NEWS
Our scoliosis and gastrointestinal guidelines booklets are available on our AussieRett website and can be freely downloaded. You can access them at http://aussierett.org.au/resources/guidelines-reports-and-books/
For access to more of our research, please visit our website: http://www.aussierett.org.au
Find summaries of our recent research publications at: http://aussierett.org.au/our-research/research-snapshots
Follow our team on Facebook: http://www.facebook.com/aussierett

PUBLICATIONS IN 2016
Members of the AussieRett team have published the following papers in 2016. Snapshots of these papers are available for you to read on our website under ‘Our Research’, or you can request full copies by emailing our team at: aussierett@telethonkids.org.au


WELCOME
Welcome to our Christmas newsletter for 2016! Wishing you all Season’s Greetings and a Happy New Year from Helen, Jenny, Amy, Kingsley and the team at the Australian Rett Syndrome Study. Have a safe, joyous, and relaxing festive season.

Thank You to All!
A big thank you to all of the families of children and adults with Rett Syndrome, for continuing to support our research by participating in the AussieRett study. Thanks very much to the members of our Parent Consumer Reference Group for your input and contribution to Rett syndrome research this year. We will continue meeting with AussieRett families so that our work is shaped by their ideas and experiences. If you are a parent who is interested in joining our teleconference meetings, please email us at aussierett@telethonkids.org.au

Throughout 2016 we have received several wonderful donations from the Rett syndrome community and beyond. Thank you to all who have contributed, and a special thank you to Taleah and Kathryn Watson and family, who held a fundraising fair to raise money for Rett syndrome.

Conference reports
Earlier this year, Helen attended the 8th World Rett Syndrome Congress in Kazan, Russia where she gave a plenary presentation entitled “Rett syndrome journeys: What Russia and the World can learn from the experience of International and population-based registries of Rett syndrome”. Helen also had the privilege of providing consultations to Russian families who had a child with Rett syndrome at a clinic held at the Children’s Hospital in Kazan. This was a humbling experience and demonstrated how far advanced management of Rett syndrome in Australia is in comparison to other countries in the world.

Later in the year, our team attended the 2016 IASSIDD (International Association for the Scientific Study of Intellectual and Developmental Disabilities) 15th World Congress in Melbourne where we had the opportunity to present our research on quality of life, sleep problems, as well as the need for interpersonos and development to study individual rare disorders. Helen’s presentation focused on the need in Rett syndrome for an international register such as InterRett to complement the Australian population-based data provided by yourselves. Helen also participated in a panel discussion regarding the genetic architecture of neurodevelopmental disabilities.

Keep in touch with AussieRett CONNECT
Every AussieRett family helps to improve our knowledge and research into Rett syndrome. Please let us know if your contact details change. We don’t want to lose you! If you would like to contact us or contribute to the next AussieRett CONNECT, please contact:
AussieRett
PO Box 855, WEST PERTH WA 6872
Email: aussierett@telethonkids.org.au
Web: http://aussierett.org.au
Phone: 08 9489 7774/7790
Mobile: 0419 956 946

We wish you a very happy and safe holiday period.
RECENT RESEARCH

Rett syndrome Fifty years on
Along with Dr Stuart Cobb from the University of Glasgow, Helen and Jenny were privileged to be invited to write a review for Nature Reviews Neurology to commemorate the 50 year anniversary of Andreas Rett’s first description of Rett syndrome in 1966. We were able to describe the clinical and biological research, that has over the last half century contributed to the immense volume of knowledge about Rett syndrome we now have today. In particular we had the opportunity to map out the milestones which have occurred in the course of this journey. We could not forget to acknowledge the roles of family organisations such as Rettsyndrome.org, originally know as IRSA, and its founder Mrs Kathy Hunter. This is a very prestigious journal and we hope the paper will reinforce the very important research progress that has been made and importantly help to raise awareness.

Bone Health
We recently developed clinical guidelines for managing bone health in order to help doctors and families reduce the risk of fractures in those with Rett syndrome. Increased physical activity, even for those unable to walk independently (where body weight supported treadmill or assisted walking is recommended) is very important. If calcium intake is sub-optimal, increased dietary calcium or administration of calcium supplements is recommended and similarly, if Vitamin D levels are low, Vitamin D supplements are recommended. We hope that families will share these publicly available guidelines with their daughter’s general practitioner and specialists.

http://journals.plos.org/plosone/article?id=10.1371/journal.pone.0146824

Sleep Problems
Sleep problems can have many impacts on children and families. Families, mainly from our international database, InterRett, contributed to our investigation of sleep problems and their specific determinants. Sleep problems in Rett syndrome occurred much more frequently than in the general population. Night waking was the most common sleep problem with nearly half of the girls and women currently waking frequently at night. Getting to sleep and staying asleep was most challenging for younger children and those with a p.Arg294* mutation. Severe seizure activity and being unable to walk were associated with a greater likelihood of daytime napping.

Quality of life
Last year, we started a program of research about quality of life (QOL). Firstly, 21 AussieRett families participated in an interview study about their daughter’s QOL. From the very rich set of interview data, we found that the important areas of QOL related to wellbeing, daily function and activities, and to community immersion. Two of the areas were new to the literature - 1) the security of routines had a greater likelihood of daytime napping. Sleep problems in Rett syndrome occurred much more frequently than in the general population. Night waking was the most common sleep problem with nearly half of the girls and women currently waking frequently at night. Getting to sleep and staying asleep was most challenging for younger children and those with a p.Arg294* mutation. Severe seizure activity and being unable to walk were associated with a greater likelihood of daytime napping.

Quality of life
Last year, we started a program of research about quality of life (QOL). Firstly, 21 AussieRett families participated in an interview study about their daughter’s QOL. From the very rich set of interview data, we found that the important areas of QOL related to wellbeing, daily function and activities, and to community immersion. Two of the areas were new to the literature - 1) the security of routines had a greater likelihood of daytime napping. Sleep problems in Rett syndrome occurred much more frequently than in the general population. Night waking was the most common sleep problem with nearly half of the girls and women currently waking frequently at night. Getting to sleep and staying asleep was most challenging for younger children and those with a p.Arg294* mutation. Severe seizure activity and being unable to walk were associated with a greater likelihood of daytime napping.

QOL comprises the important things in life and understanding this is crucial for us to be able to measure and then plan how to support QOL. As the first step, the interview data enabled us to construct a new QOL measure and we are currently “road-testing” it in a short online questionnaire. This will tell us if it does what we want and need it do to! This measure will allow much clearer identification of support needs and we look forward to putting it to use in a new family questionnaire for 2017.

Females with Rett syndrome communicate and what factors influence successful communication?
This year, team member Anna Urbanowicz completed her PhD research on the topic of communication in Rett syndrome. Her research described the communication abilities of girls and women with Rett syndrome and investigated factors that were positively and negatively associated with communication outcomes.

What was found?
• During interviews parents reported their daughters were able to express discomfort and pleasure, and make requests and choices using a variety of modalities including body movements and eye gaze. They also reported that mobility and knowledge of the communication partner influenced communication.
• Data provided by Australian and international families showed that the majority (89%) acquired speech-language abilities in the form of babble or words. Of those, most (85%) then experienced a regression in those abilities. Those with a p.Arg133Cys mutation were the most likely to use one of more words, prior to and after speech-language regression.
• Many females with Rett syndrome communicate using eye gaze or gestures. AussieRett questionnaire data (n=151) found that school-aged children had the highest scores for eye gaze. Females with better gross motor abilities had higher scores for the use of both eye gaze and gestures. The use of eye gaze did not vary across mutation groups, but those with a C-terminal deletion had the highest scores for use of gestures.
• Video footage provided in our larger video study found most 82.8% of the girls and women made a choice, most using eye gaze. Of those who made a choice, 50% did so within 8 seconds.

What does this mean for girls and women with Rett syndrome and their families?
• Females with Rett syndrome share communication strengths including the use of eye gaze and the ability to make choices.
• Communication interventions should target communicative strengths, such as the use of eye gaze, and factors shown to impact communication, including the skills of communication partners.
Anna is now working with a group of international researchers and speech-language pathologists to develop clinical guidelines for the management of communication in individuals with Rett syndrome. Just like our previous guidelines for scoliosis, gastrointestinal disorders and bone health, the communication guidelines are being developed using a consensus approach with clinicians, researchers and families. The guidelines will ensure consistent information is provided to families around the world. The guidelines will be completed by the end of this year.

For more information on this research please email Anna at: a.urbanowicz@uq.edu.au