ, into human embryonic stem cells (hESCs) using adeno-associated virus recombination and differentiated them into human-induced neurons (iNs) [68]. These cells were compared to cortical

neurons generated from NRXN1α KO mice [68]. The second study generated iPSCs from three NRXN1 deletion SCZ patients and three controls, and again differentiated them into human iNs [77]. These cells were compared to mouse embryonic stem cells (mESCs) from NRXN1 KO mice [77]. Both studies showed that heterozygous NRXN1 deletions were able to impair neurotransmitter release and synaptic function, and increase the levels of the synaptic scaffolding protein, CASK, in human iNs but not in mice cells [68,77]. Therefore, these studies provide evidence that NRXN1 deletions exhibit a major synaptic transmission phenotype in humans and are thus meaningful at a pathophysiological level.

In summary, these studies demonstrate a link between *NRXNs* and neuropsychiatric disorders such as ASD and SCZ, especially involving full or partial deletions of these genes. *NRXNs* have also been associated with BD and BPD. In addition, protein expression studies have shown changes in neurexin expression in animal models of epilepsy/seizures and Fragile X syndrome.

# 5. Role of neurexins in neurodegenerative disorders

Additionally, literature-based searches provided proof for the involvement of neurexins in various neurodegenerative disorders, and these studies are listed in table 2 and discussed below.

#### 5.1. Human studies

Studies examining cerebrospinal fluid (CSF) from AD patients have observed lowered expression of neurexin 1 [81], as well as neurexin 2 $\alpha$  and neurexin 3 $\alpha$  [85]. In addition, it was found that these changes precede the neurodegeneration markers as they were observed in the preclinical stage 1 of AD [85]. Moreover, A $\beta_{42}$  fibrils in CSF were found to bind to neurexin 1, 2 and 3 as well as proteoglycans and growth factors [83]. Levels of the synaptic proteins neuronal pentraxin 2 (NPTX2), GluA4containing glutamate (AMPA4), neuroligin 1 and neurexin 2 $\alpha$ are also declined in plasma neuron-derived exomes and this decline was associated with AD progression [82]. Neurexin 3 protein expression has also been seen to be specifically downregulated in blood samples of AD patients [84].

Another expression analysis on CSF from MS patients identified neurexin  $2\alpha$  levels as a potential biomarker for the disorder [98], while a genetic analysis found that a mutant miRNA, MIR8485, overexpresses neurexin, which leads to a calcium overload in pre-synapses [99]. It was hypothesized that this could induce neurodegeneration in MS [99].

A study examining gene expression in brain tissue samples of patients with PD found that genes related to nerve function, such as protocadherin-8 (PCDH8) and neurexin 3, were downregulated [109].

Two studies on mild cognitive impairment (MCI) found promising results. MCI is a milder form of dementia that is considered the intermediate state of cognitive decline between normal ageing and dementia [114]. Berchtold *et al.* [95] found that neurexin 1 and neurexin 2 are upregulated in MCI. In addition, neurexin 1 expression was found to be associated with longitudinal phenotypes in MCI, but not in AD [96].

One study examined neurexins in order to identify genes that are differentially regulated by HIV encephalitis [94]. This

Alzheimer's disease neuroimaging initiative; AMPA4, GluA4-containing glutamate; CSF, cerebrospinal fluid; EAE, experimental autoimmune encephalomyelitis; ELISA, enzyme-linked immunosorbent assay; FTD-GWAS, frontotemporal dementia genome-wide association study; GEO, gene expression omnibus; GWAS, genome-wide association study; HIV, human immunodeficiency virus; HYPERGENES, European Network for Genetic-Epidemiological Studies; LC-MS/MS, neuronal pentraxin 2; ONIND, other non-inflammatory neurological disease; PCDH8, protocadherin-8; PPMI, Parkinson's Progression Markers Initiative; qRT-PCR, quantitative real-time PCR; RAP-PCR, reverse arbitrarily primed PCR; rMOG, rat Table 2. List of studies that have implicated neurexin genes in neurodegenerative disorders and ageing. 6-0HDA, 6-hydroxydopamine; ACP-RT–PCR, annealing control primer reverse transcriptase–polymerase chain reaction; ADNI liquid chromatography mass spectrometry/mass spectrometry; LC-SRM, liquid chromatography single reaction monitoring; MAP, Rush Memory and Ageing Project; MR, magnetic resonance; MRI, magnetic resonance imaging; NPTX2, myelin oligodendrocyte glycoprotein; RRMS, relapsing-remitting MS; RT-PCR, reverse transcriptase-PCR; SNP, single nucleotide polymorphism; UV-CLIP, ultraviolet cross-linking and immunoprecipitation; WES, whole-exome sequencing.

ference	[0	2]	4] 3]	6]	7] (Continued.)	11 10
main finding	NRXN3 (KIAA0743) is associated with temporal lobe structure in AD [8 patients	the concentrations of the synaptic proteins neurexin 1 and neuronal PTX1, [8 as well as neurofascin, were significantly lowered in AD CSF significantly lower levels of the synaptic proteins NPTX2, AMPA4, [8 neuroligin 1 and neurexin $2\alpha$ in the plasma neuron-derived exomes correlate with AD progression	Ap <sub>42</sub> fibrils in AD CSF are involved in binding to proteoglycans, growth [ <sup>1</sup> factors and neuron-associated proteins, such as neurexin 1, 2 and 3 neurexin 3 expression is downregulated in AD [ <sup>1</sup>	levels of neurexin 2 $\alpha$ and neurexin 3 $\alpha$ , as well as other synaptic proteins [3 are decreased in preclinical AD CSF SYNAPTIC proteins, including neurexin 1, 2 and 3, undergo altered [8 expression in ageing and AD	neurexin 3 is downregulated in AD-transgenic mice with impaired [8 memory, but not in normal aged mice with impaired memory	alsocietypublishing.org/journal/rsob <i>Open Biol.</i> <b>11</b> : 210091
methods	obtained neuroimaging and genetic data from 818 subjects as part of ADNI performed a GWAS with 546,314 SNPs using temporal lobe and hippocampal volume as quantitative phenotypes	collected CF samples from 10 AD patients and 15 healthy controls analysis using LC-MS/MS collected blood and CF samples from 28 AD patients and 28 controls extracted plasma neuron-derived exomes CD81, NPTX2, AMPA4, neuroligin 1 and neurexin 2α proteins were quantified using ELISAs	collected CFF samples from six AD patients and five non-AD patients binding assays were performed to determine which proteins in CFF bind to Aβ <sub>42</sub> fibrils and/or protofibrils selected data from 263 AD patients and 151 non-demented controls sampled from the religious orders study performed RNA expression profiling	collected CSF samples from AD patients and cognitively normal controls (three stage study with different <i>n</i> for each stage) performed LC-MS/MS and LC-SRM obtained frozen brain samples from 26 AD cases and 55 non-AD controls from National Institute on Ageing Alzheimer's disease brain banks used microarrays to evaluate expression profiles of 340 synaptic genes	mice were divided into four groups, with four mice in each group: memory intact AD-transgenic mice, memory impaired AD-transgenic mice, memory intact aged mice and memory impaired aged mice performed proteomics on the hippocampus of each mouse	
type of study	GWAS of brain structure	protein expression analysis protein expression analysis	protein expression analysis transcriptome and RNA expression analysis	protein expression analysis microarray analysis	animal study	
neurexin gene	NRXN3	NRXN1 NRXN2œ	NRXN1, 2 and 3 NRXN3	NRXN2& and NRXN3& NRXN1, 2 and 3	NRXN3	
disorder/disease process	Alzheimer's disease (AD)	AD AD	AD AD	AD AD and ageing	AD and ageing	

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Table	

disorder/disease process	neurexin gene	type of study	methods	main finding	reference
AD and ageing	NRXN1 and 3	microarray analysis	performed a microarray analysis on 47 post-mortem brain samples from cognitively intact aged individuals from the MAP study identified 48 microarrays from the public GEO: 16 young cases, 18 cognitively intact aged cases and 14 AD cases analysed data to identify genes related to physical activity, ageing and AD	neurexin 1 and 3 have decreased expression in ageing and AD but have increased expression in association with late-life physical activity	[88]
ageing aneinn	NRXN3 NRXN2 NRXN2	animal study methylation analysis	cerebella were removed from three adult C57BL/6 J mice and three aged C57BL/6Jnia mice RNA was extracted and sequenced monordas were nurified from PRMCs	neurexin 3 is downregulated in the cerebellum of aged mice	[89]
ageing	NKXN2	methylation analysis	monocytes were purthed from PBMLs analysis of methylation was performed on genomic DNA from monocytes	CpG sites associated with <i>NKP1, NKXN2</i> and mik-29b-2 are hypomethylated in monocytes during ageing	[06]
ageing	NRXN1	animal study	28 Swiss albino mice were divided into four groups by age: young, adult, middle age and old molecular techniques were used to analyse neurexin 1 and neuroligin 3 expression	neurexin 1 and neuroligin 3 are differentially expressed in cerebral cortex and hippocampus during different stages of ageing, which might be responsible for alterations in synaptic plasticity during ageing	[19]
ageing	NRXN2 and 3	transcriptome analysis	collected data of 2202 post-mortem human brain samples of neurologically healthy individuals with different ages Calculated signal expression of genes	neurexin 2 and 3 are downregulated in ageing	[92]
amyotrophic lateral sclerosis (ALS)	NRXN1	cell culture and expression analysis	performed UV-CLIP experiments on SH-SYSY cells to find TDP-43 targets validated these results on lumbar spinal cords from 4 ALS patients and 4 controls using RT-PCR	neurexin 1 and other TDP-43 targets are dysregulated in ALS	[93]
HIV encephalitis	NRXN1	microarray analysis	received cortical brain tissue from 13 HIV patients: eight with HIV encephalitis and five without extracted total RNA performed microarray analysis	neurexin 1 is downregulated in HIV encephalitts	[94]
mild cognitive impairment (MCI)	NRXN1 and 2	microarray analysis	obtained frozen brain samples from 16 MCI cases, 25 AD cases and 24 aged controls from National Institute on Aging Alzheimer's Disease brain banks extracted total RNA performed a microarray analysis	neurexin 1 and 2 are upregulated in MCI	[56]
					(Continued.)

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Table

disorder/disease	neurexin				
process	gene	type of study	methods	main finding	reference
MCI and AD	NRXN1	association study	obtained brain MR images of 400 MCI subjects, 400 AD subjects and 200	neurexin 1 expression is associated with longitudinal phenotypes in MCI,	[96]
			aged controls from the ADNI database	but not in AD	
			obtained genotype data for 510 of these subjects from the ADNI database		
			performed an association study		
multiple sclerosis (MS)	NRXN3	animal study	induced EAE in 17 rats by injecting rM0G	neurexin 3 is downregulated in grey matter of EAE-induced rats	[67]
			six control rats were treated with saline		
			extracted total RNA		
			used a cDNA expression array		
MS	NRXN2 <i>c</i> c	protein expression	collected CSF samples from 37 RRMS patients, 50 patients with ONIND	neurexin $2\alpha$ in CF is a potential biomarker for MS	[88]
		analysis	and patients with non-neurological (orthopaedic) diseases		
			analysis using LC-MS/MS		
MS	NRXN1	genetic analysis	collected blood from a female patient with RRMS	overexpression of neurexin 1 by mutant MIR8485 leads to calcium overload	[66]
			performed WES and screened for mutations	in pre-synapses. This could induce neurodegeneration in MS	
MS	NRXN1	cell culture	treated THP-1 cells with ceramides to induce hypermethylation of DNA	ceramide-induced hypermethylation of DNA was associated with decreased	[100]
			isolated genomic DNA	transcript levels of neurexin 1 in cultured human monocytes	
			measured levels of neurexin 1, FZD7 and TP63 using qRT-PCR		
neurodegeneration	NRXN3	animal study	45 DA(RT1 <sup>av1</sup> ) and 45 PVG(RT1 <sup>c</sup> ) adult rats	neurexin 3 is downregulated in rats with axonal damage caused by ventral	[101]
			subjected rats to ventral root avulsion	root avulsion	
			extracted total RNA		
			used a cDNA expression assay and performed RT–PCR		
neurodegeneration	NRXN3	animal study	three experimental groups with five ICR mice each	neurexin 3 is downregulated in the hippocampus of mice treated with	[102]
			injected kainic acid into ICR mice	kainic acid, an inducer of neurodegeneration	
			extracted total RNA from the hippocampus		
			performed ACP-RT-PCR and sequenced the PCR products		
neurodegeneration	NRXN1	bioinformatics and cell	analysed cross-linking, immunoprecipitation and sequencing data from the	a novel TDP-43 binding miRNA, miR-NID1 (miR-8485), represses neurexin	[103]
		culture	ArrayExpress archive to identify RNAs bound to TDP-43 in human and	1 expression and may play a role in neurodegeneration	
			mouse brains		
			quantitative RT–PCR was used to measure mRNA expression		
					(Continued.)

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Table	

disorder/disease	neurexin				
process	gene	type of study	methods	main finding	reference
neurodegeneration	NRXN1B	cell culture	transfected rat hippocampal neurons to overexpress acetylcholinesterase	excessive glycosylated acetylcholinesterase could competitively disrupt	[104]
			performed a co-immunoprecipitation assay with neurexin 1 and	neurexin 1 $\beta$ -neuroligin junctions and impair the integrity of	
			acetylcholinesterase	glutamatergic synapses	
			co-transfected HEK-293 cells to express neurexin 1 $\beta$ and neuroligin 1 and		
			cultured these cells in acetylcholinesterase conditioned media		
			performed a co-immunoprecipitation assay with neurexin 1ß and		
			neuroligin 1		
neurotoxicity	NRXN3B	animal study	groups of 3 Sprague–Dawley rats were treated with sarin via intra-	sarin exposure causes a persistent downregulation of neurexin $1\beta$ and	[105]
			muscular injection	breakdown of the blood-brain barrier	
			rats were sacrificed 15 min or 3 months after sarin exposure		
			dissected brains and extracted total RNA		
			performed a microarray analysis		
neurotoxicity	NRXN2α	animal study	wild-type zebrafish were repeatedly exposed to domoic acid via	neurexin $2\alpha$ was upregulated in zebrafish two weeks after exposure to	[106]
			intracoelomic injection	domoic acid, a neurotoxin	
			dissected brains and extracted total RNA		
			performed a microarray analysis		
Parkinson's disease	NRXN1	cell culture	cultured SH-SY5Y cells and primary mouse mesencephalic cells	downregulation of neurexin 1 mRNA and protein was observed in the 6-	[107]
(PD)			treated cells with 6-0HDA	OHDA-induced cell culture models of PD	
			performed RAP-PCR and analysed the PCR products using RT-PCR and		
			qRT–PCR		
PD	NRXN2	animal study	transgenic mice were assigned to 4 treatment groups with 20 mice per	transgenic mice overexpressing $lpha$ -synuclein have increased levels of	[108]
			group	neurexin 2	
			cholesterol oximes were administered in food pellets	chronic administration of cholesterol oximes to these mice decreased	
				neurexin 2 levels	
			TH+ neurons were isolated from the substantia nigra and subjected to a		
			transcriptome analysis		
PD	NRXN3	genetics analysis	obtained genomic data of 29 PD samples and 18 controls from the GEO	genes related to nerve function, such as PCDH8 and neurexin 3, are	[109]
			database	downregulated in PD brain tissue samples	
			analysed the data to identify disease-related genes and differential gene		
			expression		
					(Continued.)

	lieurexin				
process	gene	type of study	methods	main finding	reference
PD	NRXN1	animal study	adult Wistar rats were divided into five treatment groups, with 6–8 rats in	neurexin 1 is significantly decreased in the striatum of 6-0HDA-induced	[110]
			each group	rats	
			experimental groups had 6-OHDA brain injections with or without different	treatment with allopregnanolone attenuates this and other molecular	
			concentrations of allopregnanolone	changes	
			western blots were performed to evaluate the levels of the synaptic		
			proteins PSD95 and neurexin 1 in the striatum		
PD	NRXN1	RNA expression	MRI data from 149 PD patients and 64 healthy controls were obtained	neurexin 1 does not have an expression pattern that predicts regional	[111]
		analysis	from the PPMI database	atrophy in PD	
			17 genes of interest implicated in PD were selected for whole-brain		
			expression analysis		
PD	NRXN1	animal study	adult Wistar rats were divided into seven treatment groups, with seven	neurexin 1 expression is decreased in the striatum of 6-OHDA-induced rats	[112]
			rats in each group		
			experimental groups had 6-0HDA brain injections with or without different	6-OHDA rats treated with apelin-13 showed increased neurexin 1	
			concentrations of apelin-13	expression in the striatum	
			western blots were performed to evaluate the levels of the synaptic		
			proteins PSD95, neurexin 1 and neuroligin in the striatum		
spinal muscular	NRXN2α	animal study	used HB9:D3cpv/MN-transgenic zebrafish and Smn-/-/SMN2 mice	SMN-deficiency downregulates neurexin $2lpha$ expression and alters its	[113]
atrophy (SMA)			isolated total RNA from both models	splicing in zebrafish and mouse models of SMA	
			performed a microarray analyses and qRT–PCR		

Table 2. (Continued.)

microarray study showed that neurexin 1 is downregulated in HIV encephalitis.

Finally, González-Velasco *et al.* [92] showed that neurexin 2 and neurexin 3 mRNA levels are downregulated in ageing. Another study found that neurexin 1, 2 and 3 underwent altered expression in both AD and ageing [86]. A more recent study from the same group confirmed decreased expression of both neurexin 1 and neurexin 3 in AD and ageing [88]. Interestingly, they also found that late-life physical activity is associated with increased expression of these proteins [88].

#### 5.1.1. Association study

A GWAS performed by Stein *et al.* [80] showed that the SNP rs7155434 within *NRXN3* is associated with temporal lobe structure in AD patients. Temporal lobe volume deficits are a known risk factor for AD; therefore, this study potentially implicates *NRXN3* with AD risk [80].

## 5.2. In vitro and in vivo models of disease

Several studies involving cell culture and/or rodent disease models have also shown differences in the expression of neurexin proteins. Three studies showed that neurexin 1 is downregulated in PD. One of these measured neurexin 1 mRNA in two 6-OHDA (6-hydroxydopamine)-induced cell culture models; one using human neuroblastoma (SH-SY5Y) cells and the other using primary mouse mesencephalic cells [107]. The other studies used a 6-OHDA-induced rat model of PD and both saw a decrease in neurexin 1 in the striatum [110,112]. In addition, these studies showed that treatment with apelin-13 [112] or allopregnanolone [110] is able to attenuate this change. Apelin-13 is an endogenous ligand for APJ [115] that has been investigated as a potential protective neuropeptide due to the role of the apelin-APJ system in neuronal survival [116], while allopregnanolone is a reduced metabolite of progesterone [117] and has reduced CSF levels in PD patients [118]. Freeze et al. [111], however, noted that the expression pattern of neurexin 1 does not predict regional atrophy in PD. This suggests that neurexin 1 is not a marker for PD; however, it does not exclude it as an important protein in PD pathogenesis. Another study in PD-transgenic mice overexpressing α-synuclein found that neurexin 2 expression was also upregulated [108]. In addition, chronic administration of cholesterol oximes was able to increase the transcription of cytoprotective genes and undo transcriptome alterations, including the alteration of neurexin 2 expression [108].

Two studies using induced models of MS implicated neurexins in this disorder. One study induced experimental autoimmune encephalomyelitis (EAE) in rats and observed downregulation of neurexin 3 [97]. This is a commonly used model that mimics certain aspects of MS. The other study used an *in vitro* model of MS, cultured human monocytes, and observed an association between ceramide-induced hypermethylation of DNA and neurexin 1 mRNA [100].

An animal study performed by Neuner *et al.* [87] showed that neurexin 3 is downregulated in AD-transgenic mice, but not in normal aged mice with impaired memory. However, Popesco *et al.* [89] found that neurexin 3 is downregulated in the cerebellum of aged mice. Another study found that levels of both neurexin 1 and neuroligin 3 are differentially expressed in cerebral cortex and hippocampus of mice and

that these expression levels change during different stages of ageing [91]. They predicted that this may be responsible for the changes in synaptic plasticity observed with age [91]. Finally, a DNA methylation study by Tserel *et al.* [90] showed that CpG sites associated with *NRP1*, *NRXN2* and miR-29b-2 are hypomethylated in monocytes during ageing.

To date, only one study has examined neurexins in amyotrophic lateral sclerosis (ALS) and spinal muscular atrophy (SMA). In a cell culture model of ALS, neurexin 1 and other RNA targets of TDP-43 were dysregulated [93]. TDP-43 is a component of the cytoplasmic inclusion bodies present in ALS patients [93]. Fragments of TDP-43 are ubiquitinated, hyperphosphorylated and then accumulate in neurons and glia [119]. In zebrafish and mouse KO models of SMA, the *SMN*-deficiency downregulated neurexin  $2\alpha$  expression and altered its splicing [113]. SMA is associated with mutation or deletions in the *SMN* gene [120] and lack of the SMN protein causes degeneration and results in anterior horn cell dysfunction.

### 5.3. Models of induced neurodegeneration and toxicity

Several studies investigated neurexins in models of neurodegeneration or toxicity instead of studying a specific neurodegenerative disease.

Four studies examined the role of neurexins in models of induced neurodegeneration. Two of these studies hypothesized that neurexin 1 could play a role in neurodegeneration. The first study showed that a novel TDP-43 binding miRNA, miR-NID1 (miR-8485) is able to repress neurexin 1 and predicted that this could play a role in neurodegeneration [103]. Xiang et al. [104] found in vitro that excessive glycosylated acetylcholinesterase could competitively disrupt the neurexin 1<sub>β</sub>-neuroligin junctions and impair the integrity of glutamatergic synapses, which could lead to neurodegeneration. The other two studies showed that neurexin 3 is downregulated in animal models of neurodegeneration [101,102]. Suh et al. [102] saw that neurexin 3 was downregulated in the hippocampus of mice treated with kainic acid, an inducer of neurodegeneration, while Swanberg et al. [101] found that neurexin 3 is downregulated in rats with axonal damage caused by ventral root avulsion.

Two studies were conducted in animal models of neurotoxicity. One study exposed zebrafish to chronic, low levels of the neurotoxin domoic acid and saw an upregulation of neurexin  $2\alpha$  after two weeks [106]. The other study exposed rats to acute doses of sarin, which caused a persistent downregulation of neurexin 1 $\beta$  and breakdown of the blood–brain barrier [105].

In summary, multiple studies have shown changes in neurexin expression in AD, ALS, MS, PD and SMA. Many of these studies have observed downregulation of protein expression for neurexin 1, 2 and 3 in these disorders. Similarly, downregulation of neurexin 1, 2 and 3 were observed in disorders such as HIV encephalitis and MCI and in studies on ageing, in models of neuronal toxicity, and animal models of MS and ALS.

# 6. Concluding remarks

A clear link between synaptic dysfunction and neurodegenerative as well as neuropsychiatric disorders has been established in recent years. Our literature-based searches revealed several

studies that have linked CNVs, deletions or expression changes in neurexins to different disorders. The evidence is most compelling for a role of neurexins in neuropsychiatric disorders, particularly in regard to the involvement of neurexin 1 in ASD and SCZ. Currently, there is comparatively less evidence for the involvement of neurexins in neurodegenerative disorders. Although there have been some studies that have suggested that neurexins may be important in these disorders, at this stage more experimental data are still needed to draw concrete conclusions. Therefore, it is apparent that more targeted studies in various disorders involving these genes as well as the proteins they encode are warranted. In terms of their broader biological and physiological functions, the neurexins function as molecular inducers, are involved in iron and protein binding, and play a role in cell-to-cell communication and response to stimuli, consequently making them critical for normal cell functioning. Furthermore, these proteins interact with various other proteins such as the neuroligins and the LRRTM proteins identified via protein interaction networks. This implicates the neurexins' involvement in synaptic integrity and functioning making them promising candidates as disease genes for a wide range of brain pathologies.

In summary, this review serves to highlight the potential importance of the neurexin genes and proteins in human disease and recommends that more targeted studies on these genes and proteins are warranted. Furthermore, with the wealth of exomic and genomic sequences and genome-wide transcriptomic datasets now available, it has become plausible to interrogate them for their involvement in various human disorders, on a scale not previously possible. In addition, the human neurexin protein structures urgently need to be solved to understand the function and infer accurate protein– protein interactions as well as to understand the effect of mutations on the protein structure. Ultimately, improved knowledge on synapses and their individual components are necessary to develop novel therapeutic approaches for the emerging and exciting field of synaptopathies.

Data accessibility. This article has no additional data.

Authors' contributions. K.C. performed the research and wrote the first draft of the manuscript. M.H. and R.C. assisted with some of the analysis. K.C. and S.B. conceptualized the study. All authors critically reviewed and edited the manuscript.

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