2015

Cross-Sectional Survey of Relevant Literatures as to the Current Proposed Disease Mechanisms and Treatments of Amyotrophic Lateral Sclerosis (ALS)

Zachary Sanford

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Recommended Citation
DOI: http://dx.doi.org/10.18590/mjm.2015.vol1.iss1.3
Available at: https://mds.marshall.edu/mjm/vol1/iss1/3

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Author Footnote: I would like to thank Drs. Janet Clark and Ole Mortensen of the Drexel University College of Medicine department of Pharmacology and Physiology for their time and tutelage as well as presenting neuropharmacology in an exciting and engaging format.
References with DOI


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Cross-Sectional Survey of Relevant Literatures as to the Current Proposed Disease Mechanisms and Treatments of Amyotrophic Lateral Sclerosis (ALS)
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The author has no conflict of interest to disclose.

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Abstract

Amyotrophic Lateral Sclerosis (ALS), more commonly referred to as Lou Gehrig’s disease, is a progressively degenerative neuromuscular disorder affecting both the upper and lower motor neurons and preferentially affecting males in their forties to seventies. Although the pathology of ALS has been clearly elucidated elsewhere, the precise mechanisms by which the disease progresses and the means by which it is acquired are still poorly understood areas of medicine. Current genomic and proteomic studies in human and animal models have yielded exciting and promising new findings that may help elucidate this pathology. It is the purpose of this review article to discuss the most relevant proposed mechanisms in current medical literature available from NCBI’s PubMed database as well as to highlight past, present, and future pharmacologic intervention therapies which have experienced varying degrees of success. This is by no means an exhaustive assessment of the current literature available; however it should suffice as a thorough review of the most salient points of modern ALS research.

Keywords:  ALS, Amyotrophic lateral sclerosis, mechanism, pathology, pharmacology

Introduction

Amyotrophic Lateral Sclerosis (ALS), more commonly referred to as Lou Gehrig’s disease, is a progressively degenerative neuromuscular disorder affecting both the upper and lower motor neurons and preferentially affecting males in their forties to seventies. Disease pathogenesis selectively destroys upper motor neurons (UMN) of the primary motor cortex and corticospinal tracts in addition to lower motor neurons (LMN) of the spinal anterior horns and pontomedullary nuclei. The destruction of these motor neurons results in a loss of coordination for voluntary movement, often manifesting in a combination of UMN symptoms including muscle spasticity, hyperreflexia, increased deep tendon reflexes, and appearance of immature reflexes such as Babinski sign in conjunction with LMN symptoms such as progressive weakness, muscle hypotonia and atrophy, fasciculations, dysarthria, dysphagia and eventually respiratory failure. Of these symptoms, respiratory failure provides the greatest danger to the patient, with complications including secondary bacterial pneumonia compromising patients within two or three years of diagnosis. While the initial manifestation of these symptoms is seemingly random in distribution, disease progression often follows along contiguous neuroanatomic tracts to varying degrees of severity. The resulting UMN and LMN symptoms spread at differing rates and manifest as a consequence of complex somatotopy, creating complex motor phenotypes that are a hallmark of the disease.

The disease can further be subdivided into separate categories based on the method of inheritance or the location of symptom onset. ALS can present either sporadically (sporadic ALS, SALS) or along familial lines (Familial ALS, FALS) with a greater incidence of sporadic cases in the United States. Furthermore, symptoms of neurodegeneration can be first noticed either in the extremities (so-called extremity onset ALS) or the tongue (bulbar onset ALS). Extremity onset ALS is a degeneration of motor neurons enervating the arms or legs of afflicted individuals and can first include localized fasciculations, stiffness, and weakness. Patients with affected upper limbs will notice difficulty completing tasks that require manual dexterity.
whereas those with lower limb onset will notice irregularities in gate and a distinct dropped foot appearance.\(^1\) Disease progression is often less rapid in those presenting with extremity ALS than in bulbar cases. Those afflicted with bulbar onset ALS first notice difficulty speaking or swallowing, and symptoms quickly spread to the intercostal muscles and the diaphragm,\(^1\) posing serious respiratory risks that are the most frequent cause of death in ALS patients.\(^2\)

**Lack of Consensus**

Presently there is a surprising lack of consensus among the medical community as to how to categorize familial versus sporadic ALS.\(^5\) In an international study among leaders in ALS research spanning Europe, India, Australia, and North America there were inconsistencies in how the distinction between the two categories were defined. While some felt the criterion for FALS should include first-degree relatives including parents or siblings, others felt the need for second-degree (such as first or second cousins) or even any family relationship to warrant the title FALS.\(^5\) This distinction goes beyond nomenclature, as the protocols for diagnosing the two subfamilies differs substantially. When asked for comment, participating neurologists claimed to conduct screens for likely candidate genes implicated in ALS more frequently in FALS rather than SALS cases an overwhelming 67.0% to 10.3%, respectively.\(^5\)

Further complicating the situation is the lack of clear and agreed upon mechanisms of disease progression among the medical community. To date there have been over twelve loci and eight genes\(^6\) proposed as causative agents for FALS and among them exist hundreds of variations. Not surprisingly, the data available on such a diverse field of study is vast, with a literary search conducted on NCBI’s PubMed database yielding close to 19,000 unique entries for amyotrophic lateral sclerosis. Searches were subsequently narrowed to include the terms “pathology” and “pharmacology,” resulting in 1,034 results. Of these, entries were broadly selected to represent unique aspects of the body of literature as a whole. It is the purpose of this review article to therefore discuss the most relevant proposed mechanisms in current medical literature as well as to highlight pharmacologic interventional therapies with demonstrated or potential future success in treating ALS.

**Suggestive Trends in Diagnosis**

Among the difficulties in treating patients with ALS is quickly and effectively confirming a diagnosis while ruling out other possible disease processes. Effective clinical indicators are contested on the grounds of validity due to an incomplete understanding of the disease mechanism however firm agreement can be reached with regards to symptomology. For this reason, studies assessing the validity of measuring tongue strength as a prognosticator of bulbar onset ALS have yielded strong statistical significance.\(^7\) The effective use of this technique can help serve as a valuable tool in early, noninvasive detection of bulbar onset ALS rather than waiting for more advanced symptoms such as dysarthria and dysphagia which, once evidenced, are already severe enough to strongly affect patient survivability. Of the patients whose tongue strengths were assessed and who later developed ALS, a statistically significant proportion deteriorated quickly regardless of site of onset, indicating a rapid spread of lower motor neuron degeneration is best marked by assessment of tongue strength.\(^7,8\)
On the other side of the clinical spectrum are highly advanced technologic measurements using functional Magnetic Resonance Imagery (fMRI) to assess neuronal activation as a function of disease progression. Interestingly, analysis of these techniques suggests there is an observable signal change in cortical blood oxygen levels during tasks requiring motor control of the hand specifically in ALS patients as measured against controls. These changes reflect a neuroplasticity heretofore unknown in ALS patients, although limited to those with slowly progressing disease symptomology and present only in initial phases of the disease. Still, these findings suggest either that slow disease progression affords patients the time to reorganize their own somatotopy or, conversely, that because patients are able to demonstrate efficient neuroplasticity they are able to delay the progression of the disease. Whichever the conclusion, the proven use of fMRI to assess these neuronal rearrangements proves promising for future study.

Not all advancements in the understanding of ALS have been noninvasive; attempts at isolating novel markers from cerebral spinal fluid (CSF) in order to obtain objective clinical markers for disease progression remain one of the forefronts of ALS research. Degeneration of the blood-CSF barrier is observed in 46% of ALS patients and offers a suggestive area of clinical assessment. Still, markers of blood brain barrier impairment, blood-spinal cord barrier impairment, blood-CSF barrier impairment, neuroaxonal degeneration, oxidative stress, neurotransmission, inflammation, immune activation, and glial activation have not yet yielded clinically relevant information in the diagnosis of ALS. While many of these markers are present in cases of ALS, they are not exclusive to this disease and have as yet presented a unique and reproducible clinical profile. However advancements have been made in other neurodegenerative diseases, with reductions in CSF cAMP and cGMP concentrations in Creutzfeldt-Jakob disease but not in ALS.

**Disease Modeling**

Clinical presentations afford unique opportunities to track the progression of disease while occasionally affording the chance to test FDA-approved experimental drugs. However in the interest of discovery, safer, non-human models must be selected for study in order to fully elucidate disease mechanisms and assess pharmacologic intervention therapies. The gold standard in ALS research remains the SOD1-G93A transgenic mouse model for preclinical drug studies. G93A mice possess gain-of-function mutations in the Superoxide Dismutase (SOD1) gene that enhances the generation of damaging oxygen radicals by Cu,Zn Superoxide Dismutase (Cu,Zn SOD). These mutations accurately reflect symptoms present in human FALS patients while also affording the luxury of working with a small, relatively short-lived animal model.

More recently, Pembroke Welsh Corgis and Boxers with mutations in SOD1 have been proven to illustrate degenerative myelopathy and peripheral neuropathy that can serve as a potential spontaneous model for human ALS. Homozygous silent mutations in canine SOD1:c.118 are able to recapitulate symptoms seen in the human disease process and can therefore offer superior alternatives to the murine model with regards to surgical study and evaluation of pharmacologic intervention. SOD mutants of the species *Caenorhabditis elegans* have also been generated for invertebrate study. *C. elegans* with pan-neuronal G85R SOD1 expression demonstrate significantly impaired locomotion and SOD1 aggregates characteristic of human
FALS patients with SOD1 mutations. These invertebrate models offer cheaper alternatives to the higher order research animals and are free of many of the ethical concerns present when working with mammals. They are limited in their application, though, as the Federal Food and Drug Administration requires drug testing to be performed on mammals before reaching the public.

**Proposed Mechanisms for ALS Pathogenesis**

Many of the proposed mechanisms currently under investigation focus on aberrations in neuronal transcription and translation events (Table 1). Irregularities in gene sequences either directly inherited or through predisposition to environmental insult, shift cellular machinery away from normal cellular chemistry toward dysfunctional proteins, hallmark cytosolic aggregations and plaques containing translationally inert RNA, and dysregulations in repair mechanisms. As no definitive etiology for ALS has been established, any or all of these mechanisms potentially represent processes in the pathology of disease and may contribute to a greater interwoven narrative. The following mechanisms are listed from most well-understood to the least elucidated although there is significant overlap in their purported functionality.
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Transactive Response DNA-Binding Protein of 43 kDa (TDP-43)

TDP-43 is a nuclear nucleic acid binding protein primarily involved in RNA translational events and metabolism that when irregularly aggregated in the cytosol represents the major pathological finding in ALS cell studies.\textsuperscript{16-18} Pathologic accumulations demonstrate polyubiquination, hyperphosphorylation, and irregular cleavage that likely contributes to the abnormal localization in affected neurons. Once in the cytosol, mutant low molecular weight TDP-43 forms β-pleated sheets not consistent with wild type phenotype\textsuperscript{19} that are highly dependent on C-terminal domains demonstrating unusual stability and resistance to proteasome degradation.\textsuperscript{20} Granule accumulation is further promoted by microtubular dysfunction.\textsuperscript{21} Silent mutations in the 342–366 region affect RNA splicing regulation promoting the formation of stress granules whose exact function is as yet poorly understood. Current research as to whether these aggregates are protective or the hallmark of cellular damage are currently under investigation, as are studies to determine whether a loss of function or a gain of function mutation is responsible for this phenotype. What is known is that a polyubiquinated TDP-43 aggregate is present in 5% of FALS and 0.5-2% of SALS cases.\textsuperscript{22}

Regional differences in composition of TDP-43 stress granules between neuronal and glial intracytoplasmic inclusions in the spinal cord and medulla oblongata of SALS patients and those found in the basal ganglia and hippocampus indicate the possibility of heterogeneity in function or pathology depending on location in the central nervous system. Among these heterogeneities is the presence of Smurf2, an upstream effector of E3 ubiquitin ligase that may impact lower motor neuron function\textsuperscript{23} and studies suggesting TDP-43 operates independently of adenosine deaminase on RNA 2 (ADAR2).\textsuperscript{24}

It has been implicated that other proteinaceous aggregates present during times of cellular strain serve as scaffolding for the formation of the cytosolic TDP-43 aggregates.\textsuperscript{25} These aggregates may follow a prion-disease-like pathology of replication, acting as an infectious transmissible protein species that behaves as a template for misfolding of nascent wild type proteins.\textsuperscript{19,26,27} This theory is further supported by the presence of described prion-related glutamine/asparagine (Q/N) rich domain preferentially seen in prion disease and also expressed in proteins in ALS patients with mutant TDP-43. Introduction of these proteins in human embryonic kidney 293 (HEK293) cells was able to demonstrate prion seed-like behavior, serving as a template from which nascent wild type proteins misfolded to mutant conformations.\textsuperscript{19,28} Further studies in murine models with inducible cytosolic TDP-43 aggregates showed statistically significant correlation between increasing number of aggregates and ALS symptomatology, with a reversal of these symptoms seen on restoration of wild type TDP-43 function.\textsuperscript{29}

Work in \textit{C. elegans} models has suggested TDP-43 aggregates serve as a nidus for calcium-dependent neurodegeneration based on activation of killer aspartyl proteasomes. These models illustrate that reductions in intracellular calcium release, specifically via loss of function mutations in calreticulin, calnexin, or the ryanodine receptor or the introduction of intracellular calcium via ethylene glycol tetraacetic acid or dantroline are neuroprotective while increases in intracellular calcium via thapsigargin further exacerbate neurotoxicity.\textsuperscript{30} Other mouse studies
mimicking nascent TDP-43 function in deficient cell lines was able to produce similar findings, with poly(A)-binding protein nuclear 1 protein specifically demonstrating promising results.\textsuperscript{18,31}

TDP-43 has been tentatively linked to a mechanism involving C9orf72 hexanucleotide expansion and Fus due to their involvement in RNA binding and transport\textsuperscript{32} although these mechanisms are as yet poorly understood. Data co-localizing TDP-43 with FUS aggregates and not with SOD1 has also been presented.\textsuperscript{21} Continued study into the pathogenesis of these aggregates is ongoing\textsuperscript{33,34} but is made more difficult with inconsistencies in the expected phenotypes of genotyped individuals. In a study by Mosca et al. two blood relatives with identical genotypes for the TDP-43 locus evidenced markedly different phenotypes; one sibling had severe onset ALS whereas the other (and the two other heterozygotes in the family) remained unaffected.\textsuperscript{35} Additionally, TDP-43 aggregates may not inherently be pathologic, having been located in adrenal medulla samples of individuals with and without ALS.\textsuperscript{36} Studies have begun elucidating the possibility that there are potential variations in TDP-43 conformations that can act as strains in pathogenesis with varying impact on cellular toxicity and disease pathogenicity.\textsuperscript{19}

**Cu/Zn Superoxide Dismutase 1 (SOD1)**

Mutant SOD1 proteins have been implicated in extracellular neuronal damage in a number of central nervous system diseases and are the traditional model for ALS disease progression. Intracellular damage is seen in clinical preparations however there is an equally likely mechanism depicting extracellular neuronal damage in mutant of SOD1 totaling over 140 variants. Studies have implicated that one of these variants, apoSOD1, undergoes abnormal folding to expose its zinc binding site,\textsuperscript{37} depleting cellular zinc stores and disrupting zinc homeostasis. In vitro studies have proven that apoSOD1 expression reduces viability of target cells but that this is not caused by the accumulation of cell aggregates as was seen in the RNA binding proteins listed previously. Further illustrating the zinc hypothesis are results showing abatement of cell destruction by the addition of exogenous zinc and exacerbation by the addition of known zinc chelators.\textsuperscript{38} These misfolded SOD1 mutants are increasingly becoming the target for drug therapy\textsuperscript{39} due primarily to their speculated early involvement in a thorough and complex oxidative damage cascade. Monoclonal antibody experiments targeting these abnormally folded SOD1 mutants\textsuperscript{40} have already been demonstrated in murine and human tissues.

The oxidative stress mechanism of nitrous oxide and other radical species put forth by proponents of the SOD1 theory of ALS pathogenesis are well documented in the literature\textsuperscript{41} and accommodates a wide variety of SOD1 mutants with an equally wide array of cellular effects.\textsuperscript{42} Among these varieties is a hyperoxidized variant of SOD1 found to be linked directly to certain cases of bulbar SALS\textsuperscript{43} as well as variants that resulted in decreased O-glycosylation with β-N-Nacetylgulosamine (O-GlcNAc), an agent linked to competitive inhibition of cellular phosphorylation.\textsuperscript{44} The implication of this reduction in intracellular O-GlcNAc is a corresponding alteration in kinase activity which would cause highly irregular and cell-wide effects. Many of these mutants have phenotypic effects directly influenced by copy number of their aberrant proteins\textsuperscript{45} which are best assessed by RT-qPCR.
Similar to TDP-43 and its interrelatedness with Fus and C9orf72 hexanucleotide expansion, SOD1 mutations have been implicated in pathways with abnormal RNA binding proteins. Methodologies proposed by Pokrishevsky et al. purport TDP-43 or Fus aggregates in the cytosol facilitate wild type SOD1 conformational changes, forming new and unusual conformations. This mechanism mirrors the already established pattern of prion diseases, where a misfolded protein serves as a template for further misfoldings. This theory is generally accepted among the ALS community as a likely mechanism to explain SALS with SOD1 abnormalities. SOD has also been implicated as an intermediate in the ER damage cascade, with mutant SOD1 inappropriately utilizing PDI. To even further complicate the picture of a complete SOD1 cascade, fibroblast growth factor 2 deficiency prolonged survival and improved motor performance in ALS mouse models positive for mutant SOD1 protein. Accompanying this deficiency was an associated up-regulation of glial derived neurotrophic factor and ciliary neurotrophic factor, two motor system-related neurotrophic factors.

**Fused in Sarcoma (FUS)**

FUS is a nuclear RNA binding protein involved in transcription, post-translational modifications including RNA splicing and microRNA processing, and DNA repair. Aberrations in FUS also represent one of the primary hallmarks of ALS cellular pathology, identified in 4-6% of FALS and 0.7-1.8% SALS cases. These mutations result in a hyperosmolar cytosolic environment where localization of mutant aggregates can occur. These aggregates form stress granules similar to and occasionally co-expressed with TDP-43 and SOD1, where local RNA molecules play an as yet undetermined role in granule stability. Furthermore, FUS irregularities disrupt correct localization of RNA binding proteins, disrupting nuclear cellular events. One current theory posits the disruption in FUS localization prevents correct DNA damage repair, resulting in accumulation of injurious insults to motor neurons and culminating in stereotyped ALS symptomology. FUS mutants also express deficiencies in autoregulation, leading to an increased vulnerability to cellular stress and degeneration manifesting as axon withdrawal, synapse dysfunction, and muscular denervation.

Animal models in *Drosophila* have demonstrated that mutations in FUS analog Cabeza (Caz) causes in vivo toxicity by disrupting neuromuscular junctions and inducing apoptosis in motor neurons. These effects are caused by a gain in function mutation resulting in overexpression of the gene, although these effects rely on the expression of the protein in the nucleus. Mutations in the *Drosophila* homolog Caz caused retinal degeneration, wing defects, decreased viability, but most importantly locomotive impairment in motor neurons and a disruption of presynaptic terminals at the neuromuscular junction culminating in motor neuron apoptosis. Toxicity is directly correlated to a functional C-terminus with overexpression as well as deletion resulting in similar disruptions of locomotion and neuromuscular junction abnormalities seen in human ALS patients. Apoptosis of motor neuron cell bodies was only observed in FUS overexpression models. C-terminal FUS knockout mice have corroborated the evidence for cytosolic aggregation seen in *Drosophila* but were unable to recreate ALS symptomatology. Additional murine models have been able to demonstrate neuroinflammation, denervation of neuromuscular junctions, reductions in functional dendritic cell architecture and total cell number, and premature host death when gain or loss of function mutations were introduced to FUS.
Differences are observed in clinical models versus those seen in Drosophila, as variants in human FUS have been proven to signify endoplasmic reticular (ER) stress. Co-localization with ER markers was seen in human lumbar spinal cord sections taken from ALS patients. These markers include protein disulfide-isomerase (PDI) as well as calreticulin, whose aggregates were seen with FUS in motor neuron cytosols, having translocated from the nucleus. These findings were consistent with both SALS as well as FALS and are related to the pathophysiology of SOD1 mutations. Patient-derived induced pluripotent stem cells transformed into spinal cord neural cells represent one model of FUS study, where stress granule aggregation has been correlated to volume of cytosolic mutant FUS concentrations. Utilizing similar models, DNA damage has been linked to cytoplasmic accumulations of mutant FUS mediated by phosphorylation of its N terminus by the DNA-dependent protein kinase.

In contrast to the models discussing aberrant cytosolic inclusion as a means for pathology, forced nuclear aggregation of FUS has been able to demonstrate neuronal cell death in some in vitro models. One study has isolated nuclear mutant FUS aggregates from ALS fibroblasts that proved resistant to sodium dodecyl sulfate denaturation, supporting the possibility that there are multiple ways to alter FUS functionality to cause neurodegeneration. Similar to TDP-43, FUS has also been implicated in prion disease-like pathology with FUS stress granules serving as proteinaceous templates for misfolding of nascent FUS proteins. This area of study is not yet fully explored and represents an ongoing area of research.

**Chromosome 9 Open Reading Frame 72 (C9orf72) Hexanucleotide Repeat Expansion**

Repeats of the hexanucleotide GGGGCC within the non-coding portion of C9orf72 on chromosome 9p21 have been reported to cause chromosome 9-linked FALS and frontotemporal dementia (FTD) and represent the most common genetic variant of ALS. In an international cross-sectional study evaluating 4448 ALS patients, 37% of individuals with FALS are in possession of this mutation, having as many as 700-1600 copies of the repeat sequence compared to controls who only have 23 copies. Comparable studies have been conducted in French (46% FALS, 8% SALS), Italian (46% FALS), Greek (50% FALS, 8.2% SALS), Turkish (18.3% FALS, 3.1% SALS), Slavic (5.9% SALS), and Russian (15% FALS, 2.5% SALS) populations testing positive for the GGGGCC repeat. Studies on Chinese SALS populations have yielded results indicating this sequence is not correlated to their patient populations, potentially describing a predilection for the C9orf72 hexanucleotide repeat for Caucasians.

Phylogenetic analyses suggest that the hexanucleotide first originated in Finland, with all present day GGGGCC sequences in C9orf72 owing to a single mutation event 1500 years ago and representing 46.4% of FALS and 21.1% SALS of the Finnish population. The hexanucleotide repeat leads to the downregulation of the expression of an alternatively spliced C9orf72 transcript and to the formation of aberrant nuclear RNA foci. The increasing evidence for the presence of this sequence in FALS cases combined with the late age of onset for symptoms has led some studies to evaluate the possibility that this irregularity in RNA processing caused by the hexanucleotide expansion may be initially corrected by conventional cellular repair mechanisms only to eventually overwhelm the system in later life. This expansion itself may not be inherently irregular but instead trigger pathogenicity in differentially
methylated states\textsuperscript{80,81} and has also demonstrated a concordance with abnormal localization of TDP-43, a hallmark of ALS neuronal pathology.\textsuperscript{58} Murine models with induced GGGGCC sequences have also been constructed, evidencing a gain of function relationship theorized elsewhere.\textsuperscript{82,83}

Although this sequence provides a possible marker for over one third of FALS cases, the ethical and moral implications of how to determine when patients should or should not be screened for this repeat are difficult at best to determine. The concerns about incomplete penetrance raised by Majounie et al. highlight an average age of disease onset of 57 with patients only beginning to have symptom onset well into their nineties.\textsuperscript{69} The exact determinants, if there are any, as to whether or not one should be screened for this repeat are as yet undetermined.\textsuperscript{84} Still, the screen can offer confirmation of disease in the event of a suspected diagnosis. Ongoing research into reproducible biomarkers of this region and novel therapeutic targets are a developing area of study.\textsuperscript{85}

**P56S-vesicle-associated membrane protein-associated protein B (VAPB)**

VAPB is a type II integral membrane protein of the ER with the free N terminal region projecting into the cytosol and is involved in ER to Golgi transport as well as cellular stress response pathways.\textsuperscript{86-90} Murine models have illustrated loss of function mutations altering the unfolded protein response pathway, reducing functional myotubule formation\textsuperscript{91} and this model has been illustrated in human samples elsewhere.\textsuperscript{86,92} Additional murine studies have demonstrated progressive hyperactivity and motor impairment due to corticospinal and spinal motor neuron destruction in VAPB transgenic mice.\textsuperscript{93} Autosomal dominant ALS based on a proline to serine substitution in the VAPB has been linked to nuclear envelope deformity, interrupting the transport of nucleoporin and emerin to the nuclear envelope.\textsuperscript{94} VAPB has also been shown to suppress adipocyte lipid differentiation through alterations in mRNA expression, potentially altering energy metabolism in a subset of ALS patients.\textsuperscript{86} However, despite its involvement in intracellular transport, VAPB mutations have not demonstrated alterations in protein transport or degradation.\textsuperscript{95} Of note, this mutation has not been demonstrated in SALS patients in Sweden, Portugal and Iceland.\textsuperscript{96}

**Optineurin (OPTN)**

OPTN is a protein involved in the nuclear factor kappa B (NF-\kappa B) signaling pathway. Several mutations have demonstrated inhibited NF-\kappa B activation and irregular localization of NF-\kappa B proteins to neuronal and glial cytoplasm\textsuperscript{97,98} with a 300 bp region consisting of Alu repeat sequences that are highly susceptible to novel deletions. Normal OPTN function has been shown to induce mitochondrial autophagolysosome formation and serves as an autophagy receptor, with mutations disrupting its ability to bind polyubiquinated proteins for transport to the lysosome for destruction.\textsuperscript{99-102} TDP-43 and SOD1 positive cells from SALS patients have also tested positive for OPTN mutations and suggest loss of function in OPTN as a contribution to TDP-43 accumulation.\textsuperscript{97,103-105} OPTN mutants are especially prevalent in the Japanese population, constituting the second most mutated gene product associated with SALS in Japan.\textsuperscript{106} In contrast, this mutation is exceptionally rare in British populations where it is deemed unlikely to be pathologic\textsuperscript{107} but can serve as a potential cause of rare rapidly progressive SALS in the Dutch.\textsuperscript{108}
Ewing Sarcoma Breakpoint Region 1 (EWSR1)

EWSR1 is a RNA binding protein related to FUS and TDP-43 and a member of the FET protein family. Due to the fact that this protein is in possession of a predicted prion domain located on the C-terminus, it is prone to aggregate formation in both in vitro and in vivo specimens. It has been demonstrated to have a toxic effect when expressed and was proposed as an additional candidate for ALS pathogenesis. Histologic preparations of ALS spinal cord specimens illustrated aberrant localization in the cytosol of EWSR1 as opposed to the nucleus in healthy controls. One study presented 817 patients with ALS and 1082 healthy control individuals who tested positive for mutations in EWSR1 exons 15-18, the most pathogenic mutations in the gene region. Further work is being conducted evaluating the relevance of this protein.

Spinocerebellar Ataxia Type 2 Protein (ATAXIN2)

ATAXIN2 is another RNA binding protein potentially linked to ALS pathology, with moderate expansion of CAG polyglutamine sequences correlating to increased risk of ALS. Based on a cohort of 1538 ALS patients and controls in one Italian study, polyglutamine sequences that in excess of 31 amino acids place an individual at elevated risks for SALS compared to those lacking the repeat sequence. These trends did not hold true for FALS cases. This was also demonstrated in a cohort study of 375 ALS patients of Sardinian ancestry with similar findings. Aberrant ATAXIN2 accumulations associate with TDP-42 and FUS, altering their respective cellular toxicities and serving as components of cytosolic ALS stress granule formation.

Epigenetics and Environment

Recently, studies into the impact of environment on gene expression have yielded insight into the dynamic nature of transcriptional potential. Early life exposure, for example to in utero hypoxia, heavy metals, pesticides, tobacco, alcohol, or vitamin deficiency has been implicated as potential cause for changes in the genetic transcriptome. While the precise mechanisms by which such stressors can contribute specifically to ALS are as yet poorly understood, some promising studies have suggested specific methylation events on proteins such as Kir4.1 glial cell potassium channel by DNA methyltransferase may play a role in pathogenesis. Additionally, individuals with specific single nucleotide polymorphisms in the MT gene encoding metallothionein have demonstrated environmental susceptibility to acquiring ALS.

Treatments and Interventions

To date many studies have been conducted to determine which treatment interventions pose the greatest benefit to ALS morbidity and mortality. The following presents evidence for those methods that were strongly believed to be clinically significant, beginning with those that have demonstrable improvement on symptomatology and survival and continuing with interventions that may prove promising in the future, then concluding with several notable therapies that have not proven beneficial and should be avoided in the treatment of ALS (Table 2). Due to the
chronic nature of this disease, each intervention is part of a broader scope of treatment delivered by neurologists, respiratory therapists, speech-language pathologists, physical therapists, dieticians, social workers, psychiatrists, nurses, and other allied health professionals.

<table>
<thead>
<tr>
<th>Table 2: NOTABLE INTERVENTIONS TESTED IN MODIFYING ALS PATHOLOGY AND PATIENT QUALITY OF LIFE</th>
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<td>Intervention</td>
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**Bi-Level Intermittent Positive Pressure (BiPAP)**

Due to the current limited understanding of the full mechanism of disease progression in ALS, the most important treatment to date remains the effective administration of assisted respiratory technologies. Of these, the most useful and well-documented in the literature is the use of bi-level intermittent positive pressure (BiPAP) as an effective, noninvasive, and generally well-
tolerated means of ensuring gas exchange. In a retrospective study conducted at Hahnemann Hospital, Kleopa et al. evaluated the efficacy of BiPAP intervention on patient outcomes and noted significant success in prolonging lifespan and decreasing symptom onset in ALS patient populations. Furthermore, the clinic suggests that patients receive a recommendation for BiPAP during the first clinical visit where the Forced Vital Capacity (FVC) reaches 50% of the expected normal. When used for four hours a day or longer, BiPAP helped increase survival by 14.2 months.122

Riluzole

Currently there is only one FDA-approved drug for the treatment of symptoms in ALS patients. This drug, Riluzole, is a neuroprotective drug that blocks glutamatergic neurotransmission in the CNS. It has been proven to inhibit the release of glutamic acid in vitro and in vivo, though the mechanism is still speculated and assumed to be due to inactivation of voltage-dependent sodium channels on glutamatergic nerve terminals or possibly the result of a G-protein coupled receptor cascade. Riluzole also blocks some of the postsynaptic effects of glutamic acid by noncompetitive blockade of N-methyl-D-aspartate (NMDA) receptors. The drug is generally accepted to be a neuroprotectant with strong anticonvulsant albeit sedative properties. In clinical trials, 100 mg daily dose of Riluzole was able to demonstrate a 9% gain in probability of surviving one year over control and raised median survival from 11.8 to 14.8 months. There was also modest symptom improvement in both bulbar and extremity onset ALS.124

Dextromethorphan with Ultra Low-Dose Quinidine (Nuedexta®)

Psuedobulbar affect is a prominent complaint among patients with ALS, prompting studies to be completed on the potential benefits of dextromethorphan. As a selective, noncompetitive antagonist of the N-methyl-D-aspartate subtype of the glutamate receptor studies initially attempted to determine if ALS symptomatology would be reduced although results showed no measurable decrease in rate of disease progression. In conjunction with ultra low dose quinidine to prevent rapid metabolism of dextromethorphan, Phase I studies have demonstrated improvement in pedoubarl affect. The most common side effects while taking 20 mg dextromethorphan and 10 mg quinidine although mild included light-headedness, slurred speech, and fatigue. Presently long term safety data is lacking. Due to difficulties extrapolating data from small sample sizes, this therapy is not yet FDA approved.

Botulinum Toxin (Botox®)

Botulinum toxin is a paralytic toxin that interferes with neural transmission by blocking release of acetylcholine, thus promoting muscle paralysis. In ALS patients suffering from upper esophageal sphincter hyperactivity and sialorrhea small clinical trials have demonstrated measurable increases in quality of life outcomes through botulinum administration. Specifically, patients with UES hyperactivity experienced decreases in dysphagia for up to four weeks while sialorrhea was reduced for up to two months.133

Resistance Exercise
With muscle weakness representing a significant cause of morbidity in ALS patients, surprisingly little objective data has been compiled on the effectiveness of resistance training in improving patient quality of life. Data remains largely equivocal due to existing cohort studies assessing small numbers of patients, but what data does exist is promising. One random controlled study determined improvement based on increased total ALS Functional Rating Scale as well as increases in upper and lower limb functionality and has been corroborated by studies undertaken in Canada.\textsuperscript{135-137} This is a promising area of study that will hopefully be developed more fully in the future.

**Arimoclomol**

Arimoclomol is an experimental research drug that amplifies heat shock protein gene expression during cell stress and has also been useful in prolonging the lifespan of SOD1 mutant mice. Although its exact mechanisms are poorly understood, it is assumed that Arimoclomol amplifies the active phosphorylated trimer of the transcription factor, heat shock factor-1. This could potentially increase available heat shock proteins to help chaperone the correct folding patterns in mutant SOD1 proteins.\textsuperscript{138} Efficacy studies are as yet unpublished for patients but the drug has proven to be well-tolerated and safe for human consumption at doses up to 300 mg/day with evidence strongly supporting the drug be dosed three times daily.\textsuperscript{139}

**Ghrelin**

Studies from a Korean research team have recently begun utilizing ghrelin as a potential therapeutic treatment for ALS, based on the protective effect ghrelin has on motor neurons experiencing chronic glutamate excitotoxicity. Ghrelin has been implicated in the activation of extracellular signal-regulated kinase 1/2 and phosphatidylinositol-3-kinase/Akt/glycogen synthase kinase-3β pathways. In order to confirm ghrelin’s direct involvement in these cascades, spinal cord cultures were exposed to exogenous threoxyhydroxyaspartate for three weeks in order to cause motor neuron degeneration. After this interval, application of ghrelin was able to significantly attenuate this degeneration. Ghrelin was also useful in preventing the expression of pro-inflammatory cytokines tumor necrosis factor-α and interleukin-1β, common factors in neurodegenerative inflammatory processes. From these studies, it is proposed that ghrelin is acting as a microglia deactivating factor, promoting the survival of distressed motor neurons.\textsuperscript{140}

**Nimodipine**

Nimodipine is a calcium channel blocker unsuccessful in the therapeutic treatment of ALS that was theorized to potentially antagonize excitatory amino acid receptor activation, decrease calcium entry into damaged neurons, and slow or perhaps reverse ALS. Unfortunately in a randomized, placebo-controlled, prospective, double-blind crossover study of nimodipine therapy there was no statistical significance in the rate of decline of pulmonary function or limb strength during treatment compared to control.Additionally, subjects receiving nimodipine experienced associated side effects including diarrhea, nausea, and lightheadedness.\textsuperscript{141} Researchers have abandoned this drug in ALS treatment and management at the clinical trial level, however one model utilizing SOD1(G37R) mice included nimodipine as part of a cocktail.
including minocycline and riluzole and was able to demonstrate significant delay in symptom onset, preservation in muscle strength, and increased longevity.142

**Coenzyme Q10 (CoQ10)**

CoQ10 is an antioxidant and mitochondrial cofactor and as such serves as a promising pharmacologic intervention. SOD1 mutants suffer from oxidative damage, suggesting the addition of an antioxidant may improve the clinical scenario at least slightly, and the mitochondrial damage seen in some ALS specimens offered further hope that additional mitochondrial factors would abate symptoms. However, during Phase II testing 2,700 mg were administered daily for nine months and showed no statistically significant reason to continue on to Phase III testing. There were no observed side effects in the patient population taking CoQ10, however there were no observable benefits, either.143

**Lithium**

Lithium is a monovalent ion that chemically competes with magnesium to alter activity of glycogen synthase kinase-3 beta, inositol monophosphatase, and Akt/β-arrestin2. It represented a potential candidate drug for ALS disease modification due to its effects on cellular oxidative stress, inflammatory pathways, autophagy and neurotrophism.144 A trial study in 2008 ignited interest in lithium research for ALS when functional deterioration was markedly reduced in a cohort of 16 patients over a fifteen month timeframe. However meta-analysis of 1100 ALS patients using lithium was unable to reproduce these findings.145 Phase IIb randomized trials on 67 patients and a phase III multicenter double-blinded randomized control study was concluded in 2011 that also was unable to demonstrate any significant benefit on survival outcomes.146,147 Although clinical trials have heavily supported the notion that lithium is no longer a viable research prospect for the treatment of ALS, recent studies have demonstrated that co-administration of riluzole may potentially mask potential neuroprotective effects of additional compounds, including lithium which in one trial was able to reduce ALS neurotoxicity.148

**Minocycline**

Minocycline is a member of the tetracycline class of antibiotics that has demonstrated potential neuroprotective effects as a result of its specific anti-inflammatory mechanisms and regulation of apoptosis pathways.149,150 After safety trials were established for patient use along with riluzole, clinical trials were initiated and early magnetic resonance imaging of patients on minocycline showed chemical changes in the precentral gyrus and brainstem in ALS patients.151,152 A phase III multicenter double-blinded randomized control study was completed in 2007 studying 412 ALS patients who were administered 400 mg minocycline daily. The results indicated minocycline was actually harmful to patients, conferring no beneficial effect for ALS morbidity or mortality while reducing quality of life.153,154

**Erythropoietin**

Erythropoietin (EPO) is a hormone responsible for regulation of red blood cell production and was studied in murine models as a potential disease modifying agent for ALS, initially
improving motor function and prolonging symptom onset in a dose-dependent fashion.\textsuperscript{155-157} CSF studies revealed lower EPO concentrations in ALS patients compared to controls, prompting research into potential efficacy trials.\textsuperscript{158} Although safe and well-tolerated initial studies and suggesting potential reductions in neurodegeneration, a phase III multicenter double-blinded randomized control study of 208 patients was unable to demonstrate change in the course of ALS pathology.\textsuperscript{159-161}

**Future Work**

At present there exists no cure or long-term treatment regimen for ALS. Lifespans are lengthened on an order of months with the most optimistic of therapies and the mechanism of disease pathogenesis at the cellular level is poorly understood. Promising studies on the use of autologous bone marrow mononuclear cells\textsuperscript{162} have been proven to reduce TDP-43 aggregations in limited study populations and lumbar intraspinal injection of neural stem cells\textsuperscript{163} have proven safe for continued Phase II research but as yet show no motor neuron protection. In addition to such exciting findings there are also disheartening discoveries still in their infancy. One proposed explanation for the failures of pharmacologic therapy has been an upregulation of ABC drug transporter proteins such as P-glycoprotein (P-gp) and breast cancer resistance protein (BCRP) both at mRNA and protein levels as seen in the SOD1-G93A mouse model.\textsuperscript{164} This research clearly illustrates that a cellular resistance mechanism is being utilized to some degree and presents its own set of unique challenges in future treatment strategies.

**Conclusion**

This has been an attempt at summarizing the most significant findings found presently in the literature regarding ALS disease progression, mechanism, and treatment. With the rapidly advancing rate of clinical trials and cytologic analysis there will invariably be discoveries that contribute to and potentially contradict many of the mechanisms proposed. At present the stress granule theory implicating the RNA binding proteins TDP-43, SOD1, and FUS are the most heavily implicated in ALS cellular pathology and riluzole is the only FDA approved pharmacologic intervention proven effective for disease modification and increased quality of life. Non-pharmacologic interventions such as physical therapy and BiPAP still present useful adjuncts to therapy and should be considered when treating ALS patients. As research continues to find new and more effective methods of treating ALS, these cellular pathways will become better known and serve as the foundation for improved therapies and increased survival for patients in the future.

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