REVIEW

Selective vulnerability in neurodegeneration: insights from clinical variants of Alzheimer’s disease

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ABSTRACT

Selective vulnerability in the nervous system refers to the fact that subpopulations of neurons in different brain systems may be more or less prone to abnormal function or death in response to specific types of pathological states or injury. The concept has been used extensively as a potential way of explaining differences in degeneration patterns and the clinical presentation of different neurodegenerative diseases. Yet the increasing complexity of molecular histopathology at the cellular level in neurodegenerative disorders frequently appears at odds with phenotyping based on clinically-directed, macroscopic regional brain involvement. While cross-disease comparisons can provide insights into the differential vulnerability of networks and neuronal populations, we focus here on what is known about selective vulnerability-related factors that might explain the differential phenotypic expressions of the same disease—in this case, typical and atypical forms of Alzheimer’s disease. Whereas considerable progress has been made in this area, much is yet to be elucidated; further studies comparing different phenotypic variants aimed at identifying both vulnerability and resilience factors may provide valuable insights into disease pathogenesis, and suggest novel targets for therapy.

Selective vulnerability

Selective vulnerability in the nervous system refers to the fact that subpopulations of neurons in different brain systems may be more or less prone to abnormal function or cell death in response to specific types of pathological states or injury. The factors underlying this selectivity are well known in some central nervous system diseases, and less well understood or unknown in others. For example, it is well known that certain areas in the brain depend on blood supply from only one or a few arteries due to differences in the circulatory anatomy and are, thus, more vulnerable to an arterial occlusion than other brain areas.6 There are also differences in the energy demand of different neurons: hippocampal CA1 neurons and cerebellar Purkinje cells are characterised by particularly high energy consumption and may, therefore, be more vulnerable to hypoxia and other metabolic stressors.7

In neurodegenerative diseases (NDDs), the potential mechanisms underlying selective vulnerability are complex, multifactorial, and incompletely understood. NDDs are characterised by protein misfolding (in the form of intraneuronal inclusions and/or extracellular protein aggregates) and cell death, with the specific protein(s) implicated and associated patterns of neuronal breakdown translating into a clinical phenotype.8 Yet the patterns of neuronal breakdown within a clinically-driven taxonomy may be defined according to various criteria which span several orders of magnitude: from cortical lobes to white matter tracts to soma to proteins to neurotransmitters to genes. Few of these parameters may be studied collectively in vivo, especially presymptomatically. In fact, these have been largely considered in post mortem tissue, that is, at the end of the pathological cascade. This creates a divide between concepts of selective vulnerability at the cellular and cerebral (system) levels.

In NDDs, the toxicity of extracellular protein aggregates (eg, Aβ in senile plaques in AD) is most likely distinct from the toxicity of intraneuronal protein inclusions (eg, tangles). Extracellular protein aggregates may injure neurons by leaking diffusible oligomers or proto fibrils of the aggregated protein that may interact with receptors or disrupt membranes in a harmful way, and/or by influencing microglial activation that may in some cases cause deleterious inflammation. In contrast, intraneuronal inclusions are more likely to affect normal cellular control of the synthesis, the

INTRODUCTION

While the amyloid plaque and neurofibrillary tangle pathology of Alzheimer’s disease (AD) is often considered synonymous with its most common clinical manifestation, that is, a progressive amnestic syndrome, individual patients have different constellations and degrees of cognitive symptomatology, with some having sufficiently unusual phenotypes that can be considered as having distinct disease variants.1 These include patients with prominent dysexecutive or behavioural problems (sometimes called frontal AD),2 patients presenting with word-finding difficulties and pauses in speech (logopenic variant of primary progressive aphasia, lvPPA),3 patients with various combinations of cortical visual dysfunction, apraxia and dyscalculia with relative sparing of episodic memory (posterior cortical atrophy, PCA),4 and patients with asymmetric rigidity and apraxia, extrapyramidal dysfunction and symptoms related to parietotemporo-occipital cortical involvement (cortical basal syndrome, CBS).4 How can the same broad underlying neuropathology be associated with such markedly different clinical manifestations?
folding, trafficking and degradation of proteins (proteostasis), for example, through sequestration of proteins (eg, chaperones) that would otherwise execute normal functions in the cell. Proteostasis is particularly important to tissues with limited capacity for cell renewal; this vulnerability is likely to be aggravated by the reduced expression of adenosine triphosphate-dependent chaperones with age in the brain. Mutations in the τ gene are sufficient to cause some NDDs, providing unequivocal evidence that misfolded τ in inclusions is toxic. The fact that τ tangles can misfold in a variety of ways and are associated with a range of different NDD, including AD, progressive supranuclear palsy and frontotemporal dementia, also suggests that different neuronal populations are likely to be vulnerable to specific pathological conformations.

The neurons implicated in early AD, including entorhinal cortex and hippocampal CA1 projection neurons, are particularly vulnerable to decreased glucose and oxygen delivery through the vasculature and thus to energy deprivation. Indeed, mild cognitive dysfunction, which frequently progresses to AD dementia, correlates with reduced glucose utilisation in the brain as assessed by fluorodeoxyglucose positron emission tomography (FDG-PET). In addition, synaptic transmission, endoplasmic reticulum stress and calcium homeostasis have been identified as critical factors in the early stages of the disease. This early neuropathological change in AD involves 

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(but not CSF τ) and syndrome-specific variations of atrophy, which may suggest that a more advanced Aβ pathology is associated with AD variant-specific patterns of neuronal injury.

NEURONAL INJURY IN DIFFERENT AD VARIANTS

Neuronal injury in AD variants has been extensively studied in vivo using three different technologies: cross-sectional volume loss or longitudinal atrophy measured by MRI, hypometabolism measured by FDG-PET and τ concentrations in CSF (using total-τ and/or phosphorylated τ). A consistent finding is that the clinical AD variants have different distributions of atrophy and hypometabolism which, in contrast to amyloid burden, much more closely mirror the clinical symptoms.31 34 36 The lack of anatomic specificity is one of the major limitations of fluid biomarkers in the study of the selective vulnerability concept. CSF total-τ and phosphorylated τ may be altered to a similar degree in all AD variants.35

Regarding the distribution of neuronal injury, the typical pattern of volume loss in LOAD is involvement of the medial temporal lobes, hippocampus and parietal lobes; while not always apparent on a single patient level, on a group basis there is volume loss in the posterior cingulate, precuneus and other structures involved in the default mode network. This pattern is also seen in many patients with EOAD, but early onset patients often have a more pronounced posterior cortical hypometabolism,34 and more pronounced deficits in cholinergic and other neurotransmitter systems.37 38 Atrophy in lvPPA is focused to the language-dominant left hemisphere,20 39 while patients with PCA have atrophy focused to the occipital, parietal, and occipitotemporal cortices.4 24 Irrespective of the neuropathology, patients with CBS have asymmetric frontoparietal cortical atrophy, with predominant involvement of premotor cortex, insula, and supplementary motor areas. In addition to these regions, patients with CBS-AD also have involvement of the temporoparietal lobes, with a relative sparing of hippocampus.42 (but this atrophy pattern can also be seen in patients without AD CBS40). Patients with the behavioural or the dysexecutive variant of AD have an atrophy pattern similar to typical amnestic AD, with predominant posterior atrophy rather than frontal atrophy, leading to questions on the designation of ‘frontal variant AD’.4

Despite the differences in neurodegeneration described above, it should be noted that there is a high degree of overlap between AD variants with hypometabolism in the dorsal default mode network31 and atrophy in parietotemporal regions and posterior cingulate.41 Furthermore, with disease progression, the patterns of neurodegeneration converge even further between the variants42 43 making it more difficult to detect variant-specific characteristics in advanced patients.

TAU PATHOLOGY IN DIFFERENT AD VARIANTS

Autopsy studies have found that τ pathology is more closely related than Aβ pathology to atrophy and clinical symptoms.44 The recent development of τ PET imaging now allows this to be tested in living patients. One recent case report of a patient with PCA showed a strong spatial overlap between τ pathology visualised by τ PET imaging using the 18F-AV1451 tracer and hypometabolism visualised by FDG-PET imaging,45 findings which have subsequently been replicated in larger case series.45 Notably, although CSF P-τ concentrations correlate with neurofibrillary pathology in AD,46 47 the marker does not provide information on the anatomic location of the pathology, making it less useful than τ PET imaging in the study of selective vulnerability-related issues.

OTHER PATHOLOGIES

CSF biomarkers to assess plaque and tangle pathology in AD represent an important step forward, particularly when combined with imaging to assess cerebrovascular pathology and (now) cerebral Aβ and τ. However, at autopsy, very few (if any) patients have just one ‘pure’ NDD and in particular, comorbidities including TDP-43 and α-synuclein (Lewy body) pathologies are highly prevalent. While such pathologies are unlikely to be simply coincidental, our ability to address any contributory effects in vivo is very limited,48-50 reflecting that currently available assays only roughly address total levels, showing no or only minor differences between patients and controls31 51 52; key pathogenic species do not appear to have been captured by the assays, at least not in any specific manner; and there are as yet no reliable imaging biomarkers for these inclusions. As and when biomarkers with appropriate sensitivity and specificity become available, it will be possible to test the hypothesis that accumulation of TDP-43 and α-synuclein in neurons might make them more susceptible to τ-induced or Aβ-induced damage, thereby explaining some of the selective vulnerability seen in typical and variant AD.

VARIANT-DEPENDENT INVOLVEMENT OF DIFFERENT FUNCTIONAL NETWORKS

It is notable that the different patterns of neuronal injury observed in AD variants broadly map onto established functional networks in the brain. Thus EOAD, language predominant AD, and visuospatial predominant AD typically show hypometabolism broadly focused in brain areas consistent with executive-control, language and high-order visual networks, respectively.31 34 These findings underlie a model that attempts to integrate findings from Aβ, τ, and injury studies to explain the variability among AD phenotypes,31 postulating that aggregation of Aβ is driven by the total neuronal activity in highly connected cortical hubs (which explains the diffuse and symmetric patterns of amyloid pathology), while τ pathology develops in specific vulnerable networks and, possibly facilitated by amyloid pathology, spreads transneuronally to closely related networks. As τ-mediated injury patterns more closely correlate both with specific functional networks and neuronal loss, this provides a means of explaining the clinical variability.54 If this model is correct, then the different AD variants arise due to different localisations of τ-related neuronal injury in specific functional networks. The next logical step is to identify factors that predispose specific networks to τ-mediated injury. It would also be important to examine whether pathology starts within the same hub (eg, the posterior cingulate) and then spreads differently within differently connected parts of the network, or starts within the network and then moves towards the hub, laterally spreading throughout the other interconnected hubs. One study of different primary progressive aphasias found that learning disability, but not left-handedness, was increased in patients with lvPPA (but not in semantic or non-fluent variants of PPA, which are not caused by AD),35 suggesting that these patients might have had a long-standing dysfunction or differential development in networks relevant for specific aspects of language.

RESILIENCE FACTORS

A corollary of such hypotheses is that individuals should be identifiable who have connectivity patterns inherently resistant to the initiation or spread of neurodegeneration. This may be an important factor underlying incomplete penetrance of genetic
mutations. Differential gene expressions across brain regions may be a further substrate. Akin to cancer, stochastic events at both the genetic or protein homeostatic level may generate ‘seeds’ for the propagation of neurodegeneration. However, most of these seeds may fall on ‘unfertile’ brain regions in terms of connectivity or cellular molecular resistance mechanisms. The varying prevalence of neurodegenerative disorders might reflect a more generic difference in the accessibility of brain networks. In the case of amyotrophic lateral sclerosis (ALS), the primary motor cortex may be a relatively small target to hit that is difficult to access by spread from an occipital lobe focus, but more easily via frontotemporal pathways inherently linked to human motor functions. Once breached, however, the largely mono-synaptic motor system may be easily overwhelmed, reflected in the notably more rapid progression of ALS compared to AD.

CHEMICAL ASPECTS
Could basic chemistry add anything to the selective vulnerability concept? Protein aggregation is concentration-dependent and pH-dependent, with higher concentrations and lower pH increasing the risk of aggregation. Increased production or defective clearance of aggregation-prone proteins intracellularly (eg, via autophagy) or extracellularly (eg, via perivascular drainage) of Aβ may initiate the process. However, there is no direct link between protein aggregation and toxicity. One intriguing and often overlooked aspect in this context is chemical protein ageing (non-enzymatic post-translational modifications that occur over time, particularly in water-deprived milieus, such as inclusions and aggregates and include methylation, deamidation, N-terminal racemisation and truncation), which may increase differential toxicity of the aggregated protein. One hypothesis, supported by data from the prion field as well as by results showing that brain-incubated Aβ is more potent than synthetic Aβ aggregates in regards to its infectivity and toxicity, is that Aβ build-up and toxicity may occur in two phases: the first involving build-up of a non-toxic, virtually inert Aβ reservoir, and the second involving time-dependent chemical modification and destabilisation of plaque-incubated Aβ, which eventually results in induction of microglial activation, tangle pathology and neurodegeneration (ie, gain of toxicity). If so, what factors govern this process are currently unknown, but the concept is supported by recent data showing more N-terminal truncations and pyrogglutamate modifications (both of which increase over time as a consequence of protein ageing) in apparently toxic Aβ aggregates isolated from AD brains, compared with those (probably non-toxic plaques) isolated from cognitively normal individuals who fulfilled criteria for pathological ageing at autopsy (AD-like brain changes without cognitive dysfunction). Similar experiments performed on Aβ pathology in different brain regions primarily affected in different AD presentations, for example, the hippocampus in typical AD and occipital cortex in PCA might therefore provide a means of testing this hypothesis. If gain of toxicity is simply a function of time, the localisation of the first Aβ seeds could determine where the most severe neurodegeneration and symptoms eventually appear; degenerated areas should express more aged Aβ fragment profiles than areas with plaque pathology surrounded by apparently healthy brain tissue. Such a scenario might provide important insights into selective vulnerability within a given network.

CONCLUDING REMARKS
The selective neuronal vulnerability concept tries to explain why only some neurons succumb in the presence of neuropathology, and why certain NDDs devastate certain neuronal networks while leaving others intact. Here, we have discussed this with special emphasis on clinical variants of AD. The aim has been to gather neuroimaging and neurochemical data that may provide useful leads to toxicity mechanisms in the brain in these disorders. It is clear that there is as yet unexplained dissociation between the topographical distribution of key pathologies, that is, Aβ and τ in the case of AD, and that certain neuronal networks may be more vulnerable than others. The precise cascade of pathological events underlying AD is being elucidated only recently; to date there is relatively little data in the literature on the genetic, physiological or biochemical factors influencing neuronal vulnerability or for that matter, resilience factors. While cross-disease comparisons are required to determine the factors influencing why specific proteinopathies affect certain networks, there is much to be learnt from studies of phenotypically different presentations of the same disease. We propose that large, consortia-based studies of deeply phenotyped patients with amnestic and variant forms of AD will be one way to explore selective vulnerability mechanisms in greater detail. There is value in comparing prognostic extremes within the same clinical phenotype, as well as unaffected older populations (the ‘welderly’), especially asymptomatic carriers of typically penetrant mutations. Determining the factors influencing neuronal vulnerability or resilience may in due course provide insights into novel treatment paradigms.

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