Abstracts for the 38th Human Genetics Society of Australasia Annual Scientific Meeting
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Plenaries and Orals

Plenary 1
WHAT’S THE FUSS ABOUT INCIDENTAL FINDINGS?
OPPORTUNISTIC SCREENING AND INTERNATIONAL ATTITUDES

Anna Middleton
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While whole genome/exome sequencing in a research setting may be used to explore the genetic basis of a phenotype it also offers the chance to opportunistically screen for additional results unrelated to the research project but relevant to the participants’ future medical health (termed ‘incidental findings’, IFs). There is a wealth of medical and ethics literature supporting the feedback of IFs, yet there is limited empirical work offering a voice from both professional and public stakeholders directly affected by this. I will explore the current debate about sharing IFs in both a clinical and research setting and offer results from our web-based survey that has investigated the attitudes of 6,944 individuals from across 91 countries towards searching for and sharing incidental findings from genome research studies.

Eighty percent of participants believed that incidental findings from sequencing studies should be made available to research participants if they want them. Treatability and perceived usefulness of the data were important, with 98% personally interested in learning about life-threatening conditions that were preventable. However, only 31% of participants thought genomic researchers should actively search for incidental findings that were not relevant to their research study. Genetic health professionals held the most conservative views towards all aspects of data sharing. Ours is the largest dataset published to date on such attitudes. I will share an overview of how the debate about IFs has progressed in the United Kingdom. I will also offer information about the UK government initiative to sequence 100,000 genomes of patients in the National Health Service.

Plenary 2
COMMUNICATING ABOUT GENETICS: A FAMILY AFFAIR

Carma Bylund
Hamad Medical Corporation, Qatar

Plenary 3
DEVELOPMENT AND IMPLEMENTATION OF CLINICAL GENOMICS

John Mattick
Garvan Institute of Medical Research, Sydney, NSW, Australia

Plenary 4
FATTY ACID OXIDATION DEFECTS: PHENOTYPES, OUTCOME AND THE DIAGNOSTIC DILEMMA

Ute Spiekerkoetter
Department of Pediatric and Adolescent Medicine, University Children’s Hospital, Freiburg, Germany

Plenary 5
NEWBORN SCREENING AND THE INTERFACE WITH TREATMENT

Bridget Wilcken
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Newborn screening has been a success story for over 50 years. The early diagnosis of babies with serious, treatable disorders enables appropriate neonatal management, improved outcome, and may be life-saving. There are, however, some harms, particularly for patients with mild phenotypes who may not require any treatment. With the expansion of newborn screening to encompass more and rarer disorders this is becoming a major problem. Three examples have been chosen to illustrate mild phenotypes still not well-enough understood, what research is required and what is already ongoing to better define the need for treatment in babies with these mild forms of a disorder: the ‘old’ disorders of galactosaemia (Duarte/galactosemia compound heterozygote) and congenital hypothyroidism, and a newer inclusion in screening, very-long-chain acyl-CoA dehydrogenase deficiency (VLCADD). New treatments, especially mutation-specific therapies, mean that more disorders will be considered for inclusion in screening panels. New treatments also mean that long-term follow-up of all treated patients must be incorporated in newborn screening programs. The moves towards an agreed overarching policy for newborn screening in Australia will aid attainment of these aims locally.

Plenary 6
THE IDENTIFICATION OF NEW RNA GENES IN GWAS REGIONS ASSOCIATED WITH COMPLEX DISEASES

John Mattick
Garvan Institute of Medical Research, Sydney, NSW, Australia

Plenary 7
MORBIDITY INTELLECTUAL DISABILITY MAP OF HUMAN X-CHROMOSOME

Jozef Gecz
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Human X-chromosome represents about 5% of the human genome and ~3% of the human exome. It contains 830 protein-coding...
genes (and 100s of pseudogenes and non-coding genes). Over the past 20 years a significant effort has been put into the identification of X-chromosome disease-causing DNA sequence variation, making the X one of the best studied human chromosomes. Of the protein-coding genes (~110 (13%) genes have been implicated (published and our unpublished data) in various forms of syndromic and non-syndromic intellectual disability. In addition to this multiple regions of the X-chromosome have been involved in intellectual disability through structural variation, including copy number variation, translocation and inversion. I will present an overview of the X-chromosome sequence variation in intellectual disability and how this summarized data is helping us to identify additional X-chromosome regions likely to be implicated (or not) in intellectual disability as well as redefining the role of known XLID loci. I will also discuss some of our most recent XLID gene discoveries. In synthesizing our and international efforts so far to identify the majority if not all intellectual disability (and clinically relevant in general) loci on the X-chromosome I will point to the remaining challenges as well as to the genome-wide relevance of the

Finally, in a subset of 50 individuals with severe ID, in whom micro-array-based CNV analysis and whole exome sequencing were negative, we applied whole genome sequencing. Notwithstanding the extensive genetic prescreening, a conclusive genetic diagnosis could be established in 21 patients (42%). These included 8 de novo CNVs, including single exon and intronic deletions as well as interchromosomal duplications, 12 de novo SNVs affecting the coding region and one compound heterozygous CNV causing disease in a recessive mode. These results suggest that de novo SNVs and CNVs affecting the coding region are a major cause of severe ID.

**Plenary 10**

**NON-INVASIVE CANCER GENOME SCANNING BY PLASMA DNA ANALYSIS**

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Malignant tumors release DNA into plasma. The analysis of tumor-associated nucleic acid markers in plasma enabled the development of blood-based tests for the diagnosis, monitoring and prognostication of cancers. However, cancers are highly heterogeneous in nature. Thus, different biomarkers are required for the detection of cancers of different organs or subtypes. With the advent of next-generation sequencing, whereby billions of DNA fragments can be identified and quantified in each analysis run, our group asked if the cancer genome can be decoded directly from human plasma. In our recent studies (Chan et al., Clin Chem 2012; Chan et al., PNAS 2013), we showed that copy number aberrations and methylation profiles of cancers can be detected from plasma DNA sequencing on a genome-wide manner. Because the approach detects any genomic changes that might be associated with a cancer, it can be applied to multiple cancer types. Tumor-associated genomic abnormalities were detected in plasma of patients with cancers of the liver, breast, ovary, lung, colon, nasopharynx or smooth muscles. We analysed plasma samples collected from a patient with synchronous breast and ovarian cancers. The plasma DNA profile revealed collective genetic aberrations from both the breast and ovarian cancers. These data suggest that plasma DNA sequencing is a potential tool for studying tumoral heterogeneity and may allow one to assess the total tumor burden in an individual. The long-term goal of the research is to achieve a test that could be applied for the screening and monitoring of multiple cancer types.
organizations ENIGMA (BRCA1/2) and InSiGHT (MMR). Variant nomenclature and supporting clinical and laboratory information to inform classification has been collated for ‘unclassified’ sequence variants from clinical testing and research laboratories. Data curation identified considerable inconsistency in documentation of variant nomenclature and variant classification, poor recognition of existing publically available information on variant classification, and need to publicize and implement uniform criteria for classification. Curation and generation of clinical and laboratory information has standardized and promoted variant classification. Areas highlighted for future research include the need to conduct extensive quantitative mRNA/function analysis and family studies to assess the correlation of level and type of splicing/functional aberration with risk. Results to date demonstrate the value of international collaborative studies to facilitate standardized evidence-based classification of cancer gene variants to improve clinical genetic counseling, and patient and family management.

Plenary 12
MAPPING OF SIGNIFICANT MUTATIONS IN COMMON CANCERS
David Bowtell1,2
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The development of high throughput technologies, especially next generation DNA sequencing, means that within the next 2–3 years a complete ‘parts list’ for human cancer will be available. Collaborative studies, particularly the International Cancer Genome Consortium (ICGC) and The Cancer Genome Atlas (TCGA) are uncovering driver mutations present in human cancer that are detectable at the level of point mutation, changes in DNA copy number, structural variation, transcriptional change, and/or altered pattern of methylation. Genomic analysis of human cancer is shifting from individual tumor types to pan-cancer analyses of thousands of samples across histotypes, identifying processes common to anatomically distinct cancers and resulting in a reclassification of cancer. The presentation will focus on our work on ovarian cancer and cancers of known primary. For the ovarian studies, which are part of the ICGC, I will outline findings from whole genome sequence analysis of over 100 high grade serous cancer genomes. The cancer of unknown primary work is heavily translational and is aimed at improving the clinical management, both through identification of likely site of origin and use of mutational profiling to guide cancer care.

Sutherland Lecture
SPLASHING AROUND IN THE GENE POOL: DON’T FORGET THE PATIENT!
John Christodoulou1,2
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Plenary 13
GENETICS AND GENOMICS TO ELUCIDATE CAUSES OF RARE AND COMMON DISEASES: HOLOPROSENCEPHALY AS AN EXAMPLE
Maximilian Muenke
National Human Genome Research Institute, Bethesda, MD, USA

Plenary 14
GENOFACES AND THE ROAD TO UNEMPLOYMENT
Gareth Baynam1,2, Mark Walters1, David Gilles1, Peter Class1, Mark Shriver6, Peter LeSouef7, Stefanie Kung1, Lyn Schofield1, Matthew Bellgard4, Hugh Dawkins5, Jack Goldblatt1
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Herein we present progress in non-invasive and non-irradiating 3D facial analysis that provides deeply precise objective measurements with a view to enabling clinical unmet need in diagnosis and treatment monitoring. We focus on studies in our, and collaborating, group(s), including novel approaches to asymmetry assessment in syndromic diseases, new facial signatures of rare diseases, and treatment monitoring in metabolic and non-metabolic disorders. We also outline studies exploring the overlap between normal range and syndromic facial variation and foundation methods for converting 3D facial information to text, namely elements of morphology terms. The later may facilitate automated reporting and fusion with text mining approaches for exploring human disease biology and aiding with computer assisted diagnostics.

Plenary 15
THE IDENTIFICATION OF GENE MUTATIONS IN CRANIOSYNOSTOSIS AND THEIR RELATION TO PHENOTYPE AND SURGICAL MANAGEMENT: FROM OCTAVE CROUZON TO MASSIVELY PARALLEL SEQUENCING
Tony Roscioli1, George Elakis2, Eric Lee2, Timothy Cox2, David Moon4, Mathew Wallis6, Peter Anderson2, David Davidd, Anne Turner1, Mark Gianoutsos4, Eric Hain4, and Michael F. Buckley2
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6 SA Clinical Genetics Service, SA Pathology at Women Children’s Hospital, and Department of Paediatrics, University of Adelaide, Adelaide, SA, Australia

Craniosynostosis is one of the most common craniofacial disorders encountered in clinical genetics practice. Between 30–70% of syndromic craniosynostoses are caused by mutations in hotspots in the fibroblast growth factor receptor (FGFR) genes or in the TWIST1 genes. Here we present results from molecular testing of an Australia and New Zealand cohort of 630 individuals with a diagnosis of craniosynostosis. Data were obtained by Sanger sequencing of FGFR1-3 hotspot exons and the TWIST1 gene, as well as by copy number detection of TWIST1. Of the 630 probands, there were 231 who had one of 80 distinct mutations (36%). Among the 80 mutations, 17 novel sequence variants were detected in three of the four genes screened. In addition to the proband cohort there were 96 individuals who underwent predictive or prenatal testing as part...
Epileptic encephalopathies (EEs) are characterized by severe, frequent seizures with a detrimental effect on development. The majority of EEs have a genetic basis, with significant genetic heterogeneity (> 300 genes). A small subset are treatable. Rapid investigation of children with EE is mandatory to guide therapeutic choices and reproductive counseling. To date, the investigative process requires invasive neurometabolic procedures and financially prohibitive sequential genetic testing, with a low diagnostic yield (<10%). Next-generation sequencing (NGS) provides a compelling tool to explore the genomic basis of undiagnosed EE. Recent literature has demonstrated the utility of NGS in epilepsy diagnostics with a diagnostic yield of up to 50%. However, many studies have a paucity of phenotypic information limiting genotype-phenotype correlations and clinical prognostication.

We investigated a cohort of children affected with infantile-onset EE, categorized by clinical and electrophysiological criteria. Whole exome sequencing was performed on 10 affected children using the HiSeq 2500 platform. We performed short read sequence alignment using BWA, and variant calling using the GATK Haplotype Caller following best practices approaches. Potential causative variants were identified by a tiered approach: (1) comparison with a list of EE genes, (2) comparison of all potentially pathogenic variants with current literature, (3) trio analysis to identify de novo variants. Sanger sequencing was used to confirm candidate mutations. We identified pathogenic/likely pathogenic variants in 5 out of 10 children, including a variant in a newly described EE gene. This study highlights the importance of NGS in Mendelian disorders as a rapid diagnostic technique.

Objective: Epileptic encephalopathy (EE) presents with severe seizures associated with developmental delay. A subset has effective treatments, and therefore rapid diagnosis is a priority to guide management. The investigation of children with EE is complex but the...
majority have a genetic basis, with several hundred genes known. The cost of the investigation of children with EE is immense as the results of neurometabolic and genetic disorders often display phenotypic overlap with other diseases. Numerous invasive investigations invariably include MRI scanning (with general anaesthesia), multiple EEGs, lumbar puncture for CSF metabolites and extensive serum/urine neurometabolic testing. Individual molecular genetic tests have a low sensitivity. Identifying specific gene mutations has been challenging until the advent of next generation sequencing (NGS). The current revolution in diagnostic molecular genetics through NGS now allows the testing of all genes in a cost-effective and timely fashion. We will investigate whether a genomic diagnostic approach to children with Epileptic Encephalopathy (EE) is a viable alternative to the traditional diagnostic algorithm. Design and Participants: A retrospective observational study of a cohort of 10 patients affected with EE has been performed. All patients have undergone the traditional neurometabolic investigative approach and were subsequently tested with NGS. With a health economist we have examined the cost-effectiveness of NGS to determine whether it is a viable, rapid, high-yield and a less expensive alternative to current sequential investigations. We propose, through these results, that NGS provides a comprehensive and more cost-effective management scheme for the investigation of EE.

AAG Oral 3
NEXT GENERATION SEQUENCING PROVIDES ANSWERS FOR FAMILIES AFFECTED BY FOETAL AKINESIA, ARTHROGYPOSIS, AND SEVERE CONGENITAL MYOPATHIES

Emily Todd1, Kyle Yu1, Royston Ong1, Jennie Slee1, George McGillivray1, Cathy Koyaly-Born1, Elizabeth Thompson1,2, Christopher Barnes1,2, Golnur Haliloglu1,2, Aritana Kariminejad1, Caroline Sewry1, Anita Cairns1,2, Nigel Clarke1,2,3, Alison Colley1,2, Monique Ryan1,2, Padma Sivadorai3, Mark Davis1,5, Richard Alcock1,4, Nigel Lai1,15, Gianna Ravenscroft1

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The fetal akinesia syndromes, severe congenital myopathies, and arthrogryposes are all heterogeneous disorders with prenatal or neonatal onset. Despite several causative genes being associated with these disorders, many cases remain without a genetic diagnosis. Next generation sequencing (NGS), can identify mutations in known and novel disease genes. Genomic DNA from individuals from 40 unrelated families with fetal akinesia, a severe congenital myopathy or arthrogryposis, was subjected to NGS with the aim of identifying the genetic cause of disease. Individuals were subjected to either whole exome sequencing or a custom-designed supercapture array, consisting of 277 known neuromuscular disease genes. Variants fitting the phenotype and inheritance pattern were investigated, and confirmed using Sanger sequencing.

Analysis of NGS data resulted in a genetic diagnosis for 17/40 families, including 3/14 affected by a fetal akinesia syndrome, 9/15 by a severe congenital myopathy, and 5 of 11 with arthrogryposis. Mutations were identified in eight different known neuromuscular disease genes (CHRND, CHRN, ECEL1, GBE1, MTM1, MYH3, NEB, and RYR1) and three novel neuromuscular disease genes (KLHL40, KLHL41, and Gene X). In conclusion, the use of NGS on a cohort of families with fetal akinesia, a severe congenital myopathy or arthrogryposis resulted in a diagnosis in 42% of cases. This study highlights the capabilities of NGS in determining the genetic diagnosis in heterogeneous diseases, as well as the ability to identify novel disease genes. Finally, this study has expanded the genotype-phenotype correlations for known neuromuscular disease genes.

AAG Oral 4
PERIVENTRICULAR NEURONAL HETEROOTOPIA CAUSED BY MUTATIONS AFFECTING THE DIMERIZATION DOMAIN OF FLNA: GENOTYPE: PHENOTYPE CORRELATIONS

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Filamin A, the filamentous protein encoded by the X-linked gene FLNA, crosslinks cytoskeletal actin into three dimensional networks, facilitating its role as a signaling scaffold and a mechanosensor of extrinsic shear forces. Central to these functions is the ability of FLNA to form V-shaped homodimers through its C-terminal located filamin repeat 24. Two brothers with mild periventricular heterotopia (PH), a neuronal migration disorder typically caused by loss-of-function mutations in FLNA, are described with a missense mutation (p.Gly2593Glu) inserting a large negatively charged amino acid into the hydrophobic dimerization interface of FLNA. Additional mutations in the same vicinity in four unrelated individuals conferred a more typical presentation of PH in female heterozygotes and presumptive lethality in hemizygous males, further underscoring the exceptionally mild consequences of the p.Gly2593Glu substitution. Co-immunoprecipitation, in vitro cross-linking studies and gel filtration chromatography all demonstrated that homodimerization of isolated FLNA repeat 24 is abolished by this Gly2593Glu substitution but that extended FLNA(16-24) constructs exhibit dimerization. Collectively these observations imply that other interactions apart from that mediated by the canonical repeat 24 dimerization interface can help mediate FLNA homodimerization.
CASE STUDY OF A LONG-STANDING SUPPORT GROUP

FLEISCHER1,2, LISA BRISTOWE1,3

ABSTRACT

Peer support groups for individuals at high genetic risk have had limited success in comparison with support groups for other conditions. A recent Cochrane review of familial breast cancer highlighted the need for examination of services available for such women. As members of the Ashkenazi Jewish population are at an increased risk of carrying such mutations, Jewish women identified as carriers of a breast cancer gene (BRCA1 or 2) mutation were invited to participate in an education and support program in 2006. Seven semi-structured interviews were conducted with women diagnosed with a DSD aged between 21 to 34 years. Participants were asked to discuss their wishes, needs and expectations of support services. Interviews were audiotaped, transcribed verbatim and analysed using thematic analysis.

Women living with disorders of sex development (DSD) may have both complex genetic differences and gynaecological features which have an impact on their reproductive health. These differences can also impact on fertility, body image, self-esteem and social acceptance. This qualitative study aimed to explore the psychosocial support needs of women who have a DSD. Ten semi-structured interviews were conducted with women diagnosed with a DSD aged between 21 to 34 years. Participants were asked to discuss their wishes, needs and expectations of support services. Interviews were audiotaped, transcribed verbatim and analysed using thematic analysis.

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ASGC Oral 3
PRECONCEPTION GENETIC SCREENING: REFLECTIONS ON THE FIRST 26 PATIENTS TO UNDERGO THE COUNSYL TEST IN A PRIVATE GENETIC COUNSELING SETTING.

Ron Fleischer1,2, Lisa Bristowet1,3

ABSTRACT

The Counsyl Universal Genetic Test is a commercially available genetic test that screens for up to 417 disease-causing mutations associated with up to 108 genetic disorders. It includes cystic fibrosis, spinal muscular atrophy, Fragile X syndrome, beta thalassemia, Tay-Sachs disease and a large number of rarer, autosomal recessive disorders. It is not a direct-to-consumer test and must be ordered through a genetic counselor or doctor.

We offer this test in a private genetic counseling setting where patients pay a standard fee for a consultation and then pay for any additional pathology directly to the pathology provider.

In our first 26 patients who requested Counsyl testing, we identified 13 carriers. Two were carriers for two conditions. There was one test that failed. One couple were suspected to be epidermolysis bullosa carriers and this was confirmed; one couple were hypothesized to be Connexin 26 carriers and this was also confirmed.

In series of vignettes, we will highlight the genetic counseling challenges of offering the Counsyl Universal Genetic Test and reflect on the demographics and motivations of patients requesting this test.

All patients had a heightened awareness of the risk of having a child with a disability, and saw the Counsyl test as a way to mitigate this risk.

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Findings revealed that overall participants felt satisfied with the available support systems, chiefly family support. However, soc-
al acceptance, formation of intimate relationships, coping with the physical implications, understanding reproductive options and maintaining better health outcomes were identified by participants as issues that could be improved with psychosocial support. Professional psychosocial and peer support services were identified as essential components of condition management and participants felt these should be offered as part of standard care practice. The majority of participants highlighted the value of appropriate peer relationships and wanted more support in this area. Participants recommended the development and provision of a resource inclusive of up-to-date health information, advice on fertility pathways, and contact information for both medical, psychosocial and peer support.

Concurrent 3: Australasian Society of Inborn Errors of Metabolism Oral Presentations

ASIEOral 1
MUTATIONS IN LYRM4, ENCODING IRON-SULFUR CLUSTER BIOGENESIS FACTOR ISD11, CAUSE DEFICIENCY OF MULTIPLE RESPIRATORY CHAIN COMPLEXES

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We previously reported sequencing ~1000 genes encoding the known mitochondrial proteome in 42 infants with definitive bio-
chemical OXPHOS disorders (Calvo et al., 2012 Sci Trans Med 4:118ra10). In addition to proven diagnoses, this identified muta-
tions in a number of candidate disease genes. A patient with defi-
ciency of complexes I, II and III in muscle and liver had a homozy-
gous mutation (c.203G>T, p.R68L) in LYRM4, encoding the ISD11 protein. ISD11 forms a complex with the sulfur donor NFS1 and sta-
bilizes it. Complexes I, II and III rely on iron sulfur (Fe-S) clusters for enzyme activity. Sanger sequencing identified the same mutation in his similarly affected cousin, who had a more severe phenotype, died in the neonatal period and had deficiency of Complex IV in muscle, liver and fibroblasts, the protein level of NDUFV1 was significantly reduced by 75% on Western blotting. In patient muscle, liver and fibroblasts, the protein level of NDUFV1 was significantly reduced by 75% on Western blotting. Clinical picture of Leigh syndrome.

ASIEOral 2
WHOLE EXOME SEQUENCING CONFIRMS LEIGH SYNDROME IN A PATIENT SHOWING LITTLE BIOCHEMICAL EVIDENCE OF A MITOCHONDRIAL DISORDER


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Background: Leigh syndrome is a severe neurological disorder caused by mutations in one of more than 30 genes; most of which are associated with the mitochondrial respiratory chain (RC). Aim: To identify the genetic cause of disease in a patient with an overall clinical picture of Leigh syndrome. Patient and Methods: A girl with a clinically suspected diagnosis of Leigh syndrome was born to healthy and non-consanguineous parents after a normal preg-
nancy and delivery. She was first hospitalized at 2 years of age for febrile seizures and partial left paresis. She then presented with left-sided weakness of her arm and leg, scoliosis and worsening dys-
tonia at 6 years. Magnetic resonance imaging of her brain showed symmetrical putaminal abnormalities. Her lactic acid level was elevated on brain magnetic resonance spectroscopy but normal in blood and cerebrospinal fluid. Whole exome sequencing was used to screen for likely pathogenic variations in the patient followed by mitochondrial enzyme assay and Western blotting. Results: Whole exome sequencing uncovered compound heterozygous mutations in the NADH dehydrogenase ubiquinone flavoprotein 1 [NDUFV1] (c.1162+4A>C, resulting in skipping of exon 8, and c.G640A, p.Glu214Lys), both previously associated with complex I deficiency and Leigh syndrome. Despite normal complex I enzyme activity in patient muscle, liver and fibroblasts, the protein level of NDUFV1 was significantly reduced by 75% on Western blotting. Conclusion: Whole exome sequencing can be used to provide a definitive diagnosis of a suspected mitochondrial RC disorder even in cases where RC enzyme activity is apparently normal.

ASIEOral 3
AGRESSIVE DIETARY MANAGEMENT DRAMATICALLY IMPROVES MARKED TRANSAMINITIS AND CLINICAL PROFILE IN GLYCOGEN STORAGE DISEASE TYPE IIA

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The purpose of this report is to describe an atypical, severe presenta-
tion of glycogen storage disease (GSD) IIXa and the aggressive dietary management used to dramatically improve metabolic control. GSD IIXa (resulting from PHKA2 mutations) has been widely
regarded as a benign condition that does not warrant treatment because the childhood symptoms frequently improve with age. Patients present in childhood with hepatomegaly, short stature and ketotic hypoglycaemia. Long term complications including hepatic fibrosis or cirrhosis have only been reported recently. We present a 3-year-old boy with GSD IXα who had a massively enlarged liver with fibrosis and deranged biochemistry with hypoglycaemia and ketosis at the time of diagnosis. Structured dietary interventions including regular daytime and continuous overnight feeds, frequent doses of uncooked cornstarch and protein supplementation was initiated. Diagnostic confirmation of the disorder was performed by mutational analysis of the PHKA2 gene. Regular liver function tests, home blood glucose and ketones monitoring and semi-quantitative estimation of the liver size were used to access clinical improvement. Considerable clinical and biochemical improvements including enhanced growth velocity, energy levels, overall well-being, decreased hepatomegaly, rapidly improved liver enzymes and stabilized blood glucose and ketone levels were demonstrated with the initiation of aggressive dietary therapy. These results concur with a recent report in 2013 by Tsilianidis and colleagues who proposed that GSD IXα is not always a mild condition, but instead part of an expanding phenotypic spectrum. Consideration of intensive dietary interventions is imperative to improve quality of life in these patients.

Combined malonic and methylmalonic aciduria is most commonly due to MCDD. Some cases are due to acyl-CoA synthetase deficiency (ACSD), a benign disorder. MCDD is not currently included in the newborn screening panels in Australia. Case 1: An 11-month-old boy with a history of failure to thrive and developmental delay, presented to a regional hospital after 5 days of vomiting. He had a metabolic acidosis with pH 7.08 and was retrieved to the RCH by plane. Urine organic acids showed marked ketoacidosis and malonate 6400 mmol/mol creatinine. Plasma propionylcarnitine was normal. This was typical of ACSD. A review of our unreported newborn screening data showed an elevated malonylcarnitine (C3DC) from Case 1 but not Case 2. Aneurysmal bone cyst and abdominal masses were noted in the second child. Case 2: A 1.5-year-old girl was admitted with dehydration and hypoglycaemia after vomiting. A high MCT diet and intravenous fluids were initiated. A high MCT con-
TWIN RESEARCH AND HUMAN GENETICS

ASoC & MGSA Oral 4
GENETIC AUTOPSY OF PERINATAL DEATH: DIAGNOSIS AND DISCOVERY BY WHOLE EXOME AND WHOLE GENOME SEQUENCING
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Next generation sequencing technologies are proving to be powerful tools in the identification of genetic variants underlying rare genetic disorders. Our laboratory has undertaken whole exome sequencing (WES) and whole genome sequencing (WGS) on a number of cases of perinatal death (PD). PD is defined as the death of a fetus after a pregnancy of at least 20 weeks gestation or weighing 400 g, or of a neonate within 28 days of birth. At present, the autopsy of PD requires a complex investigation involving medical imaging, laboratory investigations, and the examination of the body and placenta. Despite such careful process, a cause may not be found in approximately 22% of cases. We hypothesized that WES or WGS would act as a ‘genetic autopsy’, providing an explanation for many cases of PD and improving the definitive diagnostic yield. We also expected to identify novel monogenic disease genes with functions critical in embryonic developmental pathways.

The validity of this approach was first demonstrated using WES to identify a novel homozgyous variant in FGF22 causing neonatal lethality in a fetus from consanguineous parents. WGS has been applied to unexplained polycystic kidney disease (PKD) in two fetuses from consanguineous parents, found to share three regions of homozygosity (ROH). No known PKD genes were located within their ROH. WGS and analysis of ROH has identified only 1 outstanding candidate gene. Segregation analysis and primary functional assays are being performed to assess pathogenicity.

Concurrent 6: Australasian Association of Clinical Geneticists Oral Presentations

AAGC Oral 5
THE QUEENSLAND RENAL GENETICS MDT CLINICAL SERVICE
Chirag Patel1, Andrew Mallet2, Helen Healy3, Julie McGaughran1
1 Genetic Health Queensland, Royal Brisbane and Women’s Hospital, Brisbane, QLD, Australia
2 Department of Renal Medicine, Royal Brisbane and Women’s Hospital, Brisbane, QLD, Australia

We report the data from the first known Renal Genetics MDT clinical service in Australia, attended by a Clinical Geneticist and Nephrologist. Since August 2013, 30 patients have been seen in the clinic based on referral type: (a) known genetic kidney disease (GKD) — disease information/genetic counseling (11), (b) suspected GKD with a family history — diagnostics (9), or (c) suspected GKD without a family history — diagnostics (10). Referrals were from nephrologists (14), medical specialists (2), and GPs (14). Diagnostic categories included: ADPKD (8), non-ADPKD renal cysts (7), nephropathy (6), metabolic (2), syndromic (3), and unknown cause for renal failure (4).

All patients were given information on the diagnosis or differential diagnoses. We delineated a family history in 16 patients (53%). Genetic counseling was provided to 21 patients (70%). Genetic tests were ordered in 14 patients (47%), indications being: a) diagnostic (14%), genetic counseling (14%), or both (72%). Of these, 3 have positive results and 11 are pending. Other non-genetic investigations (imaging/urinary studies) were ordered in 6 (20%) patients for diagnostics.

So far the successful outcomes of the clinic have been: (a) disease information and genetic counseling in one clinic appointment for 21 patients, (b) new clinical diagnoses in 2 patients, (c) new clinical and molecular diagnoses in 2 patients, and (d) molecular confirmation of a clinical diagnosis for genetic counseling in 1 patient. The positive experiences of this clinic model highlight the need for mainstreaming genetics services and the development of similar clinics in locations across Australia.

AAGC Oral 6
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While there is good evidence to support the systematic screening of colorectal cancer cases at diagnosis for Lynch syndrome, its implementation in various parts of Australia is inconsistent. We have previously established, in a large retrospective study in Western Australia, that testing for microsatellite instability in colorectal cancers from patients aged <60 years was an effective initial screen to identify individuals with LS. From these findings, MSI and/or mismatch repair protein immunohistochemical screening was recommended for all newly diagnosed CRC patients aged <60 years in WA, regardless of a family history of cancer.

Four years experience with such testing (Jan 2009–December 2012) has demonstrated that in Western Australia we identify 8–10 new LS patients per year. Many of these patients did not report a family history of cancer and would not have been identified without this screening. Approximately 17% of CRC screened for MSI/IHC were identified as appropriate for referral for LS genetic testing. The incidence of germline mutations in these cases was approximately 25%. Only a minority of these cases did not attend the WA Familial Cancer Clinic for various reasons to be discussed.

Based on our experience, three key elements required for successful population-based detection of LS in Australia will be discussed. We propose the adoption of a coordinated approach at a national level that includes the above key elements. This endeavour is a priority of the recently established Inherited Cancer Connect Partnership (ICCon), a group of clinicians and scientists focused on improving the outcomes of people with rare inherited cancer syndromes.

**AACG Oral 8**

**MUTATIONS IN THE WNT SIGNALLING PATHWAY AND EYE DEVELOPMENT**

**Nicholas Pachter**, Lyn Schofield, Jack Goldblatt, Fabienne Grieu, Benhur Amanuel, Barry Iacopetta

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2. School of Paediatrics and Child Health, University of Western Australia, Perth, WA, Australia
3. Molecular Anatomical Pathology, PathWest QEI, Perth, WA, Australia
4. School of Surgery, University of Western Australia, Perth, WA, Australia

Microphthalmia and coloboma are rare genetic conditions that may be associated with other ocular malformations including anophthalmia, anterior segment dysgenesis and cataracts. Disease genes include SOX2, OTX2, CHX10, BMP4 and RAX, but in many patients the disease gene is not known. In addition to the low mutation detection rate for patients with these conditions, there is often low penetrance and variable expression which may be due to the presence of genetic modifiers. WNT signalling is important in various developmental processes, including eye development. We sequenced a candidate gene in the WNT signalling pathway in a cohort of 180 patients with cataract/microcornea, coloboma/microphthalmia and Peters anomaly conditions, revealing 3 novel mutations in 5 probands. Additional next-generation sequencing in the 5 patients did not reveal any mutations in the known disease genes. The mutations were subjected to luciferase assay using the TOPFLASH reporter system. One mutation resulted in a significant difference in WNT activity. To further evaluate the candidate gene in early eye development we generated a mutant mouse line with variable WNT activity regulated by the dosage of the candidate gene. Subsequent examination of the morphology and markers of neuroretinal, retinal pigment epithelium (RPE) and lens development of the generated mouse mutants revealed regionalization of WNT signalling activity. We also noticed reduced expression of the neuroretinal marker Chx10 and ectopic expression of the RPE marker Mitf in the mutant mouse line. Our results demonstrate the key function of WNT signalling in precise spatial and temporal in early eye development.

**AACG Oral 9**

**CLINICAL UTILITY OF THE MODIFIED MANCHESTER SCORE FOR BRCA GENE TESTING AT GENETIC SERVICES WESTERN AUSTRALIA**

**Anna Jarmolowicz**, Helen Mountain, Guicheng Zhang, Nicholas Pachter

1. Genetic Services of Western Australia, Perth, WA, Australia
2. School of Public Health, Curtin University, Perth, WA, Australia
3. Department of Paediatrics and Child Health, University of Western Australia, Perth, WA, Australia

The purpose of the study is to evaluate the prediction tool we currently utilize, the Modified Manchester Score (MMS), to determine eligibility for BRCA testing at Genetic Services Western Australia, in light of increasing test requests and the question of exception cases. Between January 1, 2009 and December 31, 2012 546 diagnostic BRCA tests were ordered. 389 cases (71.25%) met MMS testing criteria cut-off (≥ 15). The detection rate above this cut-off was 17% (66/389), with sensitivity 89% and specificity 31%. 157 (28.75%) did not meet MMS criteria; however, they were deemed eligible for testing for various clinical reasons (exception to Manchester group). In this group the detection rate was 5.1% (8/157). Common reasons for testing below the cut-off were: no contact with family, small family size, family history of breast and ovarian cancer, epithelial ovarian cancer and triple negative breast cancer pathology. Further analysis of exception cases showed that 13/65 cases of ovarian cancer and 27/92 cases of grade 3 triple negative breast cancer did not meet MMS testing criteria. 1 (7.7%) and 2 (7.4%) mutations respectively were found in these groups. Increasing the minimum MMS testing cut-off from ≥ 15 to ≥ 16, saw no decline in the sensitivity, but a slight improvement in specificity of finding a BRCA mutation. Few mutations were missed using the MMS. Testing exception cases may result in over testing, increased economic burden and psychosocial issues for patients without improved mutation detection rates.

**AACG Oral 10**

**ISOLATED AUTISM IS NOT AN INDICATION FOR SLO TESTING**

**Peter Kaub**, Peter Sharp, Janice Fletcher

1. Genetics & Molecular Pathology, SA Pathology, Women’s & Children’s Hospital, Adelaide, SA, Australia
2. Molecular Pathology, SA Pathology, Royal Adelaide Hospital, Adelaide, SA, Australia

Smith-Lemli-Opitz (SLO) syndrome is an autosomal recessive disorder of cholesterol synthesis due to deficiency in 7-dehydrocholesterol (7-DHC) reductase. SLO is associated with specific dysmorphism, microcephaly, syndactyly, eleft palate, growth retardation, genital ambiguity and intellectual disability with autistic features. The autistic features may respond to cholesterol supplementation. Increased levels of 7-DHC are a marker for SLO and...
Background: New genomic tests provide hope for a diagnosis for many children with developmental delay. However, results generated through use of these technologies increases the complexity of communication and uncertainty during genetic consultations. Little is known about the process of pediatric genetic consultations. Aim: We explored communicative processes coupled with parent and clinician experiences of consultations where children with developmental delay were referred to a genetics clinic. Methods: This qualitative project investigated consultations across four Australian states. Theoretical framework: Symbolic Interactionism — meaning is derived, created and modified through social interactions. Data: audio-recorded consultations (n = 32), parent pre-consultation surveys (n = 32), and post-consultation interviews with parents (n = 32) and clinicians (n = 11). Content, thematic and discourse analyses were completed across the data sets. Results: While the content of consultation was similar, different communication ‘styles’ were apparent and variable approaches were used when explaining genetic results. Interviews with parents revealed that overwhelmingly, they appreciated and were reassured by the consultation. The vast majority reported a positive relationship with their clinician and felt the genetic information had been explained in a very useful manner. During interviews, clinicians reported many professional challenges working in this area, both practical and emotional, especially frustrations of being unable to answer parents’ questions regarding the cause of the child’s delay. Conclusion: Detailed analyses of three complementary data sources provided rigorous and unique perspectives on impacts regarding the introduction of new genetic technologies for clinicians and parents. Findings from this study will inform best practice in this area of medical communication.

ASGC Oral 6
IT WASN’T A DISASTER OR ANYTHING: PARENTS’ EXPERIENCES OF THEIR CHILD’S UNCERTAIN CHROMOSOMAL MICROARRAY RESULT
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2 Victorian Clinical Genetics Services, VIC, Australia
3 Murdoch Children’s Research Institute, VIC, Australia

The aim of this study was to understand parents’ experiences of an uncertain chromosomal microarray (CMA) result for their child. This research utilized a qualitative approach with a phenomenological theoretical perspective. Families were ascertained via the Victorian Clinical Genetics Services and all families had a child with 16p11.2 or 15q13.3 microdeletion. Semi-structured interviews were conducted with nine parents of eight children aged between 2 and 11 years old at the time of testing. Interviews were transcribed verbatim and thematic analysis was used to identify themes within the data. Participants were unprepared for the CMA test and accompanying abnormal result. Despite a complex perception of the extent of the child’s condition and a mixed understanding of the clinical relevance of the result, they were accepting of the limitations of current medical knowledge, and appeared to have readily adapted to the result. The test result was empowering for parents in terms of access to medical, educational and financial services; however, they articulated significant unmet support needs. Participants expressed hope for the future, in particular that more information would become available over time, and were open to the possibility of further testing for their child. This research has demonstrated that parents of children who have an uncertain CMA result appear to adapt to uncertainty and limited available information. Furthermore, parents valued honest and empathic ongoing support from health professionals despite limited information. Genetic health professionals are well positioned to provide such support, aid families’ adaptation to their situation and promote empowerment.
received, with many believing additional support services would be available if their child was diagnosed. The majority of parents believed a diagnosis was of high importance, and this not associated with the length of time without a diagnosis. This study provided insight into the challenges parents face when raising a child with an undiagnosed medical condition. It is important for genetic health professionals to be aware of these factors so that they can help their clients and implement strategies to facilitate adaptation.

**ASGC Oral 8**
**POPULATION GENETIC CARRIER SCREENING FOR CYSTIC FIBROSIS, FRAGILE X SYNDROME AND SPINAL MUSCULAR ATROPHY: EXPLORING EXPERIENCES OF CARRIERS IDENTIFIED THROUGH THE VCGS REPRODUCTIVE GENETIC CARRIER SCREENING PROGRAM**

Catherine Beard1,2, Louisa Di Pietro1,4, David Amor1,4, Alison Archibald1,4

1 University of Melbourne, Melbourne, VIC, Australia
2 Murdoch Children’s Research Institute, Melbourne, VIC, Australia
3 Genetic Support Network of Victoria, Melbourne, VIC, Australia
4 Victorian Clinical Genetics Services, Melbourne, VIC, Australia

Due to advancing genetic technologies, carrier screening for multiple inherited conditions can now be offered within the population. This research aimed to explore how women experience undergoing carrier screening for three common inherited conditions: cystic fibrosis (CF), spinal muscular atrophy (SMA) and fragile X syndrome (FXS), through the Victorian Clinical Genetics Services (VCGS) Reproductive Genetic Carrier Screening (RGCS) program. Adopting a qualitative approach using phenomenology as the theoretical framework, the study utilized in-depth semi-structured interviews, which were transcribed verbatim. The transcripts were coded using thematic analysis to identify emerging themes. Eight female participants took part in this study: five received a carrier result for SMA and three for CF. The majority of participants were pregnant during screening and described the decision to have the test as straightforward. Participants experienced emotional responses such as anxiety and stress whilst waiting for their partner’s test result and also completed online research to find out more about the relevant condition during this time. Participants supported population carrier screening, preferably offered prior to conception. The findings of this study confirmed that genetic counselors (GCs) play an essential role within this program by providing support to couples after they receive a carrier result, particularly given the varying consent processes undertaken prior to screening. The provision of suitable Internet resources and GC-facilitated guidance to access reliable information would empower couples and assist the coping process. Improving awareness of the availability of population carrier screening within the community will also help improve knowledge levels and facilitate preconception screening.

**ASGC Oral 9**
**GENETIC COUNSELING CHALLENGES IN OSTEOGENESIS IMPERFECTA**

David Silence1, Rosie Fell1, Alexandra Groves2, Cheryl Cotton1, Edwina Richard2

1 Children’s Hospital at Westmead, Sydney, NSW, Australia
2 Hunter Regional Genetics, Waratah, NSW, Australia

The Osteogenesis Imperfecta syndromes, also known as ‘Brittle Bone’ group of genetic skeletal disorders, result from mutations in over 18 distinct genes loci. In 2010, the International Nosology Committee (INCDS) grouped the phenotypes arising from these mutations into 5 major groups (Osteogenesis Imperfecta types 1-5). With the exception of OI type 5, each group shows further heterogeneity. For this reason the INCDS recommended the use of a phenotypic classification. The phenotypic heterogeneity reflects the effect of the type of mutation (nonsense, missense, splicing, deletion/insertion, frameshift), the position of the mutation in the translated polypeptide and the effect on folding and functional activity of the protein as well as its trafficking from the cell. There are still as yet unexplained mechanisms which result in wide expressivity. There is both allelic heterogeneity (mutations in the two genes COL1A1 and COL1A2 genes result in at least 4 phenotypes) and allelic heterogeneity (at least 18 distinct gene loci). The phenotypic consequences of mutations in 4 of these genes may segregate in an autosomal dominant mode and 2 of these genes in an X-linked recessive mode. We have developed counseling paradigms for these 5 phenotypic groups and used these to guide genetic counseling for these ‘disorders’. We have used these paradigms to counsel new and former patients. There is a high acceptability for the concept of preimplantation genetic diagnosis PGD in dominantly inherited disorders among young people with OI.

**ASGC Oral 10**
**BEGINNING A GENETIC COUNSELING SERVICE IN A PRIVATE OBSTETRIC ULTRASOUND PRACTICE**

Alice Weeks

Specialist Women's Ultrasound, Melbourne, VIC Australia

Genetic counseling employment has traditionally been based in public hospitals. There is, however, an increasing number of genetic counselors working in private settings. The changing face of genetic counseling practice includes the expansion of employment for genetic counselors, in particular, in private health care settings. Using case examples and clinical data, this presentation outlines the experience of setting up a successful prenatal genetic counseling service in a private obstetric ultrasound practice in Melbourne. Discussion of key professional issues and challenges includes the definition of a new role, supervision, certification, and the introduction of genetic counseling to external practitioners. The aim of this presentation is to challenge and encourage genetic counselors to seek alternative and new types of employment.
conducted research/study that is not possible in Australia. My study project was a 6-week scholarship to explore the practical management strategies to improve the quality of life of Australian children with genetic metabolic disorders. This journey commenced at the Willink Metabolic Unit, Manchester Children’s Hospital, (3 weeks) and The Royal Salford Hospital (1 week) before moving to London to visit Evelina Children’s Hospital and Great Ormond Street Hospital (1 week) with the final week attending the International Congress of Inborn Errors of Metabolism in Barcelona. Key areas identified for consideration in our metabolic unit are: (1) A lifetime metabolic service, (2) a two-staged transition to adult services, (3) an effective model for enzyme replacement therapy in the home and (4) a model for delivery of respite and palliative care in the hospice setting. This experience increased my professional knowledge in these areas and resulted in a report to health administrators in how service delivery can be delivered by different models of care that may present both improvement and cost savings.

ASIEM Oral 8
OUTCOMES OF LIVER TRANSPLANTATION FOR GLYCOGEN STORAGE DISEASE TYPE 1B (GSD1B) — THE NEW ZEALAND EXPERIENCE

Rhonda Akroyd1, Helen M Evans2, Nicola Clark1, Callum Wilson1

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Background: GSD1b is associated with significant morbidity, namely hypoglycaemia, lactacidosis, hepatomegaly, neutropenia, recurrent infections, mouth ulcers, arthritis, hyperuricemia, hyperlipidaemia, and inflammatory bowel disease (IBD). Complications may include osteoporosis and hepatic tumors with potential for malignant transformation. Liver transplantation (LT) in GSD1b treats the underlying defect, many of the clinical and biochemical manifestations and the hepatic malignancy risk. Cases: Two patients with GSD1b aged sixteen years have undergone deceased donor orthotopic LT. Both had required GCSF for neutropenia and gastrostomy insertion for nutrition support. Case 1 had recurrent hypoglycaemia, intellectual disability, IBD with recurrent oesophageal strictures, requiring multiple dilatations and stenting, poor dentition, deafness and significant social issues. Case 2 had HLA B27 positive arthritis, IBD and recurrent mouth ulcers. Both cases required nutrition support using polymeric formula and uncooked cornstarch (UCCS) during the day, and overnight Case 1 UCCS boluses and Case 2 continuous polymeric formula with additional glucose polymer. Height and BMI were <0.4th and 91st centiles respectively for Case 1 and 0.4th and 75th centiles respectively for Case 2. Discussion: Both cases are doing well at 15 and 9 months post LT with improved quality of life and improved school attendance. GCSF requirement has decreased, both are enjoying eating and Case 1 is swallowing well without oesophageal structuring. Case 2 now has quiescent arthritis, IBD appears to have significantly improved in both cases. Case 1 has normalized BMI (50th centile). Thus LT is associated with excellent outcomes for patients with significant morbidity related to GSD1b.

ASIEM Oral 9
THE DEVELOPMENT OF A GUIDELINE AND EDUCATION PACKAGE FOR THE IMPLEMENTATION AND MANAGEMENT OF BH4 THERAPY FOR PATIENTS WITH PHENYLKETONURIA (PKU) IN AUSTRALASIA

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4 Auckland City Hospital, Auckland, New Zealand
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6 King Edward Memorial Hospital for Women, Perth, WA, Australia
7 Westmead Hospital, Sydney, NSW, Australia

Aim: Tetrahydrobiopterin (BH4) is a relatively new treatment option for patients with Phenylketonuria (PKU). Funding was obtained by ASIEM Dietitians to develop a guideline and education package for the implementation and management of BH4 therapy for PKU in Australasia. Process: (1) A guideline development group was formed; (2) project officer recruited; (3) guideline scope agreed; (4) stakeholders identified: patients/carers of people with PKU and health professionals involved in PKU management in Australasia; (5) clinically relevant questions formed; (6) extensive literature review completed and appraised; (7) international survey to ‘BH4 experts’ undertaken to capture practical day-to-day experience not reported in the literature; (8) specifically designed Delphi Survey for health professionals developed to formally address issues either inadequately addressed in the literature or had evidence that could lead to contradictory recommendations and to explore issues relevant to our population and clinical setting; (9) telephone survey for PKU patients/carers to ensure patient/carer needs considered; (10) interstate ethics approval gained. The Delphi Survey was sent to 54 health professionals including clinicians, dietitians, nurses, social workers and psychologists involved in Australasian PKU management. Two rounds were completed with consensus reached on any statement with >75% agreement. The telephone survey was completed with patients/carers in Queensland, Victoria and New South Wales with representation across patient groups to identify areas of concern and education requirements related to the introduction of BH4 therapy. Outcome: The results will be used to develop the guideline and education package for dissemination and implementation as appropriate across Australasia.

ASIEM Oral 10
DIETETIC CHALLENGES IN MANAGING CARNITINE-ACYLCARNITINE TRANSCLOCASE (CACT) DEFICIENCY

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2 Department of Nutrition and Dietetics, The Children’s Hospital at Westmead, Sydney, NSW, Australia
3 Westmead Hospital, Sydney, NSW, Australia

A female infant was diagnosed with carnitine-acylcarnitine translocase (CACT) deficiency following hypoglycaemia, hyperammonia, severe cardiac dysfunction and seizures in the first week of life. She was discharged home on day 23 on a low fat diet (5% total energy from long chain fat) with total energy intake of 160kcal/kg/day from modified fat formula and glucose polymer, 4-hourly feeds with no fasting, supplementary beta-hydroxybutyrate (600mg/kg/d) and carnitine. She is now 4 years old and developing normally, with growth on the 90th centile. When well she eats age-appropriate foods supplemented with MCT, relying on a bottle for a high daily oral intake of formula. Education on appropriate food choices has been well understood and a gastrostomy was sited at 9 months of age for use...
when oral intake is poor and overnight feeds. She has a history of recurrent viral infections, with each episode life threatening, some with hypoglycemia, some with bradycardia. Dietetic challenges centre around achieving her high energy needs (2 x requirement at 180–200 Kcals/kg/day, with no evidence of malabsorption) using less than ~8% LCT, when well and unwell whilst maintaining balanced eating habits. Other challenges include distance from primary metabolic team with acute local hospital management.

Concurrent 9: Australasian Society of Cytogeneticists Oral Presentations

ASoC ORAL 5
NON-INVASIVE PRENATAL TESTING (NIPT): EXPERIENCE OF THE FIRST 500 CASES AT QFG
Nicole Martin, Peter Field
Queensland Fertility Group, Brisbane, QLD, Australia

NIPT was first commercialized in 2011, with screening offered for the detection of trisomy 21. There are now 5 commercial companies offering NIPT and the range of trisomies screened has been expanded to include 13,18 and 21, with some companies offering sex chromosome abnormalities as well as triplody. Twin screening remains to be conclusively validated. NIPT uses the fragments of fetal placental DNA circulating in the maternal plasma, referred to as the fetal fraction (ff). ff shows inter-patient variation, but generally increases with gestational age. ff is affected by maternal BMI; with a greater maternal circulating volume, the lower the fetal fraction and the more difficult a diagnosis becomes, depending on the test methodology. There are 2 main methodologies employed in NIPT, a counting method, which does not distinguish between maternal and fetal fragments in the maternal plasma and a SNP method which can distinguish between maternal and fetal fragments; this latter methodology also includes a paternal SNP screen. Maternal age in the group screened ranged from 25 to 44 years. The overall ff detected ranged from 1.2% to 23.9%. Results were obtained in a ff range of 3.3% to 23.9%. The gestational age at screening ranged from 9 to 19 weeks. Recollection, due to low ff was within the range predicted by the testing company; a result was more likely to be obtained at a lower ff if a paternal DNA swab was submitted. Very elevated BMI cases failed to yield a result after recollection.

ASoC Oral 6
GENOME WIDE SNP ARRAY IS A VALUABLE TOOL IN THE INVESTIGATION OF FETAL AUTOXYLOSS.
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Array technology has been used to routinely investigate fetal autopsy cases in our laboratory since March 2011. These autopsy cases involved fetuses that died in utero or fetuses terminated due to abnormal ultrasound findings. Array CGH investigation of autopsy cases showed the incidence of a causative copy number variant (CNV) was 4.7% (4/85). Since July 2013, SNP Array has been used to investigate 89 autopsy cases. Eleven CNVs were detected with six determined to be causative (6.7%). SNP array also detects regions of homozygosity (ROHs) which can guide future DNA sequenc-

Malignant Hyperthermia (MH) is an autosomal dominant pharmacogenetic disorder that affects the skeletal muscle calcium release from the sarcoplasmic reticulum. MH is triggered in susceptible individuals through exposure to inhalational anaesthetics during general anesthesia and results in a hypermetabolic state characterized by hyperthermia, muscle rigidity, tachycardia, hypoxaemia and metabolic acidosis, which if untreated, can lead to death. During an MH episode, the myoplasmic calcium level increases rapidly due to increased flux of calcium from the sarcoplasmic reticulum to the cytosol. In approximately 70% of MH-susceptible families mutations are found in RYR1 (skeletal muscle calcium-release channel), in 1% of cases mutations are found in CACNA1S (alpha 1 S subunit of the voltage-dependent L type calcium channel). In 30% of MH-susceptible families mutations have not yet been identified. We have used whole exome sequencing or DNA capture of a limited
set of genes combined with Next Generation Sequencing to identify candidate mutations in New Zealand/Australian families. Variants with high allele frequency in the general population and synonymous variants were filtered out. High Resolution Melting analysis was used to determine segregation of the candidate mutations in the families. Lymphoblastoid cell lines were used to functionally characterize candidate RYR1 mutations.

MGSA Oral 6
TARGETED RESEQUENCING FOR EARLY INFANTILE EPILEPTIC ENCEPHALOPATHIES IN A CLINICAL DIAGNOSTIC SETTING USING MASSIVELY PARALLEL SEQUENCING
Gladys Ho1, Elizabeth Farnsworth1, Katherine Holman1, Bruce Bennett1,2, John Christodoulou2,3
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Early-onset epileptic encephalopathies are severe disorders of cognitive, sensory and motor development. Genetic testing in this field is hampered by the highly heterogeneous nature of these disorders, with marked phenotypic variability, overlapping clinical features and the involvement of multiple genetic loci. Due to the high costs and long turn-around times, traditional methods of molecular diagnosis are usually limited to a small number of candidate genes. Massively parallel sequencing (MPS aka next generation sequencing) provides an ideal platform for improving the genetic diagnosis of these disorders, by allowing a large number of genes to be tested simultaneously at a relatively low cost. Three different targeting approaches (Illumina TruSeq Custom Amplicon, Agilent Haloplex Custom Target Enrichment and Illumina TruSight Exome) have been trialled to interrogate up to 53 genes associated with disorders of early-onset seizures, Rett syndrome, neuronal ceroid lipofuscinoses or other disorders with overlapping clinical features. All three methods successfully identified candidate disease-causing variants in our patient cohort, but coverage performance varied depending on the method of targeting. Pick-up rate for patients screened for epileptic encephalopathy genes is low, with the majority of positive findings made in patients whose specific diagnoses were identified by other strong pointers (e.g., enzyme assays or characteristic clinical presentation). Cascade testing of other family members was important in the interpretation of variants of uncertain significance. The introduction of MPS into diagnostic laboratories has greatly facilitated the genetic diagnosis in patients with epileptic encephalopathy by reducing the time and cost required for testing.

MGSA Oral 7
EXOME SEQUENCING IN MOTORNEURODE DISEASE/ SPINAL MUSCULAR ATROPHY: IMPROVED DIAGNOSTICS AND COST-EFFECTIVENESS
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Motorneurone disease and spinal muscular atrophy are significant neurogenetic disorders. It is likely that the majority of motor neuron diseases have a genetic basis; however, gene mutations have not been defined in most patients. To date, the sequential testing of single genes has been slow and expensive and genetic mutations in most affected people remain to be identified. The recent availability of Next Generation Sequencing (NGS) has improved genetic testing and gene identification. We discuss NGS genetic data combined with clinical and neurophysiological assessments to provide pathophysiological insights into the motor neuron diseases. A minimum of two exomes have been performed in each family on the Illumina platform in 20 people with patterns of inheritance consistent with autosomal dominant and recessive traits. The results have identified pathogenic mutations in approximately 50% of the cohort. A potential novel gene expressed in the motor end plate has been identified in one consanguineous family for which investigations are ongoing. An analysis of whether a genomic diagnostic approach to patients with motor neurone diseases is cost-effective and alters traditional diagnostic algorithms is also assessed.

MGSA Oral 8
FAMILIAL HEMATOPOIETIC MALIGNANCIES AND MARKERS FOR THE IDENTIFICATION OF INITIATORS AND MARKERS OF DISEASE PREDISPOSITION AND PROGRESSION
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Familial cases of haematopoietic malignancy (HM) offer the opportunity to identify novel predisposition genes that likely also impact on the more common sporadic cases. They also offer affected families a tool for screening for predisposed individuals and possibilities for treatment if the identified genes or mutations are druggable. We previously discovered GATA2 as a myelodysplastic syndrome (MDS)/acute myeloid leukaemia (AML) predisposing gene which has led to the identification of ~50 families worldwide with GATA2 related disease, and a recommended standard of care for carrier individuals who develop disease.

We have performed whole exome sequencing (WES) and/or AmpliSeq 29 gene MDS/AML panel sequencing on the Ion Proton on affected and unaffected members from 50 families predisposed to HM. To date, we have sequenced over 200 samples and have not only developed a pipeline for sequence generation, alignment and variant calling (including detection of low frequency variants to assess clonal expansion), but also for variant annotation to facilitate pathogenicity calling. We have used the power of this approach to identify new germline mutations, to follow the mutational progression of disease through therapy, remission and relapse, and to identify pathogenic initiator or driver mutations in individuals prior to the onset of symptoms. This helps catalogue markers for early diagnosis of disease onset, contributes to prognostic data, enables time for development of therapeutic approaches including searching for bone marrow donors, and adds to understanding the biology of genes and genetic interactions that are common in HMs and those that unique to specific HMs.
This 90-minute workshop aims to help conference presenters take the next steps in moving their conference presentation towards an article likely to get a respectful reading from the editors and referees of their target journal. It is based on the presenter’s recent teaching text on this topic, and reflects 20 years’ experience in helping researchers write science effectively. Participants should bring with them a paper copy of a research article relevant to their own field and published in a journal they would like to target — we will analyse aspects of these during the workshop. Topics to be addressed during the workshop include: (1) Developing article writing skills by analysing articles using applied linguistics frameworks; (2) Results as ‘driver’ developing and writing your ‘take-home message’ and selecting a target journal; (3) Article structures and referee criteria — the connections; (5) Introductions: 5 ‘stages’ to a compelling justification.

Oral 1

LARS2 VARIATIONS ASSOCIATED WITH HYDROPS, LACTIC ACIDOSIS, SIDEROBLASTIC ANAEMIA AND MULTISYSTEM FAILURE

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Mutations in LARS2 have previously been associated with Perrault syndrome (premature ovarian failure and hearing loss). LARS2 encodes mitochondrial leucyl-tRNA synthetase which is required for mitochondrial protein synthesis. In this study we report LARS2 variations associated with a severe multisystem disorder. The proband was born prematurely (27 weeks) with severe lactic acidosis, hydrodrops and sideroblastic anaemia. She had multi-system complications with hyaline membrane disease, impaired cardiac function, a coagulopathy, pulmonary hypertension, progressive renal disease, and succumbed at 5 days of age. Whole exome sequencing of patient DNA revealed compound heterozygous mutations in LARS2 (c.1289C>A; p.Ala430Val and c.1565C>A; p.Thr522Asn). The c.1565C>A (p.Thr522Asn) mutation has previously been associated with Perrault syndrome and both variants are predicted to be damaging. There was some overlap of clinical features of this patient with those reported for YARS2 mutations (sideroblastic anaemia and lactic acidosis); however, unlike YARS2 patients, muscle samples did not display any clear mitochondrial RC enzyme deficiency. Western blotting of patient muscle, liver and fibroblasts did not reveal any deficiency in LARS2 or OXPHOS protein levels. Aminoaacylation assays are being undertaken to determine the effect of each of these mutations on the catalytic efficiency of LARS2. We speculate that LARS2 mutations may result in variable phenotypes. Further specific functional studies will clarify whether the LARS2 variations identified were responsible for the severe multisystem clinical phenotype seen in this baby.

Oral 2

UQCC2 AND PET100 MUTATIONS CAUSE ASSEMBLY DEFECTS OF MITOCHONDRIAL OXIDATIVE PHOSPHORYLATION COMPLEXES III AND IV

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The 5 enzyme complexes of mitochondrial oxidative phosphorylation (OXPHOS) comprise a total of more than 80 subunits but most patients with single enzyme defects lack mutations in these subunits. We used massively parallel sequencing (MPS) of patients to identify two novel OXPHOS assembly factors and proved causality via lentiviral correction studies in patient fibroblasts. First, we identified a homozygous splicing mutation in the C6orf125 gene (now renamed UQCC2) in a consanguineous Lebanese patient with complex III deficiency, severe intrauterine growth retardation, neonatal lactic acidosis and renal tubular dysfunction. Sequence-profile based orthology prediction shows UQCC2 is an ortholog of the Saccharomyces cerevisiae complex III assembly factor, Cbp6p, although its sequence has diverged substantially. UQCC2 interacts with UQCC1, the predicted ortholog of the Cbp6p binding partner Cbp3p, and both are required for synthesis &/or assembly of the mtDNA-encoded cytochrome b. Second, we studied eight complex IV-deficient patients with Leigh syndrome from six families of Lebanese origin. Complementation analysis suggested they had mutation(s) in the same gene but targeted MPS of 1,034 genes encoding known mitochondrial proteins failed to identify a likely candidate. Targeted MPS of a linkage region on chromosome 19 identified a homozygous mutation (c.3G>C, p.Met1?) in C1orf79 (now renamed PET100). We showed it is located in the mitochondrial inner membrane and forms a ~300-kDa subcomplex with complex IV subunits. We estimated that the mutation arose at least 520 years ago, explaining how the families could have different religions and different geographic origins within Lebanon.
Oral 3
QUALITY STANDARDS FOR DATABASES OF DNA SEQUENCE VARIANTS

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It is now routine practice to compare sequence variations identified during clinical genetic testing with variants recorded in a wide range of databases as well as in the scientific literature. New technologies are producing an even greater demand for analysis and interpretation, forming a substantial proportion of the diagnostic workload. Although numerous mutation databases already exist, there are few that meet the accuracy and reproducibility required for clinical diagnostics. Current databases are of variable quality and may contain errors in variant calls, non-standardized nomenclature, incomplete pathogenicity associations and limited phenotypic information linked to genomic data. These all represent limitations and risks to the quality of patient care. No standards or equivalent mechanisms exist for the accreditation of databases to ensure the accuracy and quality of uploaded data into any central repository to meet the needs of the clinical diagnostics environment. The RCPA in collaboration with the HGSA and the Human Variome Project (HVP) is developing standards for DNA sequence variation databases intended for use in the clinical environment. The framework for the development of these standards has addressed the following key areas: purpose and scope, governance, establishment, protection privacy security, content and functionality, curation, sharing, and professional use, management and workforce training. The draft standards are discussed to demonstrate and encourage the development and management of clinical grade databases, and secure sharing of accurate and appropriately curated variants and associated phenotypes. These standards will accelerate the delivery of accurate, actionable, and efficient clinical reports to improve patient management and outcomes.

Oral 4
IN THE CONTEXT OF MULTIGENIC ANALYSES OF COMPLEX GENETIC DISORDERS, SHOULD THE PATHOGENICITY OF A VARIANT BE ASSESSED IN ISOLATION FOR A CLINICAL APPLICATION?

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Inherited cardiac diseases are clinically important, genetically determined disorders, as well as significant causes of sudden death. Genetic testing for these conditions impacts on treatment protocols and clinical management options for the patient and at-risk family members. These conditions have a complex genetic pathology. They are often heterogeneous disorders with overlapping clinical symptoms, which may not always be penetrant. Determining whether variants are pathogenic, influence clinical severity or just normal variation within the genes is complex. We use a 101-gene, SureSelect capture NGS panel to screen for the arrhythmias (AP), cardiomyopathies (CP), aortopathies (ArP) and sudden death (SD) conditions. Data analysis is through a custom pipeline and the complete coding sequences of all genes are analyzed to a minimum coverage of 30X and quality score 15. We report all variants with a <1% frequency that have evidence of clinical impact. To date we have tested more than 400 proband samples, followed by segregation analyses in at-risk family members. Clinically actionable variants were identified in 71%, 81%, 68%, 80% of patients for AP, CP, ArP and SD respectively. The initial classification, a biological classification, is based on published data, in-silico and conservation analyses. Assessing the variant in the context of other gene variants identified, evidence of clinical impact and family segregation, facilitates a clinical classification specific for the family. The clinical classification has been different to the biological classification in a number of families, emphasizing the importance of this assessment in clinical applications.

Oral 5
HYPERMETHYLATION OF GADD45A IS CO-ASSOCIATED WITH IDH1, IDH2, AND TET2 MUTATIONS AND PREDICTS POOR OVERALL SURVIVAL AND CHEMORESISTANCE IN AML

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GADD45A is a tumor suppressor gene that plays cell-type dependent roles in cellular stress co-ordinating DNA repair and de-methylation, cell cycle arrest, and pro-apoptotic or pro-survival responses. Methylation of four discrete CpG islands (CpG1-4) resides in the distal promoter of GADD45A is a hallmark of many solid tumors and has been associated with impaired cell stress signaling and reduced drug response. In AML, GADD45A silencing is widespread but poorly characterized. We have shown that hypermethylation of the distal promoter of GADD45A (CpG1-4) is a common event in AML, occurring in 93 of 222 (42%) patients. GADD45A promoter hypermethylation is associated with poor survival in AML overall (median OS: 10 vs 25mths; p = .03) and in the intermediate group of patients (median OS: 11 vs 33mths; p = .04). The association of GADD45A hypermethylation with poor overall survival was also validated in an independent patient cohort. Subsequent exome capture and Sequenom-based mutation screening has revealed a striking overlap between GADD45A hypermethylation and IDH1/2 and TET2 mutations (78%; p < .0001). This co-association suggests that GADD45A may be a key functional target of the DNA demethylation pathway governed by IDH1/2 and TET2. Consistent with this, we show in patients and AML cell lines that re-activation of GADD45A can be achieved through treatment with mutant IDH inhibitors and the hypomethylating agents including Decitabine, resulting in increased chemosensitivity to Daunorubicin. In summary, GADD45A promoter hypermethylation is a novel functional biomarker of outcome in AML and its silencing is critical to chemoresistance.
Hyperammonemia in children is frequently caused by urea cycle disorders or organic acidurias. We recently identified mitochondrial carbonic anhydrase VA (CAVA) deficiency, due to mutations in the CAV5A gene, as another cause of hyperammonemia in early childhood. CAVA is one of several carbonic anhydrase enzymes catalyzing the conversion of carbon dioxide to bicarbonate for subsequent use in bicarbonate-requiring enzymes such as carbamoylphosphate synthetase 1 (CPS1), pyruvate carboxylase, propionyl-CoA carboxylase and 3-methylcrotonyl-CoA carboxylase. We illustrate this new disorder with the case of a male who presented on day 4 of life with lethargy, weight loss, jaundice and tachypnea. Ammonia was 422 μmol/L (<50) with normal urine orotic acid, features suggestive of CPS1 or NAGS deficiency. However, lactate was increased at 8.1 mmol/L and urine organic acid screening showed increases in ketones, lactate, 3-hydroxypropion, 3-hydroxyisovaleric and 3-methylcrotonylglycine. Treatment with carglumic acid normalized ammonia levels and other biochemical abnormalities within 48 hours. The patient has remained well up to last contact at 11 months with normal metabolite levels. He was homozygous for a CAV5A splice site mutation (c.555G>A) leading to skipping of exon 4. While the metabolite levels are not as pronounced as those seen in multiple carboxylase deficiencies, CAVA deficiency can be recognized by careful interpretation of biochemical profiles provided samples are collected during hyperammonemic episodes. It responds well to treatment with a good prognosis and should be considered as part of the differential diagnosis of hyperammonemias.

Concurrent 13: Free Communications

Oral 8
USING TRU SIGHT PANELS TO CREATE DISEASE-SPECIFIC GENE PANELS FOR MASSIVELY PARALLEL SEQUENCING TESTING
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Massively parallel sequencing (MPS, aka next generation sequencing) is a powerful tool for diagnostic genetic medicine, decreasing the time and cost required for testing a large number of genes associated with overlapping phenotypes. For Mendelian disorders with a limited number of causative genes, disease-targeted gene panels may have greater clinical utility over whole exome or whole genome sequencing, by allowing greater depth of sequencing coverage and reducing the complexity of variant interpretation. On the other hand, the design and validation of a panel for every disorder of interest are both cost and time-prohibitive, especially when discovery of new candidate genes entails constant revision of the panel with each addition. Using the Illumina TruSight panels (Inherited Disorders, Exome and One), we defined a variety of disease panels focusing on genes with known clinical phenotypes. Sequencing at average read depth of 300x provided over 97% coverage over the entire panel. Point mutations and medium sized insertions and deletions (10-50 bp) were identified, providing genotype diagnoses in patients with aortic aneurysm, lipofuscinosis, progressive encephalopathy, developmental eye disorders, mitochondrial disorders, renal disorders, osteogenesis imperfecta, Fanconi anemia, autoinflammatory disorders and various metabolic disorders. TruSight panels represent a versatile option for screening genes associated with a wide range of different clinical phenotypes with the use of a single test method. Not only does this reduce the time and expense of testing, it allows for greater flexibility in reviewing subpanels of genes, maximizing coverage over genes of interest, and minimizing incidental findings.
WHOLE EXOME SEQUENCING IN GENETIC CONDITIONS CAUSED BY LARGE AND/OR MULTIPLE GENES

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Whole exome sequencing (WES) is not commonly used for determining previously identified genetic diagnoses as most laboratories favor targeted capture of candidate genes. However, such approaches cannot easily incorporate new gene discoveries. We evaluated the performance of ‘off-the-shelf’ WES capture for individuals with Marfan syndrome (MFS) and autosomal dominant osteogenesis imperfecta (OI) — caused by a small number of large genes; phaeochromocytomas (PCC) and paragangliomas (PGL) — caused by numerous genes; and short rib polydactyly syndromes (SRP), associated with large and numerous genes. WES was performed using Illumina TruSeq and Roche NimbleGen capture platforms and Illumina HiSeq2000 sequencing technology. The MFS/OI/PCC/PGL samples had been tested previously and WES results were assessed blindly and compared post-analysis. WES identified previously reported mutations in 13/13 OI cases; 9/10 MFS cases and 11/11 PCC/PGL patients. In the SRP group recessive mutations were detected in known genes in 11 of 13 cases. One further case had a single deleterious mutation in WDR60 but a second mutation was not identified. A new candidate gene was identified in the final case. WES proved to be a sensitive, efficient, and rapid means of sequencing multiple genes; however, successful mutation identification requires careful consideration of platform selection and a variety of bioinformatics approaches.

NEUROGENETIC DISEASE DIAGNOSTICS BY TARGETED CAPTURE AND NEXT GENERATION SEQUENCING

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We recently discovered a non-coding regulatory change in HCFC1 that associated with X-linked intellectual disability (XLID) in a large multigenerational family (MRX3). The single base change abolished binding of the YY1 transcription factor at a highly conserved site in the 5’UTR of HCFC1, resulting in loss of HCFC1 transcriptional repression. We employed ex-ovo models of embryonic neural development to show that this change was likely pathogenic; overexpression of HCFC1 caused cell cycle exit of neural progenitor cells (NPCs), and reductions in neurite growth of hippocampal neurons. Intriguingly, loss-of-function HCFC1 mutations have now also been reported to cause severe neurodevelopmental delay. We have identified four additional non-recurrent missense variants in HCFC1 that segregate (where evidence is available) with ID in four families, and sought functional validation of pathogenicity. HCFC1 is a modular protein that harbors transcriptional activation and repressive activities, and mutations may manifest as either gain or loss of function. We investigated the effects of both loss- and mutation of HCFC1 function on the behavior of neural cells using ex-ovo assays. In contrast to over-expression, reduced Hcfc1 expression promoted the cell cycling of NPCs at the expense of differentiation, and enhanced neurite growth of neurons. All four variants were tested for their ability to affect neurite growth when overexpressed in wild-type neurons, and in neurons depleted of endogenous Hcfc1. Our work supplies information on the pathogenicity of HCFC1 variants found in patients with ID, and identifies relevant disease mechanisms that converge on cells present during embryonic stages of brain development.
We present the case of a 16-week fetus with multiple congenital anomalies, including intrauterine growth restriction, a turricaphalike skull shape, paramedian cleft of the upper lip extending into the palate, hypoplasia of the nasal bridge and nose, four limbbastoidial postaxial hexadactyly, hydrocephalus, brain heterotopia, ventriculoespinal deficit (VSD), adrenal hypoplasia, focally prominent hepatic ductal plates and early renal cystic dysplastic change. Skeletal changes included shortened and curved long bones, short ribs, mild platyspondyly and spur-like projections of the acetabular roof. The combination of features suggested a skeletal ciliopathy like Jeune syndrome. Whole exome sequencing on fetal DNA identified compound heterozygous mutations in IFT172, encoding the IFT-B component of the intrflagellar transport complex. Simultaneously, other groups also identified mutations in IFT172 in similar cases, resulting in its identification as a new ciliopathy gene. An increasing number of skeletal ciliopathy genes are being identified, many acting in an autosomal recessive/compound heterozygous fashion with significant implications for future children in affected families. This case is presented to highlight the clinical and radiological signs that are most suggestive of a diagnosis in this spectrum and assist in directing appropriate genetic testing. It also, along with other recent breakthroughs in Jeune syndrome, highlights that exome sequencing may prove to be the most cost-effective and efficient means of determining a diagnosis.

Oral 13
SUBMICROSCOPIC DUPLICATIONS AT Xp11.2 CONTRIBUTE TO INTELLECTUAL DISABILITY
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Copy number variations (CNVs) are a common cause of intellectual disability (ID). Determining the pathogenicity of CNVs remains challenging. Here we report three males with ID with submicroscopic duplications at Xp11.2. Using the Affymetrix Cytoscan HD array we have established the extent of these duplicated regions, encompassing two known ID genes IQSEC2 and KDM5C. No other known pathogenic CNVs were identified in these patients. Patient 1 presented with mild learning difficulties and autistic features. He has two affected maternal cousins. The 400kb duplication identified disrupts the longest isoform of IQSEC2 and extending to through FAM156A. Patient 2 had global developmental delay, severe expressive speech delay and a maternally inherited 579kb duplication that extends proximal of IQSEC2 and distal of SSX2. Patient 3 is 11 yrs old, with severe ID and no words. He is not ambulant and has dysmorphic features. His 1,441kb duplication involves, in addition to the region of case 1 and 2 also HUWE1 (known ID gene) and extends to a region distal of XAGE1D. Clinically this patient is more severe than other reported SMC1A and HUWE1 patients. The duplication arose de novo. Lymphoblastic cell lines from patient-3 showed significantly elevated levels of SMCIA and KDM5C transcripts, while patient 2 had only KDM5C mRNA elevated, consistent with the extent of their CNVs. Our data suggests that submicroscopic duplications at Xp11.2 containing known ID genes IQSEC2 and KDM5C might cause ID; however, it remains to be established which one (or both) are dosage sensitive in males. Interestingly, both genes escape X-inactivation.
mechanism of generating cellular conditions conducive to progression or maintenance of the AML phenotype.

Concurrent 15: Free Communications

Oral 15
ETHICS AND GENOMIC MEDICINE: TO FEAR OR TO FRIEND?
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This study describes and critiques the ethical issues arising in genomic medicine across the lifespan. However, before delving too far into the finer points of the ethical issues, I will first examine what exactly is novel about genomic medicine from an ethical perspective. Why, for example, can’t we simply reapply existing ethical concepts and theories? Drawing on features such as scale and pervasiveness of genomic information, I will demonstrate why current ethical concepts are either over-stretched or of no use at all. Then, using relevant case studies from across the human lifespan, the ethical issues arising in genomic medicine will be described and analyzed. Particular attention will be paid to the concept of consent and its limitations in genomic medicine. Additionally, the dogma of genetic exceptionalism will be challenged. Finally, suggestions will be made for new ethical concepts that might be of particular use in genomic medicine. These will include the concept of family wellbeing and personal responsibility for health. With genomics in its infancy in Australasian health care, the timing is ideal for an ethical analysis that takes account of real-life clinical and research contexts. Ethical analysis of emerging techniques of genomics and their application will help ensure this technology is best used to benefit all of us. Results from this work will be relevant to researchers, clinicians and policy-makers working in Australasian genomic medicine.

Oral 16
A SHORT TIME, BUT A LOVELY LITTLE SHORT TIME: BEREAVED PARENTS’ EXPERIENCES OF HAVING A CHILD WITH SPINAL MUSCULAR ATROPHY TYPE 1
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Spinal muscular atrophy type 1 is a relatively common, untreatable and invariably fatal neuromuscular disorder of early childhood. Family support and genetic counseling form a vital part of the management for families affected by this condition. There is no empirical evidence of how best to support SMA type 1 families during this time. This qualitative study undertook thematic analysis of 11 in-depth interviews with 13 bereaved parents of children with SMA type 1. While individuals’ experiences were unique, common themes emerged from the data, including: experiencing shock and anticipatory grief at the time of diagnosis, processing feelings of responsibility and helplessness, experiencing multiple losses including the loss of future reproductive freedom, regaining control by making decisions about the child’s life and death, finding peace in the dying process, and feeling well supported. These findings reveal a person/family-centered perspective of the psychosocial impact of having a child with SMA type 1. Health professionals can best support such families by offering grief-specific support beginning from the time of diagnosis, participating in joint decision-making to empower parents, and acknowledging the multiple losses parents may experience.

Oral 17
COMPARING PREFERENCES OF PARENTS, PEDIATRICIANS AND GENETIC HEALTH PROFESSIONALS FOR CHROMOSOMAL MICROARRAY RESULTS
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Background: Genomic chromosomal microarray (CMA) testing for childhood investigations has increased diagnostic yields. However, CMAs also increase detection of incidental findings (IFs) and variants of unknown and uncertain clinical significance (VUS). Elucidating patient disclosure preferences may help clinicians anticipate the type of results about which patients want to be informed. Methods: A questionnaire, using hypothetical scenarios, was designed to investigate and compare the perspectives of parents, pediatricians and genetic health professionals for result disclosure. Quantitative data were analysed using ANOVA and Kruskal Wallis tests. Open text data were analysed using content analysis. Results: 147 parents, 159 pediatricians and 69 genetic health professionals participated and at least 89% of respondents in each category certainly or probably favored disclosure of VUS as well as variants of certain clinical significance, with the lowest percentage being amongst parents, who were less sure of their disclosure preferences. There was consensus among respondent groups that knowledge of a variant of certain clinical significance would provide more practical and emotional utility compared to VUS. Parents demonstrated some different perspectives to health professionals; for example, they placed more emphasis on using knowledge of a VUS when considering future pregnancies (K.Wallis: p < .001). Conclusion: This study, together with a previous study investigating the opinions of a subset of these respondents for disclosure of IFs (Turbitt et al., EJHG 2014) is the first Australian exploration of preferences for genomic result disclosure, with implications for clinical practice.

Oral 18
CHILD HEALTH AFTER PREIMPLANTATION GENETIC DIAGNOSIS
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Despite the increasing use of Preimplantation Genetic Diagnosis (PGD), there is little follow-up of children conceived using PGD beyond 2 years of age. We examined the health, wellbeing and development of school-aged children (aged 5–8 years old) conceived following PGD. A retrospective cohort was undertaken. Children conceived after in-vitro fertilization (IVF) with PGD (exposed cohort) and children conceived after IVF without PGD (unexposed cohort) at two IVF clinics in Melbourne, born between 1999 and 2008, were recruited with a 1:2 ratio. Mothers of the children completed a questionnaire asking child-specific questions regarding health and wellbeing, mental health, development, educational achievement and family-specific questions regarding family functioning and parent-child attachment. We recruited 155 and 303 participants in the exposed and unexposed cohorts respectively. There were no differences between the cohorts with regards to most child and birth outcomes, birth defects, or maternal variables. However, compared to the unexposed cohort, children in the exposed cohort were more likely to have been delivered by a caesarean section.
(χ² = 11.3, p < .001), have lower peer problems in the males (t = 2.23, p = .03), and lower prosocial behavior in the females (t = 2.9, p < .01). While no significant differences between the exposed and unexposed cohorts were found for the majority of psychological scales, there were differences when compared with the normative population data. Children in the exposed cohort appeared to have more positive outcomes in many of the measures. The data suggest that PGD does not cause adverse outcomes in children.

## TWIN RESEARCH AND HUMAN GENETICS

**PRE-IMPLANTATION GENETIC DIAGNOSIS WHEN DISCLOSING CONCEPTION TO THEIR CHILD**

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**Background:** Preimplantation genetic diagnosis (PGD) is a technique which allows genetic testing on embryos created with in vitro fertilization (IVF). Historically, this has been used for couples with a history of genetic disease, but now a form of PGD known as preimplantation genetic screening (PGS) is often offered to other couples using IVF, as a way to detect any chromosomal imbalances in the embryo. Little is known about if or how parents who use PGD share this information with their children, or if they would benefit from additional support in disclosure. **Methods:** Previously collected questionnaires from 99 women who conceived with PGD/PGS and IVF and 199 women who conceived with IVF alone were analysed using inductive content analysis. Then, a questionnaire was developed based on those results and 6 women who conceived with PGD/PGS and IVF and had disclosed conception participated in in-depth interviews, which were transcribed verbatim and analysed thematically. Both samples heavily represented the opinions of women who had undergone PGS. **Results:** Among women who conceived with PGS, disclosure of IVF tends to occur early, but the use of PGS is discussed later in the child’s life, or not at all. Women who conceived with PGS questioned the relevance of that information for their child. Women who underwent PGD for familial genetic conditions may have far greater support needs than women who used PGS, as the genetic information has great relevance for the child.

## BEING A GENETIC COUNSELOR IN THE UK: TRAINING, REGISTRATION, ROLES

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This presentation will focus on the infrastructure of genetic counseling in the United Kingdom — how and why we work the way we do. There are approximately 300 genetic counselors (GCs) in the United Kingdom. Our professional body is the Association of Genetic Nurses and Counsellors and it is this group that has created our career pathway and represents the interests of genetic counselors on a national level. In order to practice clinically we have to be registered with the Genetic Counsellor Registration Board. At the moment this is a voluntary registration; as a profession we are in the process of applying for statutory regulation by the British government. Together with the HGSA we have created reciprocal registration pathways so that Australian genetic counselors can easily utilize their training and experience in the British system.

There are four different levels of genetic counselor: trainee GC, GC, principal GC and consultant GC. The majority of genetic counselors work in the National Health Service (i.e. the public sector) in the 23 regional clinical genetics services seeing patients. It is very unusual for genetic counselors to work in private practice as clinical genetics, as a service, only exists in the public sector. However, there are small numbers of genetic counselors attached to private IVF and breast cancer services. Experienced genetic counselors work autonomously with their own case load. I will explore the mechanisms for applying to work in the UK, provide an overview of the sorts of roles available and the salaries that can be expected.
through family members being mistaken or confused; only rarely do they occur through deliberate falsification. My patient’s anxiety regarding her cancer risk was based on her mother’s supposed cancer history. As arranged by my patient, her mother reported the family history on her behalf. However, the mother had been seen at the genetics service 16 years previously and irreconcilable discrepancies in the reported family histories were noted. The mother’s previous genetics appointment had been to review her cancer risk as she was requesting prophylactic mastectomies. She was now reporting multiple cancer diagnoses in herself, family members previously reported as deceased were now reported as living, and a previously reported daughter with breast cancer was an unrelated step-daughter. Without the mother’s written consent to share information, we were left with questions as to what could be disclosed to my patient. The particular characteristics of this case allowed us to avoid a controversial breach of confidentiality; however, it was necessary to explore the ethics of confidentiality to ensure we were properly prepared.

**ETHICAL CHALLENGES ASSOCIATED WITH A REQUEST FOR PRENATAL CARRIER TESTING FOR MECP2 DUPlication SYNDrome**

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Being a female carrier of an X-linked recessive condition usually has no implications for the woman’s health. Instead, implications of carrier testing relate to the future reproductive choices for that individual. Carrier testing of children is usually not recommended in the absence of an immediate medical benefit. In addition, prenatal carrier testing presents the challenge of determining the primary patient, as maintaining the future child’s autonomy concurrently denies the autonomy of the parent. In our case, a female carrier of MECP2 duplication syndrome requested prenatal carrier testing with the intention of having a termination of pregnancy for an affected male or a carrier female. While males affected by MECP2 duplication syndrome have severe to profound intellectual disability, the majority of female carriers are asymptomatic. The genetic counselor was challenged to deal with the ethical implications of this request and required to develop strategies for resolving the ethical conflict between the counselor and consultand.

**UNIQUE CHALLENGES OF THE CARDIAC GENETIC COUNSELOR**

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K is a 42-year-old female who presented after recurrent blackouts. Investigations lead to a diagnosis of long QT syndrome (LQTS). LQTS is a primary arrhythmogenic disorder affecting 1 in 3,000 of the population. It is an autosomal dominant disease with 13 genes currently known. Patients with LQTS often present with blackouts and the most serious outcome is sudden cardiac death. Patient K had clinical characteristics known to predispose to a higher risk of sudden cardiac death, and therefore an implantable cardioverter defibrillator (ICD) was recommended. Within 6 months after implantation, K disclosed suicide ideation and requested her ICD be turned off. An overview of the case will be covered with a focus on the counseling issues raised and the subsequent interventions required to address these. Discussion will also include the ethical principles of a consultand’s autonomous decision conflicting against the ethical principle of non-maleficence. Patient K presents a common scenario in the cardiac genetic setting, whereby coming to terms with the genetic issues is complicated by the issue of sudden cardiac death prevention. There is growing literature focusing on the psychosocial burden of having an ICD and dealing with ICD complications and shocks. The genetic counselor plays a key role in the multidisciplinary care of genetic heart disease patients, particularly those with ICDs.