Diagnostic and Therapeutic Potential of Exosomes in Neurodegenerative Diseases

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Neurodegenerative diseases are closely related to brain function and the progression of the diseases are irreversible. Due to brain tissue being not easy to acquire, the study of the pathophysiology of neurodegenerative disorders has many limitations—lack of reliable early biomarkers and personalized treatment. At the same time, the blood-brain barrier (BBB) limits most of the drug molecules into the damaged areas of the brain, which makes a big drop in the effect of drug treatment. Exosomes, a kind of endogenous nanoscale vesicles, play a key role in cell signaling through the transmission of genetic information and proteins between cells. Because of the ability to cross the BBB, exosomes are expected to link peripheral changes to central nervous system (CNS) events as potential biomarkers, and can even be used as a therapeutic carrier to deliver molecules specifically to CNS. Here we summarize the role of exosomes in pathophysiology, diagnosis, prognosis, and treatment of some neurodegenerative diseases (Alzheimer’s Disease, Parkinson’s Disease, Huntington’s Disease, Amyotrophic Lateral Sclerosis).

Keywords: exosomes, neurodegenerative diseases, pathophysiology, biomarkers, treatment

INTRODUCTION

Neurodegenerative disorders are a group of diseases closely related to brain function. Because brain tissue cannot be histologically examined, we have little understanding of its pathophysiological mechanisms, especially mental disorders. At present, the diagnosis of nervous system disease is mainly based on high-cost neuroimaging and biochemical examination of cerebrospinal fluid. Alzheimer’s Disease, Parkinson’s Disease, Amyotrophic Lateral Sclerosis and Huntington’s Disease are the important parts of the neuropsychiatric diseases. The course of neurodegenerative diseases is long and the onset is slow. Early diagnosis is difficult because of the lack of effective peripheral biomarkers (Cheng et al., 2015; Niu et al., 2020).

The blood-brain barrier (BBB) is a diffusion barrier essential for the normal functioning of CNS, preventing most molecules in the blood from entering the brain, and is composed of endothelial cells, astrocyte end-feet and pericytes (Ballabh et al., 2004). It has been shown that about 98% of small molecules and almost all macromolecules (such as peptides and gene drugs) cannot pass through the BBB, and in contrast, lipophilic small molecules pass more easily (Gabathuler, 2010). Therefore, most molecules must pass through a specific transporter protein or receptor in order to cross the BBB (Gabathuler, 2010). Previous studies have shown that the BBB is disrupted in...
neurodegenerative diseases, with increased permeability that is more favorable for substances to cross (Sweeney et al., 2018). However, in terms of treatment, the BBB still hinders the delivery of drugs to the CNS, making it difficult for drugs to reach the site of CNS injury because they are confined to the peripheral circulation, thus causing more drug side effects (Lakhal and Wood, 2011; Tian et al., 2018). Many researchers have attempted to increase the efficiency of drug delivery by injecting chemicals into the brain to disrupt the BBB, but this approach simultaneously disrupts the tight junctions of the BBB, leading to the invasion of harmful substances and metabolic waste, and disrupts cerebral blood flow levels, further exacerbating symptoms by affecting metabolism and neuroinflammation (Ouyang et al., 2021). How to transport therapeutic drugs more efficiently across the BBB to target sites has become a critical issue to be addressed in the treatment process.

Exosome contains DNA, RNA, proteins, lipids, and other substances, which can be circularly transmitted to adjacent and distant cells. The transfer of exosomes can lead to phenotypic changes of receptor cells (Cheng et al., 2015). In particular, miRNA is involved in a variety of pathological processes, including the growth and metastasis of tumors. Many recent studies showed that it also plays an important role in nerve inflammation (Steinbichler et al., 2017; Li et al., 2018; Pascual et al., 2020). It has been found that the production and release of exosomes depend largely on the characteristics of the mother cell and the target cell (Pegtel and Gould, 2019). The presence of some signature proteins on their surface can interact with other proteins such as integrins, which leads to the transport and fusion of exosomes, and the connection with target cells to function (Rastogi et al., 2021). Therefore, different exosomes have different contents and physiological functions with good specificity and targeting. Due to its peripheral availability and, ability to cross the BBB and superior targeting, it increases the possibility of being used as a biomarker in neurodegenerative diseases, thus manifesting its application in the diagnosis of ND (Wei et al., 2020). It can even be used as a drug carrier in the treatment of diseases (Haney et al., 2015). In conclusion, the study of exosomes in neurodegenerative diseases contributes to the early detection and diagnosis of diseases and provides new methods for the treatment of diseases.

This review summarizes the role of exosomes in some age-related neurodegenerative diseases (Alzheimer's Disease, Parkinson's Disease, Huntington's Disease, Amyotrophic Lateral Sclerosis) and reveals the present the latest research progress.

EXOSOME

Exosomes were first described in 1981 by Trams, E. G et al. (Trams et al., 1981). They are nanoscale vesicles (30–100 nm in diameter) released by different types of cells under specific stimuli, including neurons and glial cells. In the early 1980s, the complex process of exosome generation was proposed, including the formation and release of multivesicular bodies (MVB) (Colombo et al., 2014). However, some researchers have pointed out that exosomes are not limited to one mode of generation, but can also be generated by budding of the plasma membrane (Pegtel and Gould, 2019; Figure 1). Exosomes are widely distributed in saliva, plasma, cerebrospinal fluid, milk, and other body fluids found in a variety of body fluids and contain proteins, nucleic acids, and lipids from mother cells (Geng et al., 2019; Saeedi et al., 2019; Sun et al., 2019; Figure 1). The size, shape, and density of exosomes are determined by the proteins, lipids, and other substances they contain, and are highly variable (Pegtel and Gould, 2019). Escola et al. found that co-stimulatory molecule CD86 and several tetraspan proteins (including CD37, CD53, CD63, CD81, and CD82) were highly enriched in exosomes and correlated with their immune effects (Escola et al., 1998), with CD81 and CD63 having become the most commonly used exosome marker proteins (Hemler, 2003).

In addition to this, exosomes contain many integral membrane signaling proteins, including epidermal growth factor receptor (EGFR), mast/stem cell growth factor receptor (c-Kit), vascular endothelial growth factor receptor type-2, insulin-like growth factor I receptor, T cell receptor, cytokine receptors, G protein–coupled receptors (GPCRs), Notch receptors and so on, which can act as surface signaling molecules for cellular transmission of functional receptors and signaling pathways (Pegtel and Gould, 2019). In addition, the lipid content of exosomes is not the same as that of their mother cells, with cholesterol, sphingomyelin, glycosphingolipids and phosphatidylserine being enriched in exosomes two to three times as much as in their mother cells (Skotland et al., 2017).

Exosomes have an important biological role in vivo (Pegtel and Gould, 2019), and some researchers have categorized this into the following three points: (1) The organism can selectively remove some protein substances from the plasma membrane through the pathway of exosome generation, such as the vesicular secretion of neurodegenerative amyloidogenic proteins (Quek and Hill, 2017). (2) Exosomes are important components of the extracellular matrix that mediate amyloid aggregation, plaque and tangle formation, growth and spreading in neurodegenerative disease. (3) Exosomes mediate intercellular signaling and messaging. Exosomes can be internalized by other cells through membrane fusion, endocytosis, or specific phagocytosis. By transferring proteins and genetic materials (such as mRNA, miRNA, rRNA, long-chain non-coding RNA, DNA, etc.) to other cells, it can change the function of receptor cells and mediate cell-to-cell communication. And bilayer lipid membrane effectively protects its contents from degradation (Wang et al., 2018; Geng et al., 2019; Koteswara et al., 2019). Exosomes contain a variety of biological substances, among which miRNAs are highly enriched in exosomes. And most miRNAs that can be obtained from serum or saliva are contained in exosomes. Exosome-derived miRNAs are considered to be more stable than cell miRNAs (Wang et al., 2018). The expression profile of exosomal miRNAs is not identical to that of mother cells, and its expression can be changed according to the change of disease status. In recent years, it has been considered as a potential biomarker (Saeedi et al., 2019). More and more attention has been paid to the role of exosomes in neurodegenerative diseases (Fries and Quevedo, 2018). Especially
for neurodegenerative diseases, the study of exosomal miRNAs has attracted great attention.

The ability to cross BBB (Samanta et al., 2018; Gomez-Molina et al., 2019) makes it possible for exosomes to enter the brain as a drug delivery carrier. Also, due to the targeting effect and immune resistance of exosomes (Bunggulawa et al., 2018), they can further reduce the drug side effects due to drug retention in the periphery during transport as drug carriers,
further increasing the drug delivery benefits. In recent years, they are a promising drug carrier. In many types of cancer, whether in vivo or in vitro, it has been shown that they can carry drugs and target delivery to reduce the adverse reactions of drug treatment (Yang et al., 2017; Kalimuthu et al., 2018). While many preclinical studies have demonstrated the potential of exosomes to treat disease, only a handful of companies are currently conducting relevant clinical studies (Perocheau et al., 2021): Codikai Biosciences works to develop therapeutic exosomes carrying siRNAs targeting the KRAS (G12D) mutation in pancreatic cancer (Mendt et al., 2018); Avalon Globocare is developing engineered exosome therapeutics (Wang et al., 2019b).

In conclusion, exosomes, as natural carriers for transferring bioactive molecules between cells, are characterized by low immunogenicity, strong biodegradability, ability to wrap endogenous bioactive molecules, and ability to cross BBB, and are considered as a new endogenous drug delivery system (Sun et al., 2019; Zhang et al., 2019). In the nervous system, exosomes mediate cell-to-cell communication and are thought to be closely related to learning and memory, neuroinflammation, and other aspects (Wang et al., 2018; Saeedi et al., 2019).

**EXOSOMES AND ALZHEIMER’S DISEASE**

Alzheimer’s disease (AD) is a neurodegenerative diseases characterized by decreased levels of amyloid-beta (Aβ) (reduced levels of Aβ in the cerebrospinal fluid due to deposition of Aβ in the brain), increased levels of total tau or phosphorylated tau, and a reduction in the number and function of synapses (Palop et al., 2006). The increased Aβ aggregates into soluble oligomers to activate microglia to produce an inflammatory reaction and oxidative stress. Excessive Aβ produces a cascade reaction to make neurons degenerate. The abnormal phosphorylation of tau protein can form nerve fiber tangles, which leads to a decrease in neuron function and even neuronal apoptosis (Palop et al., 2006). In recent years, a large number of studies have shown that exosomes are closely related to the occurrence and progression of AD, bringing hope for the treatment (Figure 2).

In 2006, some studies found that exosome is associated with the release of Aβ and can aggravate the brain pathology of AD by promoting the aggregation of Aβ (Rajendran et al., 2006; Dinkins et al., 2016). It can be used as a carrier to transfer toxic substances (such as phosphorylated tau protein and Aβ) between neurons (Kaur et al., 2021). Inhibiting exosome synthesis can significantly reduce the proliferation of tau protein (Wang et al., 2017; Sardar Sinha et al., 2018; Winston et al., 2019). However, the cellular origin of the active exosomes has not been determined, and both microglia and astrocytes have been proposed (Asai et al., 2015; Rosas-Hernandez et al., 2019). In addition, it has been proposed that with the progression of AD, the level of functionally specific synaptic proteins in plasma neural-derived exosomes (NDE) decreases. And synaptic proteins in NDE may be effective preclinical indicators and progression indicators of AD (Goetzl et al., 2016, 2018).

In recent years, there have been many studies on AD biomarkers in exosomes in CSF, blood, and in vitro cultures, mainly involving proteins and miRNAs (Table 1). In terms of proteins, HSP70 (Goetzl et al., 2015; Chanteloup et al., 2019) has received a great deal of attention in addition to the various forms of Aβ extracted in various body fluids (Watson et al., 2019). In addition, the findings of exosomal miRNAs as biomarkers are variable and lack uniform and authoritative conclusions.

In treatment, because exosomes can cross the BBB, its application in targeted therapy has attracted much attention. Relative to conventional AAV (adeno-associated virus), exo-AAV (AAV combine with exosomes) is more efficient at gene delivery to the brain at low vector doses. The ability of exo-AAV of evading neutralizing antibodies and transducing CNS after peripheral delivery makes it electively ingested by neurons (Hudry et al., 2016; Perets et al., 2019). In addition, some researchers combined curcumin and exosomes to inhibit tau phosphorylation and activate GSK-3/ AKT pathway, to prevent neuronal death and relief symptoms (Wang et al., 2019a).

Besides, exosomes injected into the mouse models from different cells play different roles in treatment. For example:

1. The exosomes from mesenchymal stromal cells (MSCs) or hypoxia-preconditioned MSCs could improve cognitive function (the learning and memory) by saving synaptic dysfunction and regulating inflammatory response through regulation of miR-21 (Cui et al., 2018).

2. Treatment with exosomes from adipose-derived stem cells (ADSC-Exo) resulted in decreased Aβ42 and Aβ40 levels, increased apoptotic molecule levels (such as p53, pro-caspase-3, decreased Bcl-2 protein), and decreased apoptosis of neurons. In addition, neurite growth was also increased during the treatment (Lee et al., 2018).

3. Exosomes from human umbilical cord mesenchymal stem cells showed the ability to repair cognitive dysfunction, help clear Aβ deposition in the brain, and reduce neuroinflammation and relief symptoms by regulating the activation of mouse brain microglia cells (Ding et al., 2018).

**EXOSOMES AND PARKINSON’S DISEASE**

Parkinson’s disease (PD) is the second most common neurodegenerative diseases in the world after Alzheimer’s disease (Lebouvier et al., 2009). The main pathological changes were degeneration and death of dopaminergic neurons in substantia nigra, a significant decrease of DA in the striatum, and eosinophilic inclusion bodies appearance in the cytoplasm of residual neurons in substantia nigra, namely Lewy body, α-synuclein (α-syn) is its main ingredients. Recent studies have shown that exosomes play a certain role in the pathogenesis, diagnosis, and treatment of PD (Howitt and Hill, 2016).

As an important component of the Lewy bodies, α-syn are transferred from intracellular to extracellular by being packaged into exosomes or directly released (Lee, 2005). Exosomes provide
an ideal environment for α-syn polymerization (Grey et al., 2015). Exosomes can transfer toxic forms of α-syn between a variety of cells, such as astrocytes and microglia, which can lead to the exacerbation of PD. α-syn deposited in glial cells induces inflammation and its further transmission promotes the degeneration of neurons and exacerbates the development of PD (Chistiakov and Chistiakov, 2017). α-syn in NDE is considered an important biomarker of PD and associated with deterioration of motor symptoms (Si et al., 2019; Niu et al., 2020). Some researchers have found that the content of α-syn in NDE of non-tremor patients is higher than that of tremor patients, so it may also be used to identify different types of motor types in PD (Si et al., 2019).  

In terms of biomarkers, the value of α-syn in exosomes in diagnosis has been pointed out in many studies (Si et al., 2019; Fu et al., 2020; Niu et al., 2020; Zheng et al., 2021), and it has also been shown that exosomal α-syn can be used to differentiate PD from multiple system atrophy (Dutta et al., 2021). There are some other studies on exosomes in serum and CSF of PD patients (Table 2).

In terms of treatment, exosomes can be used as drug therapy carriers for PD and have a natural brain targeted ability. By using the blood exosomes as a delivery system, the distribution of dopamine in the brain has increased more than 15-fold. Compared with free dopamine after intravenous administration, dopamine-loaded exosomes show better therapeutic efficacy and lower systemic toxicity in a PD mouse model (Qu et al., 2018). Otherwise, the release of catalase can reduce cell death by protecting neurons from oxidative damage. In vitro and in vivo trials of PD, therapeutic catalase mRNA delivery by exosomes attenuated neurotoxicity and neuroinflammation (Haney et al., 2015; Kojima et al., 2018).

### TABLE 1 | The study of exosome contents in AD biomarkers.

| Molecule | Object | Source | Expression | Potential target/mechanism | Significance | References |
|----------|--------|--------|------------|-----------------------------|--------------|------------|
| Cathepsin D, LAMP-1, ubiquitinylated protein | Human | serum | Up | Autophagic-lysosomal dysfunction | It is a pathological change that can appear 10 years before the onset of AD. | Goedt et al., 2015 |
| HSP70 | Human | serum | Down | HSP70 acts on proteins that accumulate in the brain. | May mark the extent of synaptic dysfunction or neurodegeneration. | Goedt et al., 2015; Chanteloup et al., 2019 |
| SNAP-25 | Human | serum | Down | | It is related to the disease progression of AD and directly reflects the characteristic of synaptic loss during the progression of AD. | Agiardi et al., 2019 |
| miR-9-5p, miR-598 | Human | CSF | Down | It plays a potential regulatory role in amyloid proteins, stress pathways, and neurotrophic signaling. | These miRNAs may be potential biomarkers for AD. | Riancho et al., 2017 |
| miR-342-3p, miR-342-5p, miR-150-5p, miR-23b-3p, miR-25b-3p | Human | serum | Down | | These miRNAs are collectively altered in the disease, rather than being a single biomarker. | Lugli et al., 2015 |
| miR-124, miR-146a, miR-155, miR-21, miR-125b | - Culture of cell | Up | | These miRNAs can be transported from cell to cell via exosome form to mediate mRNA transcription and aggravate the inflammatory response. | MIR-21 plays an important role in signal transduction between microglial cells and neurons, especially in neuroinflammation. | Fernandes et al., 2018 |
| hsa-miR-101-3p | Human | serum | Up | Hsa-miR-101 can target and regulate APP mRNA, thereby reducing APP level in hippocampal neurons and promoting Aβ accumulation. Hsa-miR-1306 can target to regulate APP mRNA and increase the synthesis of APP. The down-regulation of HSPA1-MIR-106b is associated with transforming growth factor-β signaling. | Using differential miRNAs to make a random forest model for clinical classification prediction has high sensitivity and specificity. | Cheng et al., 2016 |
| miR-125b-5p, miR-451a, miR-605-5p, miR-16-5p | Human | CRF | Down in the first stages of the disease and increase in moderate and advanced stages, Down in the Young-onset AD. | Overexpression of miR-125b-5 leads to tau hyperphosphorylation and neurotoxicity and MIR-451a plays a role in neuroinflammation. | MIR-16-5p is differentially expressed in late-onset AD and Young-onset AD and may be a special biomarker of Young-onset AD. | McKeever et al., 2018 |
TABLE 2 | The study of exosome contents in PD biomarkers.

| Molecule        | Object | Source  | Expression | Potential target/mechanism                                                                 | Results                                                                                                                                                                                                 | References               |
|-----------------|--------|---------|------------|------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------|
| miR-19b, miR-195, miR-24 | Human  | serum   | Down       | Parkin RBR E3 ubiquitin protein ligase (miR-19b), LRRK2/PARK8 (miR-19b), and ATP13A2/PARK9 (miR-24 and miR-195) | ROC curve was used to evaluate the combined diagnostic value of the three miRNAs: AUC was 0.946 (85%CI, 0.910-0.981).                                                                                       | Gao et al., 2017          |
| miR-153, miR-409-3p, miR-10a-5p, let-7g-3p, miR-1 and miR-19b-3p | Human  | CSF     | Up         | Neurotrophin signaling, mTOR signaling, Ubiquitin mediated proteolysis, Dopaminergic synapse, Glutamatergic synapse were the most prominent pathways | The sensitivity and specificity for distinguishing Parkinson’s disease from control were 94% for miR-1, 93% for miR-153, 90% for miR-409-3p, 94% for miR-19b-3p, 95% for miR-10a-5p, and 95% for let-7g-3p. | Gui et al., 2015          |
| miR-505, miR-331-5p | Human  | serum   | Down       | –                                                                                          | The ROC curve analysis AUC values of miRNA-331-5p and miR-505 were 0.849 and 0.898, respectively.                                                                                                    | Yao et al., 2018          |
| prion protein   | Human  | serum   | Up         | PrPC can increase phosphorylated α-synuclein and induce synaptic damage and calcium homeostasis. | The level of prion protein in PD plasma exosomes was significantly correlated with the level of cognitive impairment. (t = -3.185, P = 0.001)                                                                 | Leng et al., 2020         |
| DJ-1            | Human  | serum   | Up         | DJ-1 can promote disease progression by regulating α-synuclein cytotoxicity.                 | The ROC analysis results of DJ-1 in PD plasma neurogenic exosomes were as follows: AUC = 0.703, sensitivity = 79.5%, specificity = 57.5%.                                                          | Zhao et al., 2018         |

Moreover, the study on exosomes also provided a theoretical basis for other known causes of PD and clarified the specific pathological mechanism. For instance:

(1) Glucocerebrosidase gene (GBA) mutation is the most common genetic pathogenic factor in PD and is associated with decreased glucocerebrosidase activity in PD. Overexpression of GBA in vitro resulted in significantly reduced exosome secretion which is associated with the α-syn oligomers. And the reduction of glucocerebrosidase activity in vivo induced a significant increase in the number of exosomes released in the brain, thus promoting the pathological changes of PD (Papadopoulos et al., 2018).

(2) Manganese exposure is considered to be an important factor in the susceptibility of PD. It can increase the protein Rab27a and some miRNAs that regulate the release of exosomes, leading to protein aggregation, autophagy, inflammation, and hypoxia, thereby potentially promoting progressive neurodegeneration (Harischandra et al., 2018). In addition, manganese exposure promotes the secretion of α-syn in exosomes, accelerates the intercellular transmission of α-syn, and leads to dopaminergic neurotoxicity, thus causing proinflammatory and neurodegenerative reactions (Harischandra et al., 2019).

(3) Mutations in leucine-rich repetitive kinase 2 (LRRK2) enhance the level of self-phosphorylated LRRK2 protein, the most commonly known cause of hereditary PD. The level of Ser(P)-1292 LRRK2 in Urinary exosomes were found to be elevated in idiopathic Parkinson’s disease and correlated with the severity of cognitive impairment and difficulty in daily activities (Fraser et al., 2016b). Its proportion in total LRRK2 predicted LRRK2 mutation status in LRRK2 mutation carriers (Fraser et al., 2016a).

EXOSOMES AND HUNTINGTON’S DISEASE

Huntington’s disease (HD) is an autosomal dominant neurodegenerative disorder. In 1993, the international Huntington’s Disease Cooperative Research Group cloned the disease-causing gene IT15, its metabolite mutant Huntington’s protein (mHTT) which has many repeated glutamines, is easy adhesion, aggregation, and eventually lead to the death of nerve cells. Exosomes can carry and spread mHTT between cells, triggering HD-related behaviors and pathological performance (Jeon et al., 2016).

Previous studies in different HD mouse models have shown that exosomes from astrocytes (ASC-EXO) carry heat shock proteins and other neuroprotective substances, which can reduce the cytotoxicity of misfolded proteins and prevent neurodegeneration. And dysfunction of astrocytes can lead to neuronal vulnerability (Hajrasoulia et al., 2013; Nafar et al., 2016). Although mHTT does not exist in ASC-EXO, it can reduce the secretion of ASC-EXO in HD mice (Hong et al., 2017). The specific possible mechanism is as follows: (1) The n-terminal of mHTT can form aggregation in the nucleus, leading to the decrease of exosomes in astrocytes; (2) mHTT reduces the expression of the small heat shock protein α-B-crystallin [α protein mainly expressed in glial cells, mediating exosome secretion (Gangalum et al., 2016)] in astrocytes, thus reducing the secretion of exosomes in the brain and promoting the acellular autonomic neurotoxicity of HD. And when ASC-EXO is injected into the striatum of HD mice, the density of mHTT aggregation could be reduced and the overexpression of α-B-crystallin could reduce the exosome deficiency and neuropathological changes, which provided ideas for the treatment of Huntington’s disease (Hong et al., 2017).

It has been found that ASC-EXO could significantly reduce mHTT aggregation of nerve cells, up-regulate the expression of PGC-1 and phosphorylated CREB, and reduce mitochondrial...
dysfunction and cell apoptosis, indicating that ASC-EXO has the potential to treat HD (Lee et al., 2016b). In addition, it has been found that exosome-mediated hydrophobic modification of siRNA can silence HTT mRNA, which is expected to promote the development of treatment methods for Huntington’s disease and other neurodegenerative diseases (Didiot et al., 2016). Some researchers have injected exosomes loaded with excess miR-124 into the striatum of HD model mice in an attempt to improve HD-like behavior, and although the results were not satisfactory, this approach still resulted in downregulation of the target gene RE1-Silencing Transcription Factor (Lee et al., 2017). Some researchers recently found that transferring serum exosomes from young mice into an in vitro model of HD can effectively ameliorate mHTT mutations, slow down apoptosis, and promote mitochondrial biogenesis, while the shared blood circulation through parabiosis experiment confirmed the above idea (Lee et al., 2021).

EXOSOMES AND AMYOTROPHIC LATERAL SCLEROSIS

Amyotrophic lateral sclerosis (ALS) is caused by progressive weakness and atrophy of the muscles innervated by the medulla oblongata and muscles in the limbs and trunk after injury to upper and lower motor neurons. According to family history, it can be divided into sporadic ALS (sALS) and familial ALS (fALS), among which only 5 ∼ 10% are fALS. So far, more than 20 genes related to the pathogenesis of ALS have been found, among which superoxide dismutase gene (SOD1) and ubiquitinated TAR-DNA binding protein 43(TDP43) are the most studied (Chen et al., 2019).

Some ALS are caused by the misfolding of mutated SOD1, and the misfolded SOD1 is transferred through exosome dependent or independent way. In cell culture, once SOD1 misfolding happens, it can still induce misfolding of the immature cell after the initial misfolded template has been degraded for a long time (Gomes et al., 2007; Grad et al., 2014a,b; Silverman et al., 2016). Other researchers have found that NDE in the brain of ALS can cause the cytoplasmic redistribution of TDP-43, suggesting that exosomes may be involved in the transmission of TDP-43 protein lesions (Yohei et al., 2016). By analyzing serum miR-27a-3b in ALS and healthy subjects, Xu Qian et al. found that exosomal miR-27a-3b expression was down-regulated in ALS, which may serve as a detection indicator for ALS, but the exact mechanism is unclear (Xu et al., 2018).

Noriko Hayashi et al. analyzed the proteomics of exosomes from patients with sALS, it is found that three proteins increased and 11 proteins decreased in exosomes of ALS patients. The most increased protein was a new that inhibitor (NIR), which was closely related to nucleolar function. The decrease of NIR in the motor neuron nucleus of ALS patients suggested that nucleolar stress might play a role in the pathogenesis of sporadic ALS through NIR dysfunction (Hayashi et al., 2019). Chen et al. (2019) found that the level of IL-6 in ASC-Exo of sALS patients was increased, which was positively correlated with the rate of disease progression (only for patients with disease course less than 12 months), indicating that inflammation of CNS was increased, and exosomes derived from nerve cell might help to reveal the neuroinflammation of CNS of ALS patients. Besides, exosome proteins have been proved to be mainly involved in the negative regulation of cell adhesion and apoptosis. In the exosomes of ALS patients, the pro-apoptotic protein Bax and caspase-3 are down-regulated, and the antiapoptotic protein Bcl-2α is up-regulated. The protein content is related to the antiapoptotic effect of exosomes (Bonafede et al., 2019). Moreover, TDP-43 from the exosomes of patients can induce the increase of monocytes, which may increase the neuroinflammatory effect (Zondler et al., 2017). In addition, exosomes mediated the interaction mechanism between muscle and bone at the cellular level, promoted the mineralization of osteoblasts, participated in the occurrence and development of ALS, and had potential reference value for the clinical diagnosis of ALS (Xu et al., 2018).

In the treatment of ALS, ASC-Exo has been shown to save mitochondria-related dysfunction, which can reduce intracellular SOD1 aggregation, regulate the cell phenotype of ALS, and can be used as a candidate drug for ALS treatment (Lee et al., 2016a). Investigators treated an in vitro cell model of ALS with ASC-Exo and found that ASC-Exo rescued mitochondrial dysfunction and the mitochondrial membrane potential, thus suggesting that ASC-Exo may be used to treat diseases characterized by mitochondrial adaptations, such as ALS (Calabria et al., 2019). Subsequently, Roberta et al. tried to inject ASC-Exo into ALS mice transgenic for SOD1 (G93A) and found that this treatment significantly improved motor performance, protected lumbar motor neurons, neuromuscular junctions and muscles in ALS mice (Bonafede et al., 2020).

DISCUSSION

Although there are a large number of studies on exosomes in nervous system disease, mainly focusing on the pathophysiologcal mechanism, the disease development, biomarkers, and treatment (as carriers or themselves), in addition to the biomarker studies, most of the rest in the nervous system research is in the animal model or in vitro cell culture data, especially in the treatment. Applying exosomes to therapeutic safety and technical issues is a major challenge (Chen and Chopp, 2018): (1) Cell culture conditions and storage methods may have a significant impact on the contents and functions of exosomes, which requires standardization of exosome extraction methods, storage, and functional read-out systems; (2) The content, function, and activity of exosomes depend on the generating cells of origin. So, it is necessary to optimize exosome cell sources, including age, gender, comorbidities, and other factors related to the exosome-generating cells; (3) As a therapeutic method, exosomes are mainly focused on functional aspects, but the negative effects are rarely studied. In previous studies, we can see that exosomes play a certain role in promoting neurogenesis, inhibiting neuroinflammation, promoting angiogenesis, and promoting synaptic plasticity. However, in the treatment of a disease, not all the effects are meaningful for the treatment, some even bring great risks, such as whether the treatment
of other non-tumor diseases will increase the risk of cancer promotion; (4) Exosome therapy is still in the preclinical trial stage. Although it can relieve symptoms, it is still a problem whether it can promote the prognosis.

In a word, exosomes bring hope for patients with neurodegenerative diseases, which has great clinical research value, provides a new method for the diagnosis and treatment of diseases, and better clarifies the pathophysiological mechanism.

AUTHOR CONTRIBUTIONS
PG was responsible for reviewing the literature and writing the manuscript. XL and XD participated in providing ideas for the article. SL and YX participated in revising the manuscript. All authors contributed to the article and approved the submitted version.

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