Does Sporadic Amyotrophic Lateral Sclerosis Spread via Axonal Connectivities?

Introduction

Amyotrophic lateral sclerosis (ALS) is characterized by rapid progressive paralysis of the striated skeletal musculature [1, 2]. The present review focuses on sporadic disease (sALS), which constitutes the majority of cases without a known genetic mutation. Clinically similar but inherited (familial) forms that display very diverse pathologies [3–7] are not taken into account here.

The pathological process underlying sALS entails abnormal changes of an endogenous and predominantly intranuclear protein, TDP-43 (transactive response DNA-binding protein 43). Following nuclear clearance, the protein remains delocalized in the cytoplasm of susceptible nerve cells, where it undergoes defective phosphorylation and conformational change followed by ubiquitination, which ultimately prevent its re-entry into the nuclear compartment [7–11].

The pathomechanisms that lead to the dysfunction and death of involved cells are still unknown. Along with a “loss of function” attributable to the loss of intranuclear TDP-43 expression accompanied by disrupted RNA metabolism, a “gain of function” mechanism owing to a noxious effect on cells by toxic TDP-43 forms has been discussed [7, 12].

Which Nerve Cell Types Develop TDP-43-immunoreactive Inclusions in sALS?

The abnormal nuclear clearance and development of cytoplasmic TDP-43-immunopositive inclusions are confined to a few types

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ABSTRACT

The pathological process underlying sporadic amyotrophic lateral sclerosis (sALS) that is associated with the formation of cytoplasmic inclusions of a nuclear protein (TDP-43) is confined to only a few types of long-axoned projection neurons. The giant Betz pyramidal cells of the primary motor neocortex as well as large α-motor neurons of the lower brainstem and spinal cord become involved early. In the human brain, these 2 neuronal types are to a large extent interconnected by monosynaptic axonal projections. The cell nuclei of affected neurons gradually forfeit their normal expression of the protein TDP-43.

In α-motor neurons, this nuclear loss is followed by the formation of insoluble TDP-43-immunopositive inclusions in the cytoplasm, whereas in Betz cells the loss of nuclear expression remains for an unknown period of time unaccompanied by somatodendritic and/or axoplasmic aggregations. It is possible that in cortical pyramidal cells (Betz cells) the nuclear clearing initially leads to the formation of an abnormal but still soluble cytoplasmic TDP-43 which may enter the axoplasm and, following transmission via direct synaptic contacts, induces anew TDP-43 dysregulation and aggregation in recipient neurons.

The trajectory of the spreading pattern that consecutively develops during the course of sALS is consistent with the dissemination from chiefly cortical projection neurons via axonal transport through direct synaptic contacts leading to the secondary induction of TDP-43-containing inclusions within recipient nerve cells in involved subcortical regions.
of nerve cells, and these have an axon that is, in comparison to the dimensions of the soma, disproportionately long. Projection neurons with a short axon do not become involved. Motor neurons are especially vulnerable. However, non-motor cells (e.g., projection neurons of the inferior olivary nucleus, the paraventricular portion of the red nucleus, and the striatum) are also susceptible [13, 14]. Thus, the widely used term ‘motor neuron disease’ fails to encompass all aspects of the pathological process underlying sALS. The giant pyramidal cells of Betz in layer Vb of the primary motor neocortex and the large α-motor neurons of the lower brainstem and of the spinal cord anterior horn become involved early and are especially hard hit (Fig. 1) [14].

Which Nerve Cell Types Are Protected against Such Inclusions?
The resistance of many neuronal types to sALS is striking: Even among the motor neurons there are some that seldom or never develop TDP-43-positive lesions: for instance, the visceromotor preganglionic motor neurons of the dorsal glossopharyngeal and vagal nuclei as well as the parasympathetic and sympathetic nuclei in the spinal cord lateral horn [14]. Compared to the morphologically homogeneous group of somatomotor nerve cells in the spinal cord and in motor nuclei of the cranial nerves V, VII, and XI-XII that develop markedly severe pathology, the same cell type in the nuclei of the cranial nerves III, IV, and VI, which control the extrinsic eye muscles, remains virtually intact for the duration of the disease [14].

Corticofugal Projections Control Susceptible Neurons
In humans, the somatomotor neurons of the lower brainstem and spinal cord are directly controlled by corticobulbar and corticospinal projections [15, 16], whereas the nuclei of the extrinsic eye muscles are only indirectly subject to influence by the cerebral cortex and receive their contacts essentially from noncortical fibers of the medial longitudinal fascicle [17, 18]. Visceromotor neurons are subject to, above all, the control of noncortical autonomic centers and to the modulatory influence of the reticular formation and aminergic nuclei of the lower brainstem. As such, it is evident that the existence of direct (i.e., monosynaptic) control by the neocortex is one of the prerequisites for the affection of subcortical nerve cells in the sALS pathological process. In the same context, it should be pointed out that the group of so-called ‘nonthalamic nuclei with diffuse projections’ (i.e., the locus coeruleus, upper raphe nuclei, magnocellular nuclei of the basal forebrain) to the cerebral cortex
and to other regions that become severely involved in other neurodegenerative proteinopathies, such as Alzheimer’s disease and Parkinson’s disease, remain virtually intact in sALS cases. In other words, projection cells with predominantly corticopetal connectivities to the cerebral cortex, in contrast to those controlled by corticofugal projections, are not subject to pathological changes.

The Transmission of the Pathology from Cortical Pyramidal Cells to Projection Neurons of Target Regions

The inclusions that systematically develop during sALS display a characteristic topographical distribution pattern within the central nervous system. During the course of the disease, additional groups of neurons in previously uninvolved regions are drawn into the pathological process in a time-dependent manner. These changes in the distribution pattern of the lesions can be assigned to 4 neuropathological stages (Fig. 1) [14, 19]. Naturally, the duration and transition of each stage cannot be determined within the context of post-mortem cross-sectional studies; nevertheless, in vivo diffusion-tensor imaging (DTI) and diffusion-weighted imaging (DWI)-based results gained from ALS patients fit well with the proposed hypothetical sequence of neuropathological staging [20–24].

Stage 1: Abnormal changes develop not only in Betz pyramidal cells in the deep layer Vb of the primary motor cortex (Brodmann field 4) [25] but also in lower brainstem and spinal cord α-motor neurons (Fig. 1, 2) [3]. These changes commence during the first stage and continue during the subsequent stages.

Stage 2: In addition to those in the agranular motor cortex (chiefly Brodmann fields 4 and 6), lesions appear in adjacent portions of the prefrontal cerebral cortex. In the midbrain, the pathology develops in the parvocellular portion of the red nucleus, which is controlled directly via corticorubral projections [26] and, in the lower brainstem, in parts of the reticular formation and the precerebellar nuclei, such as the inferior olivary nucleus (Fig. 1). By contrast, the magnocellular portions of the red nucleus that project to the anterior horn of the spinal cord remain intact.
Stage 3. The pathology in the prefrontal fields becomes more extensive and reaches both the orbital gyri and gyrus rectus; via the superior longitudinal fascicles it also arrives at the postcentrally located sensory fields of the parietal and occipital lobes. At subcortical sites, the lesions develop in the medium-sized GABAergic projection neurons of the striatum (Fig. 1). Within the 3 subnuclei there, the accumbens nucleus (ventral striatum) becomes involved first followed by the putamen and caudate nucleus [27]. The striatum receives strong corticostriatal projections originating in layer V of the neocortex, the putamen projections arising predominantly in areas of the agranular motor cortex, and the caudate nucleus projections coming from prefrontal fields.

Stage 4. The cortical pathology expands further and enters, via the neocortical regions of the temporal lobe, the transitional zone between the temporal neo- and allocortex (transenthohinal region) and prefrontal cortex (hippocampal formation) (Fig. 1) [14]. The allocortical regions of the temporal lobe that are responsible for learning and the consolidation of memory become involved relatively late [28]. It is remarkable that the hallmark pathological changes develop only in regions that receive and are under the direct control of strong corticofugal projections (Fig. 1), whereas regions that chiefly project to the cerebral cortex, including the locus coeruleus and upper raphe group, are mostly spared [29]. This situation provides support for the idea that the driving force behind the origins and relentless progression of the TDP-43 pathology may reside in the cerebral cortex itself (beginning in the neocortical motor fields) mediated via corticofugal axons and the anterograde transport of soluble TDP-43 that is neurotoxic (Fig. 1, 2) [19, 30–32]. The damage inflicted on selected nerve cells of the cerebral cortex would be, for all intents and purposes, the source that necessarily leads to the subsequent pathological changes in regions of the lower brainstem and spinal cord innervated monosynaptically by these cortical neurons.

The Pathology in Cortical Betz Cells Differs from That in Bulbar and Spinal α-Motor Neurons

The giant pyramidal cells of Betz and the large α-motor neurons in spinal cord and lower brainstem have been well characterized morphologically. They are readily distinguishable from other cell types of the central nervous system and, thus, well-suited for delineating the development of the TDP-43 pathology in both neuronal types (Fig. 2) [25].

Beginning in stage 1, it is possible to see alongside of the many normal Betz cells and α-motor neurons displaying strong TDP-43-positive intranuclear staining an increasing number of the same cell types with reduced intranuclear TDP-43 immunostaining. In time and with disease progression, the nuclei of such cells completely lack TDP-43 immunoreactivity. However, many of the involved Betz cells also fail to display any TDP-43-positive cytoplasmic inclusions (Fig. 2) [32]. Only closer scrutiny of a larger number of these cells reveals the presence of a few that contain very subtle cytoplasmic aggregates. Thus, it can be concluded that cortical Betz cells do not completely forfeit their capacity to produce insoluble cytoplasmic inclusions; on the contrary, this capacity is only suppressed for an indeterminate interval of time or the process of TDP-43 aggregation occurs at a remarkably slow rate (Fig. 2).

The nuclei of the large bulbar and spinal cord α-motor neurons also display the phenomena of reduced TDP-43-positive intranuclear staining and, ultimately, complete TDP-43 intranuclear clearance. In contrast to the Betz cell population, however, the α-motor neurons with weak TDP-43-immunoreactive nuclei contain needle-like cytoplasmic aggregates that extend into the somatodendritic and axonal compartments of involved cells [33]. Then, the aggregates appear to coalesce into coarser skein-like inclusions (Fig. 2) [32].

Can the Pathology Be Propagated Transsynaptically from the Cortical Projection Neurons to Interconnected Projection Neurons?

In the ascending primate scale, the direct control of corticofugal tracts via monosynaptic contacts increasingly replaces the indirect influence exerted on the α-motor neurons by the cerebral cortex via intervening interneurons in lower mammals. In the human nervous system, this development reaches a provisional acme [15, 16, 29, 32, 34]. The assumption that cortical neurons (Betz cells) are the first to become involved followed by the α-motor neurons, tends to favor Charcot’s theory [35], namely, that ALS originates in the lateral funiculus (i.e., corticospinal tract, where the axons of the Betz cells are located) and only then affects the α-motor neurons in the anterior horn of the spinal cord [36]. The sources of the very first malfunctions within the protein TDP-43 in Betz cells are incompletely understood; but, it is conceivable that the further spread of the pathology to interconnected neurons takes place by means of the phylogenetically ‘new’ or recent monosynaptic contacts [9, 29, 32, 37–41]. Whether the same holds true not only for corticobulbar and corticospinal projections but also for corticorubral and corticostriatal connectivities is currently an open question.

Conclusions

We have postulated for sALS that the misfolded and abnormally phosphorylated protein TDP-43 may be present in the cytoplasm in a soluble phase in involved Betz cells displaying nuclear clearance (TDP-43 immunonegative cell nuclei) and a TDP-43-immunonegative cytoplasm. Anterograde axonal transport of the soluble protein could permit it to reach the presynaptic terminals of the Betz cells, where its transsynaptic (cell-to-cell) transmission to the corresponding α-motor neurons of the spinal cord and brainstem could lead anew therein to the pathological dysregulation and prion-like seeded propagation of TDP-43 [42]. According to this hypothesis, the emergence of cytoplasmic aggregates within target neurons would mostly interfere with the ensuing spread of the pathology to the immediately following neuron in the neuronal chain (Fig. 1).

The regional distribution pattern of the TDP-43 pathology described above is not compatible with theories of a pathological process that purportedly commences in motor neurons of the spinal cord.
cord or lower brainstem, followed, secondarily, by damage to cortical areas [43–47]: Such a proposed route of progression fails to explain the involvement in SALS of the parcellocular portion of the red nucleus and of the medium-sized GABAergic projection neurons in the striatum, none of which have any connectivities whatsoever to spinal or bulbular motor neurons (*Fig. 1*).

The concept that the disease process can be primarily influenced by the cerebral cortex is corroborated by the clinical findings reported for split-hand syndrome and split-leg phenomenon [48–54]. Clinical SALS phenotypes that eventually allow for alternative interpretations, e.g., flail-arm or flail-leg syndrome [45–47, 55], still require neuropathological confirmation [56, 57].

From this standpoint, it is essential to elucidate mechanisms that might preserve TDP-43 for an indeterminate period of time in an abnormal but soluble phase within the cytoplasm of cortical projection cells and to clarify the mechanisms by means of which neuron-to-neuron transmission occurs. Selective intervention into these mechanisms could eventually prevent the TDP-43 pathology from spreading into interconnected motor neurons.

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**Conflict of interest**

No conflict of interest has been declared by the authors.

**References**


