Proposal and point of view on targeting α-synuclein for the treatment of
 Parkinson's disease

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5 Abstract: Many scientific studies in the biochemical, genetic fields suggest that there 6 were common mechanisms, such as genes, α -synuclein protein, tau protein, oxidative 7 stress, mitochondrial dysfunction, and iron might be shared in Alzheimer disease (AD) 8 and Parkinson disease (PD). α -synuclein is suggested to have a vital role in the 9 pathogenesis of PD and is a promising therapeutic target. However, gap might always 10 exist between clinical and basic researches. The failure of recent phase III trials of the 11 anti-Amyloid- β (A β) monoclonal for AD prompts us to rethink PD therapy strategies. 12 As multiple mechanisms are involved in PD pathogenesis and their relative roles 13 might vary at different stages of this disease. Use of comprehensive prevention 14 strategies and targets at different stages of PD might be a promising way to cure or 15 prevent PD in the future.

16 Key words: Alzheimer disease; Parkinson disease; α-synuclein; Neurodegeneration;
17 Amyloid-β;

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20 1.Some common mechanisms might be shared in Alzheimer's Disease and

21 Parkinson's Disease

22 Alzheimer disease (AD) and Parkinson disease (PD) are both neurodegenerative 23 diseases, whose symptoms and pathological changes overlap each other in their late 24 phase. Whether there are common genetic risks, as well as the existence of linkage 25 disequilibrium, epigenetics and other mechanisms between AD and PD is still 26 controversial. It needs large-scale genome-wide association studies (GWAS) of 27 different mutual authentication in the future. In terms of mechanism, AD and PD all 28 belong abnormal protein folding diseases. Previous paper in Science reported that 29 Amyloid- β (A β) of AD and α -synuclein of PD as well as other abnormally folded 30 proteins had the same spatial conformation. Many mechanisms such as genes, α -31 synuclein protein, tau protein, oxidative stress, mitochondrial dysfunction, iron, and 32 locus coeruleus, involved in AD and PD may be the same. There were common 33 mechanisms, which might be shared in AD and PD, were supported by many 34 scientific studies in the biochemical, genetic, and molecular fields [1]. It is presumed 35 that AD and PD might have a common upstream pathogenic pathway.

36 2.α-synuclein provides us with hope and confidence for the treatment of PD

37 α -synuclein is a 140 amino acid neuronal protein that has been associated with 38 several neurodegenerative diseases. Many mutations in the gene coding for the α -39 synuclein protein have been identified in familial and sporadic PD since the first 40 mutation has been found in a rare familial form of PD. α -synuclein were discovered to 41 be predominantly component of the aggregated proteinaceous inclusions called Lewy 42 body (LB) found in PD and cortical Lewy body dementia (LBD). Aberrant 43 aggregation of α -synuclein has been detected in an increasing number of 44 neurodegenerative diseases, collectively known as synucleopathies. α -synuclein exists 45 physiologically in both soluble and membrane-bound states, in unstructured and α -46 helical conformations, respectively.

47 It is well known that LB is the characteristic pathological sign of PD, and fibrous 48 aggregates of α -synuclein are the main elements of LB. As the same core of A β in AD, 49 the position and roles of α -synuclein in the etiology of PD are not easy to be shaken. 50 A subsequent series of cell culture and animal experiments had confirmed the 51 affirmative role of α -synuclein in the pathogenesis of PD [2]. Many academics had 52 suggested that therapeutic strategies might target on α -synuclein for the treatment of 53 PD, which indicate that α -synuclein appears to be a very promising targets for PD. It 54 has also been discovered that duplication or triplication of the wild type a-synuclein 55 gene itself can lead to a familial form of PD [3-5]. This evidence suggests that 56 overexpression of the normal, wild type α -synuclein protein itself can lead to the 57 development of PD. α -synuclein might lie in the upstream of PD pathological process, 58 which mainly exist as three kinds of monomers, oligomers, and polymer [6]. 59 Wakabayashi found that the mainly toxic effects come from soluble oligomers, which 60 suggests that α -synuclein oligomers may be a key factor in the pathogenesis of PD. 61 Many studies have shown that α -synuclein forming LB may have dual effects. The 62 formation of LB is believed to have a protective effect (Proctective sinks), which is 63 conducive to α -synuclein oligomers fibrosis and thus reducing its toxicity effect, 64 preventing nerve cells from erosion due to α -synuclein toxicity. With the increasing of 65 α -synuclein aggregation, the aggregated α -synuclein might exhibit prion-like effect, 66 which can adopt a self-propagating conformation that causes neurodegeneration. 67 Recent evidence now suggests the possibility that α -synuclein is a prion-like protein 68 and that PD is a prion-like disease. α -synuclein in an aberrantly folded, β -sheet-rich 69 form could migrate from affected to unaffected neurons [7]. Laboratory studies also 70 confirm that α -synuclein can transfer from affected to unaffected nerve cells, where it 71 appears that the misfolded protein can act as a template to promote host α -synuclein

misfolding. Immunization therapy with human α -synuclein could reduce α -synuclein aggregate formation and attenuate neurodegeneration in human α -synuclein transgenic mice [8]. A subsequent series of experiments in vivo and vitro suggested that antibodies against α -synuclein reduce cell-to-cell transfer of the protein by directing extracellular α -synuclein to microglia [9].

77 3.Gap might always exist between clinical and basic researches

78 However, the clinical practice tells us that there often exists far distance between 79 the basis pathogenesis and clinical outcomes of the disease. There is not always 80 concordant between them. So that the results coming from basic and clinical research 81 often appear to be different, even contradictory. Just as a prerequisite for evidence-82 based medicine, medicine-based evidence is always in overwhelming status in clinical 83 practice. However, sometimes there exists paradox between evidence-base medicine 84 with practice-base evidence. For example, A β is suggested to have a vital role in the 85 pathogenesis of AD and is a major therapeutic target. Recently, phase III trials of two 86 monoclonal antibodies against A β , bapineuzumab and solanezumab, failed to 87 significantly improve clinical outcomes in patients with mild to moderate AD [10,11].

88 4. Rethinking the strategies of PD therapy

89 Overproduction and clearance disabilities of α -synuclein become an important 90 cause of PD pathogenesis. Therefore, there is no doubt to target on α -synuclein for the 91 treatment of PD. However, just like the pathology of AD, once PD pathological 92 process started, subsequent pathological reactions would gradually increase, forming 93 a vicious cycle of their own, which includes: increasing the permeability of the lipid 94 bilayer, destroying the integrity of the synaptic vesicles, interfering dopamine 95 metabolism, and axoplasmic and vesicles transportation of intracellular substances

96 from the endoplasmic reticulum to the golgi, mitochondria damage, inhibiting the 97 proteasome activity, molecular chaperone mediated autophagy and histone acetylation, 98 leading to proteasome degradation resistance, causing hyperphosphorylation of tau 99 protein, oxidative stress and neuroinflammation [12]. Most patients diagnosed with 100 PD in clinical practice have been in the middle or late stage of the disease. If only 101 targeting on α -synuclein for the treatment of PD, it is difficult to obtain an ideal effect. 102 Therefore, early diagnosis is critical for PD treatment. The further studies on the 103 prevention and treatment for PD should focus on the following points: Firstly, aiming 104 at PD high-risk population screening and early diagnosis of PD, finding out method 105 and the strategy for detection of PD high-risk groups and patients in the premotor 106 phase of PD, on the base of which, the PD diagnosis standard might be revised in the 107 future. The development of PD can be regarded as a continuous evolution process 108 consisting of the preclinical stage or asymptomatic stage (including the pre-109 physiological phase), premotor phase, the early motor symptoms (the pre-diagnostic 110 phase) and motor symptoms stage (the diagnostic phase).

111 Research shows that the brain has begun to appear relevant pathological damage 112 before motor symptoms of PD appear [13]. How to predict and diagnose PD in 113 preclinical stages of PD without symptoms is an important prerequisite for early 114 prevention of PD. Second, for patients in the middle and late stage, comprehensive 115 treatment strategies aiming at more aspects of this disease is needed. Except for 116 blocking the core pathological processes of PD, it should be endeavor to reconstruct 117 the functional balance of the neurotransmitter control loop in the brain of PD patients 118 based on existing pathology, that is called "rebuilding a new home on the ruins." 119 Third, we should actively explore effective exercise rehabilitation therapies such as 120 occupational therapy, physical therapy, and Tai Chi style motortherapy. These are

121 perhaps wonderful strategies which could improve the life quality of patients in the 122 middle and late stage. From the perspective of PD prevention, it can be mainly 123 targeted on α -synuclein production and clearance. While once the onset of PD occurs, 124 from the perspective of therapy, multiple therapeutic targets for comprehensive 125 treatment are needed. It is afraid that single target treatment is difficult to achieve the 126 desired results.

127 Inflammation is a common pathological process of various diseases, which are 128 also involved in the development and progression of PD [14]. Previous studies 129 showed that chronic inflammation in the body including α -synuclein production and 130 tau protein hyperphosphorylation could promote the occurrence and the development 131 of PD. PD pathological process will promote intensify of inflammatory response, 132 including production of inflammatory mediators and activation of glial cells. 133 Inflammation and α -synuclein could form a vicious circle, promoting and intensifying 134 each other. Suppression of the nervous system inflammatory response would 135 contribute to cut off α -synuclein-inflammation vicious cycle. Recently, researchers 136 have found that autoimmune mechanisms are involved in the pathogenesis of PD [15, 137 16], which suggests that the generalized inflammation is indispensable for the PD 138 pathogenesis process. Previous study has found that nonsteroidal anti-inflammatory 139 drugs (NSAID) for the treatment of PD has a certain significance [17,18]. Therefore, 140 as one of the comprehensive treatment strategies of PD, anti-inflammation should be 141 necessary and promising.

142 5. Looking forward the PD therapy strategies

How to successfully cure PD? The current evidence suggests that anti-αsynuclein therapies, as a preventative measure, should be given in the early stage of

145 the disease. Being different from A β to AD, α -synuclein has not yet been discovered 146 there exists a dynamic equilibrium conditions in the central and peripheral system. 147 Researchers found that the plasma levels of α -synuclein were low in normal subjects 148 and patients with PD [19]. There seemed no necessarily link between plasma levels of 149 α -synuclein with disease onset in PD patients. Therefore, not like AD, we could not 150 reach the purpose of the central α -synuclein removal by eliminating the peripheral α -151 synuclein. However, studies have found that exogenous α -synuclein is more likely to 152 be aggregated in the plasma of PD patients. It seems to imply that there might have 153 some factors which could promote α -synuclein overproduction and aggregation in the 154 internal environment of PD. Therefore, for the prevention and treatment of PD, on the 155 one hand, we should not just only focus on the clearance of α -synuclein, most 156 important of all, promote to explore and discover the root cause which results in α -157 synuclein overproduction and aggregation, thereby eliminating its upstream 158 precipitating factor.

159 Successful therapeutic management might be obtained by targeting pathogenesis 160 of PD at different disease stages. This method should effectively prevent the 161 production of α -synuclein, protection of synaptic function and inhibition of α -162 synuclein hyperphosphorylation at the preclinical stage; removal of aggregated α -163 synuclein, protection of synaptic function and neurons, and attenuation of α -synuclein 164 hyperphosphorylation at premotor phase of PD; rebuilding the functional balance of 165 the neurotransmitter control loop in the brain of PD, that is called, " rebuilding a new 166 home on the ruins" at the early stage of motor symptoms. Comprehensive prevention 167 strategies should be adopted, such as targeting α -synuclein accumulation, synaptic 168 dysfunction, α -synuclein hyperphosphorylation, synaptic function disorder, 169 neuroinflammation and oxidative stress, especially focusing on neuroprotection,

effective exercise rehabilitation and cognitive training, which cloud contribute to
improve movement disorders and cognitive dysfunction of PD, especially gait
disorder and as well as their quality of life at motor symptoms stage.

173 Conlusion

As multiple mechanisms are involved in PD pathogenesis and their relative roles might vary at different stages of this disease. Use of comprehensive prevention strategies and targets at different stages of PD might be a promising way to cure or prevent PD in the future.

- 178 Conflicts of interest
- 179 The authors declare that there is no conflict of interests regarding the publication of180 this paper.
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