Modelling amyotrophic lateral sclerosis (ALS) using mutant and CAS9/CRISSPR-corrected motor neurons from patients with C9ORF72 mutations reveals disease-specific cellular phenotypes

R. Mutihac, N. Ababneh, J. Scabera, S. Cowley, A. Talbot, Nuffield Department of Clinical Neurosciences, University of Oxford, Oxford, United Kingdom; bDepartment of Physiology Anatomy and Genetics, University of Oxford, Oxford, United Kingdom; cJames Martin Stem Cell Facility, University of Oxford, Oxford, United Kingdom

Background: The C9orf72 hexanucleotide expansion is the commonest genetic cause of ALS and Frontotemporal Dementia (FTD). In addition to cytoplasmic aggregation of phospho-TDP-43, pathological features include RNA foci and aggregations of dipeptide protein. The relative contribution of these pathologies to the disease remains unresolved.

Objective: To use human motor neurons from patients with ALS, and correction with gene editing, to resolve the key pathological features of ALS.

Methods: Induced pluripotent stem cell (iPSC) lines were generated from four ALS patients carrying the C9ORF72 repeat expansion. One line was corrected by genome editing to serve as an isogenic control. Cells were characterized functionally and pathologically.

Results: ALS/FTD iPSC line OXC9-02-02 was successfully used to target the expanded G4C2 repeat using CRISPR (clustered regularly interspaced short palindromic repeats)/Cas9-mediated homologous recombination, in the presence of plasmid DNA donor template containing a positive selection cassette. In C9orf72 iPSC-derived motor neurons, dysfunction in Ca2+ homeostasis and endoplasmic reticulum (eR) stress correlated with decreased cellular survival and reduced levels of the anti-apoptotic protein Bcl-2. Furthermore, the C9orf72 motor neurons showed evidence of abnormal protein aggregation and stress granule formation in the absence of external stress. These phenotypes were corrected by excision of the mutation by gene editing.

Conclusions: We have demonstrated that genome editing can be used to validate an ALS/FTD model system. The identification of a novel pathogenic link between C9orf72 mutations, dysregulation of calcium signalling and altered proteostasis demonstrates the value of iPSC-derived motor neurons as a cellular model for the investigation of neurodegeneration.

doi:10.1016/j.jns.2015.08.198
Background. Amyotrophic lateral sclerosis (ALS) leads to a complete limb paralysis, dysphagia and anarthria. A brain-computer interface (BCI) technology may aid ALS patients in communication and motor control.

Objective. To set up a BCI system for wireless control, by ALS patients, of a humanoid robot, with the aim to reach and grasp a glass of water.

Patients and methods. Four non-demented ALS patients were recruited. Controls were four healthy subjects, matched for demographic variables. A BCI command interpreter was used to control a humanoid robot. The task was to instruct the robot to move towards a glass of water, reach and grasp it (first item) and then bring the glass to the subject (second item). The protocol consisted of a calibration session, an online session, in which the two items are sequentially selected, and a robot session where the two items translate into high level commands. The minimal accuracy of each response and the number of errors each session were evaluated and analysed.

Results. All ALS patients completed the task (5 trials, 95.5% success). Controls performed comparably with a 100% success over the 5 trials. The minimum accuracy leading to a correct item selection for the robot movement was slightly better for ALS patients (ALS 60% vs controls 53%, p = 0.49).

Conclusions. ALS patients can successfully control a humanoid robot through a BCI system. This bears the potential to virtually restore the autonomous motion of an ALS patient, enabling him to extend his presence beyond the boundaries of his bed.

doi:10.1016/j.jins.2015.08.200

125
WFN15-1342
Motor Neuron Disease 1
Fibroblasts from patients with amyotrophic lateral sclerosis (ALS) associated with mutations in tardbp gene as model of TDP-43 proteinopathy
C. Hewamadduma¹, A. Grierson, A. Higginbottom, P. Shaw. Academic Neurology, Sheffield Institute for Translational Neurosciences, Sheffield, United Kingdom

Rationale & hypothesis: ALS is a devastating progressive neurodegenerative condition, which results in death. TARDBP encoded protein, TDP-43, has been implicated in both the sporadic and familial ALS cases. Animal models of TDP-43 have been inconclusive and the role of TDP-43 in ALS remains an enigma to date. Fibroblasts obtained from the patients carrying mutations in TARDBP gene provide a vital tool in investigation of TDP-43 due to physiological levels of TDP-43.

Methodology: Immunocytochemistry was performed on three lines of control and three different TDP-43 mutant fibroblast lines (M337V, G287V, A321V) and confocal microscopy was performed to identify general TDP-43, phosphorylated TDP-43 and anti p62 (to identify ubiquitin) localisations. Fibroblasts were also subjected to 0.5 mM arsenite and the stress response was assessed using markers of stress granules such as TIAR and HUR. Recovery after stress was also assessed.

Findings/conclusion: In keeping with findings in ALS postmortem material, relative clearing of nuclear TDP-43 was noted in mutant TDP-43 fibroblasts (p < 0.001). TDP-43 fibroblasts also showed accumulation of p62 positive aggregates (p < 0.0003), and phosphorylated TDP-43 accumulation (p < 0.001) compared to controls, suggesting that mutant TDP-43 fibroblasts share some characteristics of the surviving motor neurons from both sALS and FALS. Following exogenic stress endogenous TDP-43 localised to HUR positive stress granules (p < 0.01), formation of stress granules and their recovery were significantly impaired in mutant TDP-43 cases (p < 0.01) suggesting that dysfunction of TDP-43 dysregulates handling of exogenic stress. We suggest that this may contribute to premature degeneration of motor neurons expression mutant TDP-43 in ALS patients. Fibroblasts also form a robust and an economical platform to study TDP-43 related neurodegeneration.

I have obtained patient and Institutional Review Board (IRB) approval and local ethics committee approval.

doi:10.1016/j.jins.2015.08.201

126
WFN15-1481
Motor Neuron Disease 1
C9ORF72 repeated expansion in patients with familial amyotrophic lateral sclerosis from a Brazilian research center.
A preliminary report
G. Chadi¹, J.R. Maximino, F.M.H. Jorge, D. Callegaro. Neurology, University of Sao Paulo School of Medicine, Sao Paulo, Brazil

Background: The expanded GGGGCC hexanucleotide repeat in the non-coding region of the chromosome 9 open reading frame 72 (C9ORF72) gene is the most common genetic abnormality in familial amyotrophic lateral sclerosis (FALS).

Objective: To determine the C9ORF72 hexanucleotide repeat expansion in FALS patients from ALS Unit of Sao Paulo, Brazil.

Patients and methods: Patients with FALS from the ALS Unit of Clinics Hospital, University of Sao Paulo Medical School, Brazil have been evaluated for the presence of an expanded (GGGGCC) in C9ORF72. A repeat-primed-PCR reaction was applied to provide a qualitative assessment of the expansions. PCR products were analyzed on an ABI3730 and visualized using GeneMapper-software. A cutoff of >30 repeats combined with a typical sawtooth pattern was considered pathologic.

Results: Preliminaries results from 15 FALS patients (mean age of onset 51.40 ± 3.02 years) are shown. The repeat expansion was present in 5 FALS cases (33.3%) and 10 FALS did not present the pathologic expansion. One patient with C9ORF72 expansion presented a bulbar-onset and developed later a frontotemporal degeneration. Patients without C9ORF72 expansion had a spinal-onset disease. FALS patients with C9ORF72 expansion developed a later onset symptoms (54.80 ± 4.16 years) when compared to FALS without expansion (49.7 ± 4.07 years). A shorter lifespan was screen in C9ORF72 expansion carriers (5.0 ± 1.22 years) than C9ORF72 negative (9.10 ± 2.42 years).

Conclusion: A high frequency of C9ORF72 expansion was detected in this partial report of a small FASL sample of Sao Paulo ALS Unit. Supported by FAPESP and CNPq, Brazil.

doi:10.1016/j.jins.2015.08.202

127
WFN15-0089
Motor Neuron Disease 1
Epidemiological and clinical features of amyotrophic lateral sclerosis in Uzbekistan
D. Mirzaeva¹, A. Prohorova, H. Daminova. Neurology disease, Tashkent Medical Academy, Tashkent, Uzbekistan

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease characterized by multifactorial etiology, affection of central and
peripheral motor neurons, progressive course and onset of terminal respiratory insufficiency. The amount of patients including ones in Uzbekistan that are suffering and dying from ALS is increasing. The development of a familial ALS (FALS) shows the role of mutations of the gene producing the superoxide dismutase 1.

Data on morbidity of ALS on base of register for the period of 2013–2014 was studied. Special attention was paid to geographic epidemiology of ALS in different regions of Uzbekistan and to features of clinical manifestations of this disease in Uzbek nationality. During the analyzed period there were 3.4 cases of ALS per 100,000 per year registered. 70% were patients from Andijan valley, 30% from other regions. Gender analysis determined that 60% of patients were women (mean age 43 ± 4.2 years), 40% men (mean age 48 ± 3.4 years). Most patients associated the debut of ALS with virus disease. Disease of 80% of patients debuted by bulbar form, of 20% by spinal form. The FALS is absent in our country. The average life expectancy was 2.5 ± 0.45 years during the bulbar form and 3 ± 0.86 years during the spinal form. The onset of decompensation period on the average 6 month later after debut.

The analysis of the register of ALS in Uzbekistan demonstrated that ALS was the most common disease in Andijan valley, primary occurred in women, is not of a personal nature and mainly declares itself by bulbar form.

doi:10.1016/j.jns.2015.08.203

128
WFN15-0254
Motor Neuron Disease
Rehabilitation of blood-spinal cord-barrier toward amyotrophic lateral sclerosis therapy
B. Solomon, I. Rabinovich-Nikitin. Biotechnology-Microbiology, Tel Aviv University, Tel-Aviv, Israel

Background: ALS patients and transgenic mice expressing ALS-associated superoxide dismutase 1 (SOD1) mutations show alterations in the blood-spinal cord barrier (B-SC-B) as suggested from the reduction of levels of various tight junction proteins (TJPs) including zonula occludens-1 (ZO-1), occludin and claudin-5 between endothelial cells and early protection of the B-SC-B integrity was found to delay onset of motor-neuron impairment and degeneration.

Objective: The aim of this research was to investigate if inhibition of the axis CXCL12/CXCR4 receptor widely expressed in neurons and glial cells and modulates neuronal apoptosis, may improve motor neurons survival by increasing the expression of tight junction proteins and rehabilitation of the barrier.

Materials and methods: Transgenic mouse model of ALS were treated with AMD3100, antagonist of CXCR4. Motor function, weight changes and survival were evaluated. In a separate experiment, mice were sacrificed after one month of treatment and levels of proteins essential for the formation of the barrier in comparison with proteins that do not participate in the barrier were measured.

Results: We found that chronic administration of AMD3100 to ALS mouse model was effective in restoring the expression of tight junction proteins and considerably increase the survival, confirming the importance of early treatment for rehabilitation of the barriers to prevent infiltration of neurotoxic products and microhemorrhages.

Conclusions: These data reveal that multi-faceted action of AMD3100 may provide a novel option for ALS therapy leading to rehabilitation of B-SC-B proteins and thus preventing additional damage to motor neurons.

doi:10.1016/j.jns.2015.08.204

129
WFN15-1264
Motor Neuron Disease
Far beyond our typical dengue fever on three cases reported: weakness, visual loss and aphasia as initial clinical presentations?
A. Cronemberger-Andradea,⁎, H.R. Soares-Netoa, A.F.P. Pouza, D.D. de-Fariaa, L. Dongyangb, N.C. Bergamasco, C.A. de-Albuquerque, E.P. de-Andradeb, S.M.C.A. Silvab. aServicio de Neurología, Hospital do Servidor Público Estadual Francisco Morato de Oliveira (HSPE-FMO/ IAMSPE), São Paulo, Brazil; bServicio de Oftalmología, Hospital do Servidor Público Estadual Francisco Morato de Oliveira (HSPE-FMO/ IAMSPE), São Paulo, Brazil

Background. The mosquito-borne dengue virus human infection is endemic in Brazil and the most populated São Paulo State concentrates a great number of cases. There is a wide range of possible clinical manifestations. Secondary dengue-related diseases and complications can show up with nonspecific symptoms to more severe hemorrhagic shock. Possible neurological manifestations are also a part of initial clinical assessment of an infected patient and can define severity of affection. Objective. This cases report aims to study complications dengue fever can present with and alert health care professionals dealing with this disease.

Patients. Here we present three cases of adult patients admitted in March 2015 at our hospital during a current epidemic of dengue viral infection in the State who experienced neurological complications in the convalescent phase of infection: a man with weakness (Guilian–Barré syndrome), a woman with visual loss (unilateral maculopathy), and a woman with aphasia (post-seizure Todd’s palsy). The patients had in common a previously acute febrile disease diagnosed as dengue fever.

Results. After specific investigation and treatment, all of them had their complaints improved.

Conclusion. These various disease presentations on patients referred to neurological care should be promptly diagnosed and treated. To call attention and inform health teams on these different presentations is essential. The dissemination of new clinical guidelines for health professionals on non-tertiary services including these neurological complications might be useful and can yield better outcomes.

doi:10.1016/j.jns.2015.08.205