1. Drug Development

ALS is not one disease. Will not be solved by one treatment. Genetics discoveries point to the pathways for disease.

18 clinical trials have failed in the last decade. Failure in clinical trials is a problem, shared with other neurodegenerative disease research. Better disease modelling and better understanding of disease mechanisms will help.

Everyone is extremely optimistic about antisense oligonucleotide therapies (ASOs) for familial MND (SOD1 and C9orf72). Clinical trials beginning this year. Success treating SMA in 2016/17 gives great hope for MND.

Predictions for success in treating SOD1- and C9ORF72-related MND with ASOs range from 5-10 years.

In future, ASOs could be delivered orally (peptide-assisted delivery), or could be new morpholino antisense oligonucleotide (PMO) therapeutics. ASOs could also be useful for sporadic ALS – lowering ataxin 2.

Several other drug development avenues are being explored and any one could have a big impact.

“I believe that immunotherapy has great promise for MND.” - Prof Robert Henderson.

People with fast progressing MND have higher inflammation. Regulatory T cells correlate with a slower rate of disease progression.

Whatever causes MND results in misfolded proteins, which triggers an immune response, which causes inflammation, which causes damage to functional cells.

Cell stress leads to motor neuron death leads to motor function loss.

Motor neurons don’t die alone. Motor neuron death is non-cell autonomous, depends on a well-orchestrated dialog involving neurons, glia and T cells.

Higher cortical hyperexcitability is related to a shorter life-span; could help early diagnosis or be useful as a biomarker.

Spinal cord in ALS shows breakdown of blood-spinal cord barrier (BSCB). We need to understand better what is going on at BSCB and BBB with MND – permeability, transporters.

TDP43 gives us a target common to almost everyone with MND. TDP43 aggregates correlates with motor neuron death.
Opportunities and advances in drug development for ALS/MND

Dr Lucie Bruijn, Chief Scientist, ALS Association, USA
lucie@alsa-national.org

- Three drugs approved for ALS. Rilutek and Radicava are disease modifying. Neudexta provides pseudobulbar affect symptom relief.
- Explore therapeutics approaches as any one could make a big impact on ALS
  - small molecules
  - antisense therapy
  - biologics (stem cells, antibodies, gene therapy)
- Most promising targets are:
  - SOD1 mutations – 2% of ALS
  - C9orf72 – 10% of ALS
  - TDP43 – almost all cases of ALS
- Biggest challenge is that ALS is not one disease. Like the cancers it is a heterogeneous population. Do we think it can be solved by one treatment? I think we all agree that this is not the case.
- Failure in clinical trials not unique to ALS but shared with neurodegenerative diseases. Are we looking at correct target? Are we administering in the correct time frame?
- Other bottlenecks: disease modelling, biomarkers, clinical trial design
- Mouse models help us understand complexity of the disease. Dog model is also being widely used to test gene therapy and antisense – to help solves the problem in dogs too.
- Biomarker discovery improves diagnosis, helps stratify clinical trials.
- www.alsa.org/research/clinical-trials/

Antisense therapy & ALS Association funding:
- Antisense oligonucleotide (ASOs) therapy prevents the production of proteins in genes.
- In second clinical trial for SOD1, C9orf72 trial starting any day.
- The ALS Association has been supporting antisense therapy since 2004. From investment of $1.5 million, this mobilised an additional $100 million in research funds from industry.
- ALS Association funding leads to much more significant (x200) industry funding.
- ALS Association now has many industry partnerships (GSK, Ionis)
- ALS Association was fortunate to receive a significant boost to funding in a small time frame (Ice Bucket!).
- Best to seed fund small areas
- Genetics discoveries point to the pathways for disease.
- ALS Association has committed to investing in large data – clinical and environmental information. Really need to understand more about the whole of a person living with MND and not aspects in isolation.
- Why fund similar initiatives? Many came from different angles and bring unique opportunities.
Gene silencing therapy for ALS and beyond
Prof Don Cleveland
Ludwig Institute and Dept Cellular and Molecular Medicine, University of California San Diego, USA
dcleveland@ucsd.edu

- Development of “designer DNA” drugs about to be in trial for ALS.
- First causative gene discovered in 1993 – SOD1. When mouse model created, led to worldwide agreement that mutations provoke disease through a toxic property, not loss of activity.
- We think SOD1 is in every cell. By silencing it in motor neurons, get disease later. Deleted in astrocytes, mice lived twice as long. Deleted in microglia, lived three times as long. Deleted from oligodendrocytes, delayed onset. Conclusion that disease is non-cell autonomous, SOD1 within motor neurons and oligodendrocytes drive onset.
- Strategy for ASO delivery = inject directly into cerebral spinal fluid, so crosses BBB.
- **Inject at onset, you double survival after onset.**
- C9ORF72: Largest cause of inherited MND and FTD discovered Sept 2011. Create mice that express c9 expansion. Mice develop FTD-like behavioural abnormalities but not MND.
- What happens if lower c9 levels? Not much in nervous system. Loss of function of c9 associated with gain of toxicity. Have to lower the bad RNA but not the good RNA.
- Injecting single dose of ASO lowers RNA level in c9 mice. Did it before disease onset, one dose, suppressed acquisition of cognitive phenotype with a single dose.
- **Targets sense strand RNA (not antisense)** because can’t measure antisense.
- When people diagnosed, already have RNA toxicity. If you can turn down RNA, you turn down translation product. You won’t get dead neurons back but have a chance of getting damaged neurons back.
- Expecting to get to human trial in second quarter 2018 – just 6.5 years after the gene was discovered!
- ALS Association invested right from the beginning when no-one else would. Was not initiated with corporate money.

Success in other neurodegenerative diseases:
- ASOs will be widely used in other neurodegenerative conditions also.
- it’s possible to silence any gene we have tested to 95%.
- Huntington’s disease:
  - Reduce mutant huntingtin synthesis in cells = long-term, partial disease reversal in mice, stopping further loss of brain mass in another mouse model. Sustained gene suppression throughout the nervous system after ASO injection at base of spinal cord. Single dose injection led to 3-month efficacy of target mRNA suppression.
- Spinal muscular atrophy (SMA):
  - affects one in 90,000 children.
  - 2016 ASO injection trial stopped at halfway point because all efficacy endpoints were met.
  - Spinraza (for SMA) injected intrathecal injection (in babies – highly invasive).
- Alzheimers:
  - A clinical trial of ASO therapy to lower tau launched in 2017.
- 10-year predictions:
  - **ALS and FTD – lowering c9 (2Q, 2018)**
  - **sporadic ALS – lowering ataxin 2**
  - spinal cerebellar ataxias – lowering mutant ataxins
• chronic brain injury – lowering tau
• Parkinson’s – lowering alpha-synuclein
• AD – lowering tau
• glioblastoma – lower cell growth genes

CNS delivery of oligonucleotide for treatment of neurodegenerative diseases
Fazel Shabanpoor
Florey Institute of Neuroscience, Australia
Fazel.shabanpoor@unimelb.edu.au

• How can we deliver antisense oligonucleotides (ASOs)? Development of peptides as a delivery vector.
• ASOs are a single string of nucleic acids, interfere with gene expression by altering RNA function. Approved for SMA, Duchene Muscular Dystrophy, Huntington’s, ALS (SOD1, C9). Also in clinical trial for cancer. Really useful technology.
• All treatments so far require large doses, repeating dosing, invasive injections. Require repeat doses because big, bulky molecules. Need to cross BBB, neuronal cell membrane.
• Other vectors for delivery:
  o nanoparticles
  o liposome
  o antibodies
  o peptides
• Peptide-assisted delivery of ASOs: easy to synthesise and modify, more selective targeting, low cost of manufacturing. Downsides: short half-life.
• Peptide ASO – what would the final product look like? A daily dose? If can improve half-life to 1-2 hours that determines dosing. Won’t have to do it every day.

Genomic structural variations in ALS genomic regions
Prof Anthony Akkari
Head MND Genetics and Therapeutics Research, Perron Institute, Australia
anthony.akkari@perron.uwa.edu.au

• Currently only 2 therapeutics approved for MND that have a limited effect (riluzole and edaravone).
• 18 clinical trials have failed in the last decade alone.
• We have no diagnostics to predict who will respond to the drug and how to select those patients.
• Working on developing new morpholino antisense oligonucleotide (PMO) therapeutics and novel biomarkers called structural variations (SVs) that improve chances of success.
• PMO therapeutics are suitable for a proportion of ALS patients. A PMO drug developed for Duchenne Muscular Dystrophy received FDA approval in 2016.
• High failure rate will continue if we continue using past approaches to ALS therapeutics development. Antisense oligonucleotide technologies are essential.
NEUROINFLAMMATION

Suppressing neuroinflammation – cell-based therapy
Prof Stanley Appel, Houston Methodist Neurological Institute, USA
sappel@houstonmethodist.org

- Each time a new gene is implicated in MND, it gives us a new process that might be impaired. As you look at each process, you see that cells seek ‘help’ from other cells. Signalling beyond the motor neuron contributes to the cell’s death, because it appears motor neurons don’t die alone.
- In early stage MND, neuroprotective M2 microglia and T-reg lymphocytes slow disease progression.
- As disease accelerates: motor neurons release ‘danger signals’ that promote microglial activation to M1 proinflammatory state, downregulate neuroprotective T-regs, upregulate proinflammatory Th1 lymphocytes. This accelerates disease progression.
- In a mouse model of mSOD1 ALS, there are slow and fast progressing types. The fast progressing have higher inflammation.
- T-cells suppress neuroinflammation in the mouse model. In humans, regulatory T cells (T-regs) correlate with a slower rate of disease progression in pwMND. These “T-regs” are protective in pwMND.
- FoxP3 cells are also dysfunctional in pwMND.
- Total immunosuppression doesn’t work. You need to change the ratio of “bad guys” to “good guys”, to increase the regulatory T cells.
- Infusions of patients own T-regs on 3 pwMND normalised the rate of progression. Doing it again 16 weeks later (once a month for four months) again stabilised progression. Progression increased after treatment stopped (possible disease acceleration). So this was a positive result but not an effective long-term therapy because it lasted only 2-6 weeks. Only 3 patients so far and not placebo controlled.
- Prof Stanley Appel says pilot study has shown that cell-based therapy to suppress neuroinflammation is safe & well-tolerated: infusions of autologous T-reg cells restore immune cell function.
- A placebo controlled trial is now planned for six months, with dose escalation and infusions for one year.

Associate Professor Anthony White
QIMR Berghofer Medical Research Institute, Australia
Tony.white@qimrberghofer.edu.au

- Neuroinflammation is a major contributor to MND, largely mediated by microglia, an immune cell of the CNS
- Microglia are brain macrophages that enter our brain in the embryonic stage. They respond to cell death. The c9orf72 region is important to microglial cells.
- Microglial targeted drugs have not yet translated into successful clinical outcomes. This is because there are differences between animal and human microglia, and microglia are highly sensitive to their local environment.
- To overcome these issue we are establishing a new cell model of human microglia generated from human peripheral blood monocytes (hiMG)

John Lee
School of Biomedical Sciences, University of Queensland
j.lee9@uq.edu.au

- The complement system is an innate immune system, a component of blood.
- C5a and C5aR1 are the most potent inflammatory mediators. The C5aR1 protein increases in the microglia in the spinal cord of SOD1 mice in the later stages of MND.
In mice, the genetic absence of C5aR1 increased survival and improved motor function. The C5aR1 protein can also be knocked out in a mouse model with an irreversible drug – PMX205 – that can be taken orally and crosses the blood-brain barrier. The PMX205 treatment improved muscle strength. They also collected blood from pwMND and showed that C5a levels were elevated in plasma and leukocytes. C5aR1 is also elevated in pwMND monocyte populations.

Prof Luis Berbeito
Institut Pasteur de Montvideo, Uruguay
barbeito@pasteur.edu.uy
- Masitinib is a drug that is currently in clinical trials for asthma, MS, AD and ALS.
- It targets mast cells. It can control microphages and microglia.
- Mast cells infiltrate the muscles of humans with MND. Masitinib prevents this mast cell and macrophage infiltration into muscles in rats with ALS.
- There is hope for this drug class (tyrosine kinase inhibitors) for slowing MND progression.

STEM CELLS

Stem cells in MND research and treatment – an update
Prof Kevin Eggan
Dept of Stem Cell and Regenerative Biology, Harvard University; Director of the Stem Cell Program, Broad Institute, USA
eggan@mcb.harvard.edu
- Stem cells can differentiate into other types of cell. There are many types of stem cells. There are already many effective stem cell therapies eg bone marrow transplants.
- “Mesenchymal” stem cells derived from fibroblasts are the basis of the Brainstorm/NurOwn trial, injected into ALS patients.
- What are the oldest cells in your body? Neurons in your nervous system. Neurons don’t replicate themselves. Substantial ramifications for how brain repairs itself. Once a neuron becomes differentiated, it will never replace.
2. Disease Models and Mechanisms

GENETICS

How repeats get translated in c9orf72 ALS
Prof Peter Todd , University of Michigan Medical School, USA
petertod@med.umich.edu

- A repeat expansion of the gene C9orf72 triggers the production of toxic proteins by a process known as Repeat Associated Non-AUG (RAN) translation.
- Have C9orf72 repeat when born, may get bigger over life, what drives penetrance when get to be 50 or 60 or 70? Stress could be a driver.
- When cells undergo stresses (viral, oxidative, nutrient limitation etc), triggers stress granule formation (toxic proteins).
- RAN translation of C9ORF72 proteins kills neurones and causes neurodegeneration.
- Further research is needed to identify methods of blocking the RAN translation of C9ORF72 as a therapy and therefore stopping neurodegeneration.
- This type of RAN translation happens in other repeat expansion diseases. If you develop a drug that blocks this, you have a drug that can block multiple neurodegenerative diseases.

CORTICAL DYSFUNCTION/HYPEREXCITABILITY

Cortical dysfunction identifies regions of onset in ALS
Thanuja Dharmadasa, University of Sydeny
Nudgie6@gmail.com

- The hand is the most common site of onset of ALS – it’s a vulnerable area for neurodegeneration because it’s so highly evolved.
- Bulbar-onset patients show the greatest degree of cortical dysfunction (cortical hyperexcitability).
- Prof Dharmadasa measured cortical hyperexcitability using MRI across all four limbs.
- Measuring early cortical dysfunction marks the onset region and differences in the prognosis. This suggests that higher cortical hyperexcitability is related to a shorter life-span.
- Prominent cortical hyperexcitability was found in hand region of the motor cortex across all MND subgroups. The hand is the most commonly reported onset site in ALS. Is the hand a critical trigger in MND pathogenesis?
- PLS type do not show any cortical hyperexcitability.
- Cortical hyperexcitability is a focal phenomenon associated with a discreet area of onset. Opportunity to apply regional therapies to stop spread.
- Findings could be a useful prognosis biomarker and therapy approach aimed at modifying /reducing cortical hyperexcitability.

Mehdi van den Bos
Westmead Hospital, Australia
mehdivandenbos@gmail.com

- Cortical hyperexcitability is a key point in the cascade of neuronal death.
- Probing using Transcranial Magnetic Stimulation found that short intracortical inhibition (SICI) and short intracortical facilitation (SICF) shift towards and excitatory state early in the disease.
- Findings could help early diagnosis of cortical dysfunction or be useful as a biomarker.
BLOOD-BRAIN BARRIER / BLOOD-SPINAL CORD BARRIER

Loss of blood-spinal cord barrier integrity displays regional patterning in ALS
Emma Scotter, University of Auckland, NZ
Emma.scotter@auckland.ac.nz

- **TDP43 gives us a target common to almost everyone with MND.** Deposits in brain, spinal cord. All NZ banked MND brains have TDP43 in motor cortex.
- **TDP43 aggregates correlates with motor neuron death.** Map TDP43 across spinal cord, maps against hot spots of motor neuron loss.
- Breakdown of blood-spinal cord barrier (BSCB) might worsen outcomes for pwMND. **Spinal cord in ALS shows BSCB disturbance.** Neurotoxic blood components found in cord. Proposed to be a mechanism that may influence course of ALS.
- We don’t yet know what is causing this leakage in the spinal cord. Probably some deficit in micro-circulation. Speculate that BSCB leakage develops and resolves first in the cervical and lumbar cord, persisting in the thoracic cord at the end of life.
- Thank you to the people who donated brain and spinal cord to our research.

The blood-brain barrier: An obstacle for CNS drug delivery impacted by MND
Dr Joseph Nicolazzo, Monash Institute of Pharmaceutical Sciences
Joseph.nicolazzo@monash.edu

- **Overview of factors affecting transport of therapeutics across the blood-brain barrier (BBB).**
- About 640km of micro-vessels in the brain. Endothelial cells surround these, creating the BBB, or ‘neuro-vascular unit’. In a healthy BBB, movement between these cells is impossible. Proteins are very well organised and have tight junctions.
- It’s not impossible for molecules to get into the brain. Some transporters take glucose and other small endogenous molecules across. Peptides, antibodies can also be trafficked across the BBB (like insulin). Many scientists trying to get drugs to mimic these to get large molecules across the BBB – if changing structure of the drug doesn’t change its efficacy.
- **Could MND have an impact on BBB transport of drugs?** Blood has been found in spinal cord of pwMND. Suggestive of paracellular permeability and dysfunctional tight junctions. If large molecules detected in brain of pwMND could mean improved uptake of large molecule drugs.
- Need to be mindful of co-existing conditions, non-MND drugs could start to go into brain. Concern from pharmacological perspective.
- **We need to understand better what is going on at BBB with MND – permeability, transporters.**