Species-independent identification of known and novel recurrent genomic entities in multiple cancer patients

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MEETING ABSTRACTS

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O1

The metabolomics approach to autism: identification of biomarkers for early detection of autism spectrum disorder
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Human Genomics 2016, 10(Suppl 1):O1

Objectives
From the first description by Leo Kanner [1], autism has been an enigmatic neurobehavioral phenomenon. The new genetic/genomic technologies of the past decade have not as productive as originally anticipated in unveiling the mysteries of autism. The specific etiology of the majority of cases of autism spectrum disorder (ASD) is unknown, although numerous genetic/genomic variants and alterations of diverse cellular functions have been reported. Prompted by this failure, we have investigated whether the metabolomics approach might yield results which could simultaneously lead to a blood-based screening/diagnostic test and to treatment options.

Methods
Plasma samples from a clinically well-defined cohort of 100 male individuals, ages 2-16+ years, with ASD and 32 age-matched typically developing (TD) controls were subjected to global metabolomic analysis.

Results
We have identified more than 25 plasma metabolites among the approximately 650 metabolites analyzed, representing over 70 biochemical pathways, that can discriminate children with ASD as young as 2 years from children that are developing typically. The discriminating power was greatest in the 2-year age group and weaker in older age groups. The initial findings were validated in a second cohort of 83 children, males and females, ages 2-10 years, with ASD and 76 age and gender-matched TD children. The discriminating metabolites were associated with several key biochemical pathways suggestive of potential contributions of increased oxidative stress, mitochondrial dysfunction, inflammation and immune dysregulation in ASD. Further, targeted quantitative analysis of a subset of discriminating metabolites using tandem mass spectrometry provided a reliable laboratory method to detect children with ASD.

Conclusion
Metabolic profiling appears to be a robust technique to identify children with ASD ages 2-10 years and provides insights into the altered metabolic pathways in ASD, which could lead to treatment strategies.

References

Disclosure of interest
None declared.

O2

Phenome-wide association study for smoking- and drinking-associated genes in 26,394 American women with African, Asian, European, and Hispanic descents
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Human Genomics 2016, 10(Suppl 1):O2

Objectives
To uncover novel traits associated with nicotine and alcohol use genetics, we performed a phenome-wide association study in a large multi-ethnic cohort.

Methods
We investigated 7,688 African-Americans (AFR), 1,133 Asian-Americans (ASN), 14,081 European-Americans (EUR), and 3,492 Hispanic-Americans (HISP) from the Women’s Health Initiative, analyzing risk alleles located in the CHRNA5–CHRNA3 locus (rs8034191, rs1051730, rs12914385, rs2036527, and rs16969968) for nicotine-related traits and ADH1B (rs1229984 and rs2066702) and ALDH2 (rs671) for alcohol-related traits with respect to anthropometric characteristics, dietary habits, social status, psychological circumstances, reproductive history, health conditions, and nicotine- and alcohol-related traits.

Results
The investigated loci resulted associated with novel traits: rs1229984 were associated with family income (p=4.1*10−6), having a pet (p=6.5*10−6), and smoked at least 100 cigarettes (p=7.3*10−6), and number of term pregnancies (p=1.1*10−6) in EUR; rs1051730 and rs16969968 showed a suggestive association with “High cholesterol requiring pills ever” (p=3.8*10−5 and p=1.8*10−5) in trans-ethnic meta-analysis. We also replicated the known associations: rs8034191 was associated with cigarettes per day (CIGSDAY, p=3.4*10−5), smoking status (p=2.7*10−3), and “smoked at least 100 cigarettes” (p=7.3*10−6) in EUR; rs1051730 and rs16969968 were associated with CIGSDAY (p=9.1*10−8 and p=1.1*10−7) and lung cancer (p=7.3*10−3 and p=9.9*10−3) in EUR; rs2036527 was associated with CIGSDAY (p=3.5*10−5) in AFR; rs1229984 showed associations for alcohol servings (ALC, p=2.9*10−4), beer servings (p=3.1*10−4), wine servings (WINE, p=3.9*10−4), liquor servings (p=5.5*10−5), dietary alcohol (DIETALC, p=6.1*10−4), “Drinks alcohol (age 50)” (p=9.3*10−4) in EUR and for ALC (p=5.2*10−5) and DIETALC (p=9.6*10−5) in HISP; rs671 resulted associated with alcohol intake (p=3.8*10−5), DIETALC (p=1.9*10−5), ALC (p=1.3*10−5), WINE (p=1.1*10−5) and “Drank 12 alcoholic beverages ever” (p=1.2*10−5) in ASN.

Conclusion
We provided novel genetic data regarding the consequences of smoking and drinking behaviors and confirmed ethnic differences in their genetic predisposition.

Disclosure of interest
None declared.
Effects of prenatal environment, genotype and DNA methylation on birth weight and subsequent postnatal outcomes: findings from GUSTO, an Asian birth cohort

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Human Genomics 2016, 10(Suppl 1):O3

Objectives
Prenatal environment and genetic polymorphism can have a lasting impact on offspring’s metabolic function by perturbing its epigenome. Birth weight is often used as a surrogate for the overall quality of the intrauterine environment. We present the first neonate epigenome-wide association study in an Asian mother-offspring cohort, that interrogates the effects of prenatal environment variables, umbilical cord DNA methylation and SNPs, on birth weight.

Methods
In GUSTO, a prospective mother-offspring cohort study (N=987), we examined the associations between DNA methylation, SNPs, birth weight and 11 prenatal environment variables. First, we investigated the association between perinatal methylation and birth weight to identify sites of variability in methylation. Second, we interrogated the contribution of genetic and prenatal environmental factors on this variability in the epigenome. Finally, we examined whether these methylation marks at birth were associated with offspring size and adiposity in early childhood.

Results
Methylation levels at 50 CpGs were significantly associated with birth weight, and a subset of these CpGs was located in genes and miRNA known to be involved in metabolic pathways/disorders. We further examined the influence of environmental and genetic factors on methylation at these 50 CpG sites. Sixteen CpGs were associated with both, an additional 24 CpGs were associated with only environmental factors, while only 3 CpGs were associated with genetic factors alone. Environmental factors associated with methylation were predominantly maternal-adiposity-related (pre-pregnancy body mass index, pregnancy weight gain and maternal glucose levels). Methylation levels at half of these CpGs were also associated with offspring size and adiposity in early childhood.

Conclusion
Developmental pathways to obesity begin before birth and involve genetic, epigenetic and environmental factors.

Disclosure of interest
X. Lin: None declared, I. Y. Lim: None declared, Y. Wu: None declared, A. L. Teh: None declared, L. Chen: None declared, J. M. Aris: None declared, S. E. Soh: None declared, M. T. Tint: None declared, J. L. MacLacsa: None declared, F. Yap: None declared, K. Kwek: None declared, S. M. Saw: None declared, M. S. Kabor: None declared, M. J. Meaney: None declared, K. M. Godfrey Conflict with: KMG has received reimbursement for speaking at conferences sponsored by companies selling nutritional products. He is part of an academic consortium that has received research funding from Abbott Nutrition, Nestec and Danone, Y. S. Chong Conflict with: YSC has received reimbursement for speaking at conferences sponsored by companies selling nutritional products. He is part of an academic consortium that has received research funding from Abbott Nutrition, Nestec and Danone, J. D. Holbrook: None declared, Y. S. Lee: None declared, P. D. Gluckman: None declared, N. Karnani: None declared.

High-throughput identification of specific QT interval modulating enhancers at the SCN5A locus

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Objectives
Genome-wide association studies (GWAS) have indicated that sequence variation in cis-regulatory elements (CRE) plays important roles in common disease risk/trait variation, but identification of these causal variants has remained a major challenge in complex trait genetics. Here, we performed reporter assays for all common variants at the QT interval associated SCN5A GWAS locus, with the goal of identifying the underlying causal variants.

Methods
A target region of ~500kb at SCN5A was defined based on recombination hotspots (rate>10cM/Mb; HapMap) flanking the 5 independent QT interval GWAS hits. Within the target region, all common variants (minor allele frequency >5%) from the 1000 Genomes European ancestry populations in moderate linkage disequilibrium (r2>0.3) with any of the 5 GWAS hits were selected. Both alleles of these variants were amplified with flanking sequences and cloned upstream of a minimal promoter driven firefly luciferase gene in pGL4.23. Human cardiomyocyte cells, AC16, were transfected with test constructs and Renilla luciferase vector (for transfection normalization) in triplicate and luciferase assays were performed 24h later. Reporter assays on a subset of variants were repeated for assessing allelic difference in regulatory activity. All cloning and reporter assays were performed in 96- and 24-well plates.

Results
Of a total 121 variants selected, 112 variants in 104 amplicons passed primer design (amplicon size 256-617bp; median 397bp), and we successfully cloned both alleles for 106 variants in 98 amplicons. In reporter assays, compared to empty vector, 24 and 40 amplicons showed enhancer (>2-fold) and suppressor (<0.5-fold) activities in AC16 cells, respectively. Of these only 4 were observed as open chromatin regions in heart tissue in NIH Epigenomics data. Overall, 12 variants showed nominally significant allelic difference (P<0.05) in reporter activity and were repeated with 18 replicates and 7 variants were identified to have repeated significant allelic difference in regulatory activity.

Conclusion
Independent of the available epigenomic data, which are of limited relevance, an unbiased in vitro reporter screen for CREs overlapping all common variants associated with QT interval at the SCN5A GWAS locus identified 7 common cis-regulatory variants. Our immediate next goals are to a) evaluate the effect of deleting these 7 CREs on SCN5A expression in AC16 cells and b) identify the trans-acting factors regulating their functions.

Disclosure of interest
None declared.

Identification of extracellular matrix components inducing cancer cell migration in the supernatant of cultivated mesenchymal stem cells

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Human Genomics 2016, 10(Suppl 1):O5

Objectives
High-throughput identification of extracellular matrix components inducing cancer cell migration in the supernatant of cultivated mesenchymal stem cells

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Human Genomics 2016, 10(Suppl 1):O5
Objectives

Some cancers show a strong tendency to metastasize to bone, a tissue of mesenchymal origin and a prominent site of mesenchymal stem cells (MSC) residing in the stem cell niche. With bone metastasis formation being one of the most detrimental steps in cancer progression, a better understanding of how bone metastases are initially formed is key to successfully targeting bone metastasis of, for example, prostate cancer. Recent reports have suggested that bone-metastasizing cancers might mimic the process of homing of hematopoietic stem cells to their bone niche.

Methods

In order to understand the role of MSC in metastasis formation, we investigated the interaction of primary human bone marrow MSC with established cancer cell lines able to metastatize to bone. With a trans-well migration assay we could show that MSC induced a rapid migration response of prostate and breast cancer cell lines already within two hours after start of the experiment. In order to identify factors stimulating cancer cell migration, MSC cell culture supernatant was separated by size exclusion and ion exchange chromatography. Migratory fractions then were further analyzed by mass spectrometry and antibody array analysis.

Results

With this approach we identified the extracellular matrix proteins type I and type III collagen, fibronectin and laminin 421 as potential drivers of cancer cell migration, which was confirmed by using recombinant proteins. RNAi experiments showed that the cancer cell extracellular matrix receptor beta 1 integrin obviously plays a pivotal role for cell migration.

Conclusion

From our results we conclude that the extracellular matrix as it is produced by MSC obviously plays a crucial role for cancer metastasis and therefore might be a promising anti-cancer drug target.

Disclosure of interest

None declared.

O6

Single cell allele specific expression (ASE) IN T21 and common trisomies: a novel approach to understand DOWN syndrome and other aneuploidies

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Human Genomics 2016, 10(Suppl 1):O6

Objectives

Trisomy 21 is a model disorder of altered gene expression. We have previously used a pair of monozygotic twins discordant for trisomy 21 to study the global dysregulation of gene expression, without the noise due to genetic variation among individuals (Nature:508; 345–350;2014). The majority of previous studies focused on aneuploidies were conducted on cell populations or tissues. Our study focusing on gene and allelic expression behaviour of single cells (SC), aims to reveal biological insights regarding the cellular impact of aneuploidy and uncover the mechanisms of gene dosage.

Methods

We estimated the allele specific expression (ASE) from RNAseq of ~1000 single cells in different aneuploidies. We used 352 SC fibroblasts (173 Normal and 179 T21 cells) from the pair of monozygotic twins discordant for T21, 166 SC from a mosaic T21, 176 SC from a mosaic T18, 151 SC from a mosaic T8, and 146 SC from a mosaic T13.

Results

In the monozygotic twins, a considerable number of heterozygous sites at the non-chr21 genome showed monoallelic expression (MAE); (Normal: 73.5 % monoallelic in 564,668 observations, and T21: 78.7 % monoallelic in 549,799 observations). There was also considerable MAE for chr21 sites in Normal and, surprisingly, in T21 cells as well (Normal: 63.3 % monoallelic in 5,009 observations, and T21: 72.8 % monoallelic in 6,456 observations). We classified the genes on chr21 in 3 classes according to the level of the aggregate MAE of their corresponding sites (9 monoallelic, 29 intermediate, 2 biallelic). Similar results, i.e. extensive MAE on the supernumerary chromosome genes, were also observed in the other aneuploidies.

Conclusion

We hypothesize that for genes with monoallelic expression, the abnormal gene dosage induced by the aneuploid chromosome is maybe due to the number of cells expressing the gene. This difference in the fraction of expressing cells could contribute to the development and the variability of phenotypes in aneuploidies. This study provides a new fundamental understanding of the allele specific expression in T21 and other aneuploidies.

Disclosure of interest

None declared.

O7

Role of microRNA in LCL to iPSC reprogramming

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Human Genomics 2016, 10(Suppl 1):O7

Objectives

A large number of EBV immortalized lymphoblastoid cell lines (LCLs) have been generated and maintained in genetic/epidemiological studies as a perpetual source of DNA and as a surrogate in-vitro cell model. Recent successes in reprogramming LCLs into induced pluripotent stem cells (iPSCs) have paved the way to generate more relevant in-vitro disease models using this existing bio-resource. However the effects of EBV encoded oncoproteins on cellular transcription and function make LCLs a unique biomaterial to re-programme. Accumulating evidence now provides support that miRNAs play a critical role in transcription factor-induced reprogramming of iPSCs.

Methods

To investigate the role of miRNAs in regulating gene expression and cellular functions during LCL to iPSC reprogramming, we performed a parallel genome-wide miRNA and mRNA expression analysis in six LCLs and their reprogrammed iPSCs.

Results

A total of 77 miRNAs and 5,228 mRNAs were significantly (FC-abs ≥ 2.0 and FDR ≤ 0.05) differentially expressed (DE) during LCL to iPSC reprogramming out of which 29 miRNAs and 2,317 mRNAs were significantly down-regulated and 48 miRNAs and 2,911mRNAs were significantly up-regulated. The down-regulated miRNAs were highly enriched for LCL specific miRNAs (miR-155, let-7a-1, miR-21, miR-142, miR103, miR-320, miR-146a-b) and the up-regulated miRNAs were highly enriched for iPSC specific miRNAs (miR-302a, miR-302c, miR-371a, miR-302b, miR-302d, miR-372, miR-373mir-92a-1, miR-92a-2, miR-92b, miR-17, miR-20a, miR-18a). Further we performed target prediction analysis for all the significantly DE miRNAs using the miRNA target prediction data bases. The 3,456 genes were predicted to be targets of the 29 miRNAs that were significantly down-regulated during LCL to iPSC reprogramming. Out of these 3,456 predicted target genes 1,023 were significantly DE during LCL to iPSC reprogramming. For the 48 miRNAs that were significantly up-regulated during LCL to iPSC reprogramming 5,063 target genes were predicted out of which 1,462 were significantly DE during LCL to iPSC reprogramming.
The significantly DE genes that were also the predicted targets of the significantly down or up regulated miRNAs were further analyzed for functional annotations and pathway analysis using Ingenuity Pathway Analysis Platform.

**Conclusion**
In summary, our analysis identifies DE miRNAs and their DE target genes and a global role of miRNAs in broad resetting of cellular transcriptome and function during LCL to iPSC reprogramming.

**Disclosure of interest**
None declared.

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**O8 Multiple enhancer variants disrupt gene regulatory network in Hirschsprung disease**

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**Objectives**
Common sequence variation in cis-regulatory elements (CREs) are the suspected etiological causes of complex disorders. We examined all common (>10% minor allele frequency) non-coding variants within a ~153kb locus surrounding the gene for receptor tyrosine kinase Ret, which is most commonly mutated in Hirschsprung disease (HSCR) or congenital aganglionicism), a form of functional intestinal obstruction in neonates (1 in 5,000 live births). We hoped to find all causal non-coding polymorphisms disrupting enhancer function leading to the disease.

**Methods**
We used human and mouse fetal gut at relevant developmental time points for transcriptional profiling, ChiP assays, transgenic enhancer assays and siRNA mediated knockdowns of relevant transcription factors.

**Results**
We demonstrate that: (i) the three polymorphisms residing in 3 distinct enhancers that increase risk of the disease by 4-, 2- and 1.7-fold. Haplotypes for these three independent variants display wide variation in risk. (ii) the three CREs are Ret enhancers with distinct temporal activities during mouse gut development; (iii) the CREs are bound by the transcription factors Rab, Gata2/3 and Sox10, respectively, each developmentally expressed concordant with its cognate enhancer activity; (iv) variants in these CREs lead to their loss of activity and reduce Ret expression; (v) Ret is a positive feedback regulator of Sox10 and Gata2/3 transcription and; (vi) additional feedback interactions affect its ligand Gdnf, co-receptor Gfra1 and signal terminator Cbl.

**Conclusion**
These results explain how individually common, small effect non-coding polymorphisms can lead to large genetic effects in HSCR, since transcription attenuation of Ret from enhancer mutations are amplified through its auto-regulation. These results implicate RET as a key rate limiting step in early enteric nervous system (ENS) development and explains why >95% of HSCR cases have at least one RET loss-of-function allele. More generally, the phenotypic impact of a complex disorder can only be understood by assessing gene effects in the context of their gene regulatory networks.

**Disclosure of interest**
None declared.

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**O9 Metabolomic profiling for the diagnosis of neurometabolic disorders**

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**Objectives**
In individuals presenting with undifferentiated phenotypes such as developmental delay, hypotonia, and seizures, the list of differential diagnoses is often very long and includes metabolic/neurometabolic, genomic, and other Mendelian disorders. Here we want to demonstrate the utility of untargeted metabolomic profiling to screen for a wide range of neurometabolic disorders.

**Methods**
Untargeted small molecule metabolomic profiling was performed as described previously [1] on plasma samples from 12 patients suspected to have a neurometabolic disorder with a presentation of seizures, developmental delay and hypotonia.

**Results**
We identified 5 different neurometabolic disorders in these 12 patients. We observed elevations of 3-methoxytyrosine and decreased levels of dopamine and vanillylmandelate in AADC deficiency, elevations of 2-pyridylidione in ABAT deficiency, elevations of succinyladenosine in ADSL deficiency, increased citrate in citrate transporter deficiency, and elevations of imidazole propionic acid, cis and trans-urocanate in urocanase deficiency. The perturbations in the metabolomic profiles of plasma from these patients are unique, specific and not previously seen in over 300 other samples analyzed as normal controls or for other indications.

**Conclusion**
The standard diagnostic test for AADC, ABAT, and ADSL deficiency is CF neurotransmitter analysis, while testing for urocanase deficiency requires an enzyme activity assay from a liver biopsy. These cases demonstrate the ability of untargeted metabolomic profiling for the functional confirmation of pathogenicity of VUS found via WES; moreover, disorders for which there is no biochemical testing or where testing is only available on CSF are able to be diagnosed in a plasma sample. This also demonstrates the utility of metabolomic profiling alone to screen for a wide range of neurometabolic disorders.

**References**

**Disclosure of interest**
None declared.

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**O10 A novel causal methylation network approach to Alzheimer’s disease**

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Human Genomics 2016, 10(Suppl 1):O10

**Objectives**
A novel causal methylation network approach to Alzheimer’s disease.
Objectives
Alzheimer’s disease (AD) is the most common progressive neurodegenerative disease and represents a major cause of disability for elderly patients. DNA methylation--are increasingly seen as playing an important role in AD development. However its causal mechanisms remain unclear. Recent studies indicate that AD develops essentially as a result of dysfunction of molecular networks. Our purpose is to develop large-scale causal methylation networks to uncover the mechanism of AD development.

Methods
We propose to use causal graphs as a major concept and a general framework for causal methylation network analysis and develop “score and search”-based methods for exact learning causal graphs of methylation networks to find the best-scoring structures for a given methylation dataset. Specifically, we develop novel functional structural equations for modeling methylation networks and use integer programming to search the network with optimal score.

Results
The proposed methods were applied to AD data with 460045 CpG sites from 748 samples. At the first stage, the methylation data of 168 gene from the pathway Alzheimer’s disease were used to create a causal network describing the connection among the methylation sites between these genes. According to the current result, 148 gene was matched and tested in the model. We identified a largest connected causation methylation network with 47 nodes and 96 edges. Most genes were confirmed to play an important role in the AD development from the literature.

Conclusion
The proposed methods provide a highly flexible general framework for causal methylation network analysis and provide more rich information than co-methylation network. The exact learning algorithms will guarantee to find optimal solutions and hence provide accurate estimations of causal graphs of methylation networks. The causal methylation networks are able to uncover the mechanism of AD development.

Disclosure of interest
None declared.

O12
Transcriptome analysis identifies genes, enhancer RNAs and repetitive elements that are recurrently deregulated across multiple cancer types
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Human Genomics 2016, 10(Suppl 1):O12

Objectives
We aim to find genes that are frequently deregulated in cancer and thus can be useful as diagnostic markers for early detection, and potentially as therapeutic targets. We focus on biomarkers with pan-cancer potential that can be applicable to multiple cancer types.

Methods
We used the Cap Analysis of Gene Expression (CAGE) profiles of 225 cancer cell lines and 339 normal primary cells from FANTOMS project. CAGE is a S’ sequence tag technology that enables promoter-level expression analysis and can be used to estimate the activity of enhancers from bidirectional transcription of enhancer RNAs. As a complementary data set, we used RNA-seq data from 14 tumor types profiled by The Cancer Genome Atlas (TCGA). In both data sets (FANTOM5 and TCGA), we performed cancer vs. normal differential expression analysis in all cancer types.

Results
We identified a set of pan-cancer markers (of both coding and non-coding transcripts) that are recurrently perturbed in both the cancer cell lines (FANTOM5) and clinical tumors (TCGA). The FANTOM5 CAGE data provided novel insights into cancer transcriptome. We used the genomic location of the CAGE TSSs to show that promoters that overlap repetitive elements (especially SINE/Alu and LTR/ERV1 elements) are often upregulated in cancer. Specifically, a little known repeat family, REP522 (~1.8Kb in size, largely palindromic, unclassified interspersed repeat), was strongly enriched for the most cancer-activated promoters. Here we present previously
Elevated mutation and widespread loss of constraint at regulatory and architectural binding sites across 11 tumour types

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Human Genomics 2016, 10(Suppl 1):O13

Objectives
Disruption of gene regulation is thought to play major roles in carcinogenesis and tumour progression. Here, we characterize the mutational profiles of diverse transcription factor binding sites (TFBSs) across 1,574 completely sequenced cancer genomes encompassing 11 tumour types. We assess the relative rates and impact of mutation at the binding sites of 87 different transcription factors (TFs) by comparing the abundance and patterns of single base substitutions within putatively functional binding sites to matched control sites.

Methods
To detect putatively regulatory binding sites in the genome, we used a combination of computational prediction and experimental data. Position weight matrices for 118 transcription factor binding motifs were used to find TFBS motif matches in the genome. We intersected these motif matches with experimentally defined open chromatin regions to define putatively functional TFBSs. Motif matches not occurring within open chromatin were used as control, putatively non-functional sites. Comparisons between these functional and control sites underlie our methods, and we develop novel metrics to assess the relative rates and functional impact of cancer mutations at putatively functional regulatory sites.

Results
We observe a strong and significant excess of mutations at functional binding sites across TFs, and show that the substitutions that accumulate in cancers are often more disruptive than those that are tolerated as germline variants. Putatively functional CTCF binding sites suffer an exceptionally high mutational load in cancer relative to control sites, and those involved in the architecture of higher order chromatin structures are the most highly mutated. The mutational load at CTCF-binding sites appears to be dominantly determined by replication timing and the mutational signature of the genome. The tandem duplicator phenotype as a distinct genomic configuration in cancer

The tandem duplicator phenotype as a distinct genomic configuration in cancer

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Human Genomics 2016, 10(Suppl 1):O15

Objectives
Next generation sequencing studies have revealed genome-wide structural variation patterns in cancer, such as chromothripsis and chromoplexy that do not engage a single discernable driver mutation, and that currently have no clinical relevance. We aimed at a

Exome sequencing provides evidence of pathogenicity for genes implicated in colorectal cancer

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Human Genomics 2016, 10(Suppl 1):O14

Objectives
In the U.S., the lifetime risk of developing colorectal cancer (CRC) is 4.5%. ~5% of cases carry an identified pathogenic variant in known causal genes. In another 20%, CRC appears inherited but no known pathogenic variant has been detected. This is partly due to lack of evidence to classify variants as pathogenic when they occur in genes that are implicated (GWAS, linkage, or pathological information), but not proven, to be associated with CRC.

Methods
To find evidence of association for 1128 suspected CRC associated genes, we compared the number of rare (MAF < 0.005), potentially disruptive variants (PDV) (stop gain (SG), splice acceptor/donor change (SA, SD), and frameshift (FS)) found in 169 CRC cases and 3524 controls. Cases included individuals from the Clinical Sequencing Exploratory Research NEXT Medicine study (CSER, N=78), Women’s Health Initiative (WHI, N=76), and Northwest Institute of Genetic Medicine Family Polyposis Study (NWIGM, N=15). Controls were selected randomly with respect to CRC from the Exome Sequencing Project (ESP), and were not known to have any Lynch associated cancers or to carry a known pathogenic CRC variant.

Results
We found a significant association between case status and rare PDV carrier status: 25% of CRC cases carried a rare PDV compared to 4% of controls (p<2e-16, OR 8.1). 96 genes had 174 rare PDVs: 92 SGs, 24 SDs, 15 SAs and 43 FSS. Rare PDVs occurred in cases only for 35 genes, in controls only for 55 genes, and in both for 6 genes. Among cases, there were 11 SGs, 2 SDs, 6 SAs and 26 FSS. Among controls, there were 81 SGs, 22 SDs, 9 SAs and 17 FSS. Case specific genes had ≤2 rare PDVs: those with 2 were MLH2 and MSH1. Interestingly, ATM, ATR and BRCA2 contained PDVs in both cases and controls.

Conclusion
We show the power of aggregate information to find support for disease association in a subset of CRC implicated genes. Additionally, we provide evidence that ATM, ATR and BRCA2 may be associated with CRC in addition to the known association with breast cancer.

Disclosure of interest
None declared.
detailed molecular characterization of one of these genomic configurations, the tandem duplicator phenotype (TDP).

Methods
We combined whole genome sequencing (WGS) data from 277 human genomes representing 11 cancer types and devised a robust genomic metric able to identify cancers with a chromosome called tandem duplicator phenotype (TDP) characterized by frequent and distributed tandem duplications (TDs).

Results
Enriched only in triple negative breast, ovarian, endometrial, and liver cancers, TDP tumors jointly exhibit TP53-mutations, low expression of BRCA1, and increased expression of DNA replication genes pointing at re-replication in a defective checkpoint environment as a plausible causal mechanism. The resultant TDs in TDP augment global oncogene expression and disrupt tumor suppressor genes. Importantly the TDP strongly correlates with cisplatin sensitivity in both triple negative breast cancer cell lines and primary patient-derived xenografts.

Conclusion
We conclude that the TDP is a common cancer chromotype that coordinately alters oncogene/tumor suppressor expression with potential as a marker for chemotherapeutic response.

Disclosure of interest
None declared.

O16
Modeling genetic interactions associated with molecular subtypes of breast cancer
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Human Genomics 2016, 10(Suppl 1):O16

Objectives
The characterization of mRNA-expression subtypes in breast cancer facilitates genomic and genetic studies to identify biological processes that drive distinct molecular subtypes and elucidates the potential feasibility of subtype-specific drug targets. However, such therapies tend to have limited efficacy, often due to unpredicted compensation in the network of mutations. Polygenic models that account for multiple somatic mutations and their interactions can potentially improve target selection and provide a more detailed view of tumor genetic architecture.

Methods
We addressed this problem with a multi-trait genetic interaction analysis of copy-number variation and gene expression data from breast cancer samples in The Cancer Genome Atlas. Modules of co-expressed genes were derived and assessed for biological function and genetic association with mutations in oncogenes and tumor suppressors. Summary module phenotypes with pleiotropic associated loci were simultaneously analyzed to infer direct genetic effects as well as effects mediated by genetic interactions for each module.

Results
We observed widespread evidence of genetic redundancy, in which two mutations combine to yield a less than additive effect that is similar to either mutation in isolation. In addition, we also identified interacting mutations that combinatorially associate with distinct modules and subtypes in a non-additive manner. These somatic mutant combinations were often predictive of molecular subtypes when single mutations were not.

Conclusion
Accounting for interactions among somatic mutations in tumor samples reveals high genetic redundancy and complex regulatory hypotheses for breast cancer subtypes. Our work demonstrates how integrative genetic and genomic analysis can be used to generate more precise hypotheses for tumor genetics, which may be used to prioritize therapeutic targets for robust tumor suppression.

Disclosure of interest
None declared.

O17
Recurrent somatic mutation in the MYC associated factor X in brain tumors
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Objectives
Activation of MYC pathway has been shown in diverse cancers. MYC Associated Factor X (MAX) plays a key role in the MYC-MAX-MAD gene regulatory network; however, its direct involvement in cancer has not yet been reported. Here we report discovery of a novel recurrent somatic mutation in MAX in brain tumors and study its effects on the development and progression of tumors.

Methods
We found a mutation on Arg 51 residue to Glu in MAX gene in a patient with bilateral thalamic pediatric astrocytoma in which the primary tumors (left and right thalams) had nearly identical mutation profiles except for the presence of the MAX R51Q only in one.

We used this unique opportunity to study the effects of this mutation on the progression and development of brain tumors. We performed differential expression on these samples to find the pathways affected by this mutation. Using ChipSeq we studied changes in the chromatin conformation in genes regulated by this network. We used CD experiments, to study how this mutation affects the affinity between Max and other proteins in this family and with DNA.

Results
We screened our dataset and found 7 cases in 180 HGAs exome sequenced by our group (3.8%) with this mutation. We also identified 14 cases in published datasets.

We found that this mutation always appears later in tumor development in subclonal fashion and is accompanied by at least one driver such as H3 K27M. Our differential expression and ChipSeq experiments revealed lack of a global effect of this mutation but specific effects on groups of genes involved in some pathways such as apoptosis. We also demonstrate that this mutation has no effect on the binding efficacy between proteins in its regulatory network, but a Max R51Q homodimer binds less efficiently to nonspecific DNA than its wild type. It however, only affects the binding between Myc/Max heterodimer to DNA in E-boxes.

Conclusion
We identify MAX as a new cancer gene, particularly relevant to brain cancer.

Our results show the possible effects of MAX mutation in promoting tumor progression and development. It also suggests the effect of this mutation on the spread of the tumor. These findings shed new light on the mechanisms underlying cancer progression and the involvement of MYC signalling in development of brain tumors which, in turn, can point us towards new targets for therapeutic approaches.
Disclosure of interest
None declared.

O18
Predictive biomarkers to metastatic pancreatic cancer treatment
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Human Genomics 2016, 10(Suppl 1):O18

Objectives
In 2015, we demonstrated that a strong family history of BRCA-related tumors portends a better prognosis in metastatic pancreatic cancer patients. We now investigate if this holds true for more recent patients treated with standard-of-care FOLFIRINOX (FNX) or Gemcitabine/nab-paclitaxel (GA). We hypothesize that targeted sequencing of these tumors for DNA repair pathway aberrations will better predict outcomes than the surrogate marker of family history, which may be subject to patient bias.

Methods
We identified patients with de novo stage 4 pancreatic cancer initially treated at MD Anderson Cancer Center with first-line FNX or GA. We excluded patients with prior surgical resection (bypass was allowed) or radiation as initial therapy, and patients with unknown family history. Survival analysis was performed using the Kaplan-Meier method.

Results
We identified 153 patients initially treated with FNX and 80 patients treated with GA. Median age of the entire cohort was 62 years (36–84). 58% were male. Median OS was 286 and 295 days, respectively. Approximately 5% of patients had a family history of 3 or more BRCA tumors (breast, ovarian, prostate, pancreas). Median OS for these patients was 469 days, as compared to 285, 268, and 296 days for patients with 0, 1, and 2 affected family members. Median survival in patients with 0–3 affected family members was 456 days and 511 days for patients on FNX and GA respectively (95% CI 240–636d, 0–1092 d). For patients with 0–2 such family members, median OS was 283 and 268 days, respectively. As expected, ECOG 0–1 and the absence of liver metastases were associated with longer survival. We identified 126 of 153 FNX patients and 51 of 80 GA patients with pathology specimens available for targeted sequencing.

Conclusion
As in our earlier report, we see a trend towards increased survival in patients with 3 or more family members with BRCA associated tumors. However, the small number of these patients precludes a definitive assessment. We believe that targeted sequencing of DNA repair pathway and associated genes will better elucidate the mechanism of survival benefit over biased family history.

References

Disclosure of interest
None declared.

O19
DDIT4 gene expression as a prognostic marker in several malignant tumors
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Human Genomics 2016, 10(Suppl 1):O19

Objectives
The DDIT4 gene (DNA-damage-inducible transcript 4) encodes a protein related to adverse environmental conditions, whose action is the inhibition of mTOR. In a recent work we found DDIT4 levels was associated with the outcome in triple negative breast cancer (J Clin Oncol 33, 2015 (suppl; abstr 1097)). There are not previous reports relating DDIT4 with prognosis of cancer patients. Our aim in this study was to explore the influence of this gene in several types of malignant tumors.

Methods
We evaluated the influence of DDIT4 expression in the outcome (either, disease-free survival or progression-free survival or overall survival). Univariate Cox regression analysis of DDIT4 in the online platforms KM Plotter (http://kmplot.com/) and SurvExpress (http://bioinformatica.mty.itesm.mx/) was done. Datasets were split based in the median of expression. Cancer types evaluated include: Acute Myeloid Leukemia, Brain Cancer, Breast Cancer, Bone Cancer, Cervical Cancer, Head and Neck Cancer, Hematological Cancer, Liver Cancer, Lung Cancer, Pancreatic Cancer, Ovarian Cancer and Prostate Cancer.

Results
A high level of DDIT4 was significantly associated with a worse outcome in breast cancer (n=4142), P=3x10−14 (HR=1.64, CI95%: 1.44-1.87); acute myeloid leukemia (n=168), P=3.47x10−5 (HR=2.31, CI95%: 1.55-3.43), glioblastoma multiforme (n=538), P=0.005809 (HR=1.31, CI95%:1.08-1.59); ovarian cancer (n=1648), P=0.0096 (HR=1.2, CI95%: 1.04-1.37); head and neck squamous cell carcinoma (n=283), P=0.03347.
(HR=1.49, CI95%: 1.03-2.15) and lung adenocarcinoma (n=866), \( P=0.0038 \) (HR=1.4, CI95%: 1.02-1.9). In contrast, a high level of DDIT4 was associated with a better prognosis in gastric cancer (n=641), \( P=1.7 \times 10^{-6} \) (HR=0.62, CI95%:0.5-0.75) and lung squamous cell carcinoma (n=675), \( P=0.0015 \) (HR=0.42, CI95%:0.24-0.73). Frequency of structural alteration of DDIT4 found at cbioportal.org, indicate that 0 to 5.1% of primary tumors where DDIT4 was related with the prognostic had mutations, however in breast tumors xenografts, gene amplification occurs in 17.4% elucidating an key role in tumor aggressiveness.

**Conclusion**

DDIT4 is a promising molecular marker for outcome in several types of cancer. In addition elucidation of DDIT4 participation in cancer aggressiveness could lead to improve the therapeutic strategies, mainly those related with mTOR inhibition.

**Disclosure of interest**

None declared.

**Methods**

We utilized approximately 200 thousand somatic genomic alterations (deletions, inversions, duplications) and more than 34 million somatic mutations from 2573 high-coverage whole genome sequencing data across 45 different cancer studies with paired normal samples. We integrated mutations, gene expression and structural alterations with TAD boundaries that we have identified from 5 different human cell lines, representing three different germ layers.

**Results**

Our analysis revealed a strong correlation between the mutational landscape and the TAD organization of the genome. In addition, we found that TAD boundaries inflicted structural alterations that not only affected nearby gene regulation but also the distribution of mutations in human cancers.

**Conclusion**

Structural alterations affecting the spatial organization of the human genome, could lead to dysregulation of genes as well as aberrant mutation distributions in human cancers.

**Disclosure of interest**

None declared.

**O20**

**Spatial organization of the genome and genomic alterations in human cancers**

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**Human Genomics 2016, 10(Suppl 1):O20**

**Objectives**

The hierarchical folding of genomic DNA within the nucleus is closely related to transcriptional regulation. Recent chromosome conformation studies have suggested that mammalian chromosomes are structured into tissue-invariant topologically associating domains (TADs) where the DNA within a domain is interacting more frequently than with regions in other domains. Genes within the same TADs represent similar gene-expression, histone-modification profiles. Therefore regions separating different TADs (boundaries) have important roles in reinforcing the stability of these domain-wide organizations. TAD boundary disruptions in human limb malformations and cancer lead to dysregulation of certain genes, due to de novo promiscuous enhancer exposure to promoters. Here we sought to identify relationship between genomic architecture and genomic alterations in human cancers.

**Methods**

Spatial organization was quantified by analyzing the genomic landscapes of 5 human primary cell cultures and 67 xenografts using ChromHMM, assessing the consistency of TADs across these different samples. Consistency and the ability of these domain-wide organization to be used as predictive biomarkers for targeted therapies will be presented.

**Conclusion**

Structural alterations affecting the genomic landscape were found to be predictive of gene transcriptional dysregulation, gene amplification, and cancer aggressiveness. The consistency of TADs across different cell lines and xenografts suggests a robust strategy for targeted therapies.
Genomic analysis reveals novel drivers and progression pathways in skin basal cell carcinoma


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Human Genomics 2016, 10(Suppl 1):O22

Objectives
Basal cell carcinoma of the skin (BCC) is the most common malignant neoplasm in humans. BCC is primarily driven by aberrant activation of the Sonic Hedgehog (Hh) pathway. However, its extensive phenotypical variation remains to be explained.

Methods
The genetic profiling of 293 BCCs revealed the highest mutation rate in MYCN (30%), PPARα (15%), STK19 (10%), LAT51 (8%), ERBB2 (4%), PIK3CA (2%), RAC1 (1%) and N/K/H-RAS (2%) in PTEN (23%), RB1 (8%) and FBXW7 (5%). In line with the mutational profiles detected by DNA sequencing, we observed activation of the Hh pathway as well as upregulation of target genes of the Hippo-YAP pathway and activation of MYCN target genes in RNAseq experiments.

Results
85% of BCCs harbored mutations in Hh pathway genes: mutually exclusive PTCH1 (73%) and SMO (20%) (P=6.6x10^-19), SUFU (8%), and in TP53 (61%). 85% of BCCs also harbored additional driver mutations in other genes implicated in BCC tumorigenesis. Recurrent driver mutations were observed in MYCN (30%), PPARα (15%), STK19 (10%), LAT51 (8%), ERBB2 (4%), PIK3CA (2%), RAC1 (1%) and N/K/H-RAS (2%). Loss of function (LoF) and deleterious missense mutations were observed in PTEN (23%), RB1 (8%) and FBXW7 (5%). In line with the mutational profiles detected by DNA sequencing, we observed activation of the Hh pathway as well as upregulation of target genes of the Hippo-YAP pathway and activation of MYCN target genes in RNAseq experiments.

Conclusion
The functional analysis of the novel tumorigenic driver mutations in MYCN, PTEN, and LAT51 demonstrates their relevance in BCC tumorigenesis and provides an expanded molecular understanding of BCC.

Disclosure of interest
None declared.
O24 Clinical validity and actionability of multigene tests for hereditary cancers in a large multi-center study

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Clinical genetic testing is rapidly evolving with the introduction of next-generation sequencing (NGS) however questions remain about these new tests. First, can NGS methods be deployed that deliver equal or improved performance vs. traditional methods on the full spectrum of (often complex) disease causing variants? Second, do expanded NGS tests provide medical benefits which outweigh the increased uncertainty that naturally follows from testing more genes in more patients? In recently published work [1,2] we tested a large panel of cancer risk genes by NGS in a clinically representative population to evaluate these questions. Here we expand upon that work and arrays hybridized according to manufacturer’s protocol. Data was obtained for 33 of the 39 samples. Review showed that 3 data were unavailable.

Results

We find 100% concordance with traditional methods on both sequence and copy-number alterations using a battery of 3 calling algorithms and associated biochemistries, we find 99.8% concordance with BRCA1/2 classifications produced by a different laboratory that uses a large proprietary database. Finally, we find that 52% of genetic test results positive for genes other than BRCA1/2 would warrant consideration of a change in care for mutation-positive patients under current medical practice guidelines.

Conclusion

In appropriately referred patients, multi-gene panel testing yields clinically relevant findings with potential management impact for substantially more patients than does BRCA1/2 testing alone.

References

1. Lincoln, JMD 2015
2. Desmond, JAMA Oncol 2015

Disclosure of interest


O25 Correlation with tumor ploidy status is essential for correct determination of genome-wide copy number changes by SNP array

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Human Genomics 2016, 10(Suppl 1):O25

Objectives

Neuroblastoma, the most common extra-cranial solid tumor in the pediatric population, is a histologically and clinically heterogeneous neoplasm. Tumors characterized only by whole chromosome changes tend to act favorably, whereas tumors with MYCN gene amplification and/or segmental chromosomal aberrations have poor outcomes. Given the clinical relevance, our aim was to verify the accuracy of OncoScan FFPE SNP Array (Affymetrix) in assessing genome-wide copy number changes in tumor samples by comparison to other tumor ploidy testing methods.

Methods

39 neuroblastic tumors from 38 patients were analyzed using the OncoScan FFPE SNP Array at a pediatric hospital. Tumor slides were microdissected to increase tumor purity, genomic DNA was isolated and arrays hybridized according to manufacturer’s protocol. Data was analyzed using the OncoScan Console and results viewed using Chromosome Analysis Suite (ChAS) and OncoScan Nexus Express. Copy number results determined by SNP array were compared to ploidy results obtained by karyotype analysis and/or flow cytometry. Additionally, chromosome 1 and 2 aneuploidy data were used as a surrogate ploidy marker in cases where cytogenetics and flow cytometry were unavailable.

Results

Data were obtained for 33 of the 39 samples. Review showed that 3 of the 33 samples required recentering of the tumor ploidy baseline. Two samples from 2 lesions in one patient showed widely discordant copy number calls by the software, with one assigned a near tetraploid status and the other a near diploid status. Karyotype analysis confirmed a near tetraploid state. Correlation by karyotype, flow cytometry or chromosome 1 and 2 aneuploidy revealed an additional 4 discordant cases (total of 5/33 cases or 15%).
Conclusion
Despite complex algorithms used by the OncoScan software to assign copy number calls, in 15% of the analyzed cases a 2nd method was necessary to correctly assign tumor ploidy baseline. The complex chromosomal copy number changes present in tumors, in addition to tumor impurity, heterogeneity, and poor sample quality, can challenge the software's ability to correctly assign ploidy state. Our results demonstrate that a second independent method may be necessary in complex tumor cases to correctly assign ploidy that truly reflects tumor biology and that may be necessary for correct patient management.

Disclosure of interest
None declared.

O27
Mutation spectrum in a pulmonary arterial hypertension (PAH) cohort and identification of associated truncating mutations in TBX4
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Human Genomics 2016, 10(Suppl 1):O27

Objectives
To identify the genetic determinants of Pulmonary Arterial Hypertension (PAH) in a cohort of pediatric PAH patients.

Methods
We performed whole-exome sequencing (WES) in a cohort of 60 probands with PAH and family members when available (180 total individuals) without a molecular diagnosis after most of the series was screened for mutations in BMPR2. In addition, we performed WES in additional 118 singleton cases. We screened all samples for variants in known PAH associated genes and performed trio-based analysis to identify novel candidate PAH genes.

Results
We identified known and novel mutations in the known PAH genes. In addition we identified novel truncating variants in TBX4 occurring de novo or inherited from an asymptomatic parent in 5 patients and a de novo predicted deleterious nonsynonymous variant in one additional patient.

TBX4 is a transcription factor of the T-box gene family. It is expressed in a variety of tissues during early mouse development including the atrium of the heart, the limbs, and the mesenchyme of the lung and trachea. TBX4, jointly with TBX5, has been shown to interact with FGF10 during lung growth and branching. Mutations in TBX4 have been previously reported to cause small patella syndrome, an autosomal-dominant skeletal dysplasia characterized by patellar aplasia or hypoplasia. A study in 2012 [1] identified an association of TBX4 mutations with PAH in 6 patients.

Conclusion
We identified 6 different deleterious variants in TBX4 (2 inherited and 4 de novo) in our initial cohort of trios where patients were ascertained for primary pulmonary arterial hypertension, accounting for ~10% of the probands in this series. Subsequently we confirmed this association in additional 4 singleton cases, including a patient with a large intragenic microdeletion of TBX4. Our results confirm the role of TBX4 as an important cause of hereditary PAH, accounting for ~5% of our whole PAH cohort.

References

Disclosure of interest
Objectives
To identify mutations causing North Carolina macular dystrophy (NCMD, MCDR1).

Methods
We performed targeted Nex Gen sequencing of the MCDR1 region (870kb) in 8 affected individuals from 3 families representing 3 different haplotypes affected with chromosome 6 linked NCMD (MCDR1). In addition to our original 11 MCDR1 families recently published (141 total subjects), we now have an additional cohort of 23 families with the NCMD phenotype available for study (total of 367 subjects, 32 families).

Results
We initially found 14 rare variants spanning 870kb of the disease-causing allele. One of these variants (V1, ch6:1000400906) was absent from all published databases and all 261 controls, but was found in a total of 13 NCMD kindreds. This variant lies in a Dnase 1 hypersensitivity site (DHS) upstream of both the PRDM13 and CCNC genes. Sanger sequencing of 1 kb centered on V1 was performed in the remaining NCMD probands, and 2 additional novel single nucleotide variants (V2, ch6:10000987, in 6 families and V3, ch6:100041040 in 1 family) were identified in the DHS within 134 bp of the location of V1. A complete duplication of the PRDM13 gene was also discovered in a single family (V4). The 4 mutations V1 to V4 segregated perfectly in the 118 affected and 33 unaffected members of the 21 NCMD families.

Conclusion
We identified 4 rare mutations in a non-coding region, each capable of arresting human macular development by causing over expression of PRDM13. Additional families with the NCMD phenotype continue to support that these mutations are causative of MCDR1 / NCMD.

Disclosure of interest
None declared.

O29
PhenoDB and genematcher, solving unsolved whole exome sequencing data


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Human Genomics 2016, 10(Suppl 1):O29

Objectives
To identify the causative variant(s) and gene(s) of rare Mendelian phenotypes by the re-analysis of unsolved whole exome sequencing (WES) data.

Methods
To address some of these cases, we have incorporated maternal and paternal imprinting analysis and polygenic analysis to the PhenoDB Variant Analysis tool. We also analyzed WES data from 1063 samples for rare, functional variants in known imprinted genes, in the genes on pseudoautosomal regions, genes that escape X-inactivation, and genes on chromosome Y. To facilitate data sharing as well as improve the search for patients or model organisms with variants in specific candidate genes we have also been adding capabilities to GeneMatcher (www.genematcher.org). In GeneMatcher there is an option to match based upon OMIM® number, genomic location and, as of October 2015, on phenotypic features. As part of the Matchmaker Exchange (MME) (http://matchmakerexchange.org/), we have also developed an API that was implemented in August 2015 and allows the GeneMatcher users to submit their data to query PhenomeCentral and/or DECIPHER. Also, as part of the MME we have been working with other matchmaker databases on the API implementation to connect them to GeneMatcher and have been working on the version 2.0 of the API that will allow for more detailed queries.

Results
We found that the genes in the pseudoautosomal regions are not captured by the Agilent SureSelect v4 baits that we used to sequence these samples. The analysis of variants in the genes on chromosome Y identified 52 rare functional variants and the analysis of variants in the 242 imprinted genes identified 4,337 rare functional variants. These variants are being further evaluated to define causality. As of December 2015, 3,568 genes were submitted to GeneMatcher by 984 individuals from 48 countries and 1252 matches (60 matches with PhenomeCentral and 34 matches with DECIPHER).

Conclusion
The GeneMatcher approach has enabled collaborations and the description of novel Mendelian phenotypes and novel Mendelian genes like SPATAS, HNRNPK and TEL2. We expect that further use of GeneMatcher and other MME matchmaker databases will enable many new gene/phenotype connections and that the full impact this approach will be revealed in the published literature over the next years.

Disclosure of interest
None declared.

O30
Baylo-Johns Hopkins Center for Mendelian genomics: a four year review


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Human Genomics 2016, 10(Suppl 1):O30

Objectives
At its inception in 2011 the Baylo-Johns Hopkins Center for Mendelian Genomics (BHCMM), as one of three NIH funded Centers for Mendelian Genomics (CMGs), began its efforts towards: i) novel gene/mutation discovery, ii) elucidating the molecular bases of disease, iii) understanding the genetic susceptibility to disease traits, and iv) determining the genetic/genomic architecture of disease. Collectively
the CMGs have sequenced ~19,000 patient samples in collaboration with more than 500 investigators from 36 countries in the past four years. The BHCMG has learned many lessons from its contribution of 5,200 exomes ranging in 475 phenotypes and presented in 82 publications.

Methods
The Human Genome Sequencing Center (HGSC) at the Baylor College of Medicine has generated 46 TB of whole exome sequencing (WES) data using the HGSC-VCrome capture reagent and now includes ‘Spike-in PKv2’ to capture more difficult regions and additional gene targets. The HGSC-VCrome capture reagent, along with a multiplex strategy and use of full-length blocking oligos employed for hybridization, has yielded 7.7Gb of data per exome providing a coverage of 96% at 20X or greater. The Spike-in PKv2 reagent is comprised of 3,643 additional unique gene targets derived from Genetests, OMIM, selected cancer genes and Baylor Miraca Genetics Laboratory positive cases. The addition of this reagent converts >700-800 genes from partially covered to fully covered at >20X coverage.

Results
Success rates in this program have varied by phenotype and cohort collections ranging from 37% to 85%. The BHCMG has established valuable sample acquisition approaches and resources, enhanced sequencing methodology, curated a well-characterized phenotype-rich genetic database enabling genotype/phenotype relationships and encouraged collaborative efforts (i.e. GeneMatcher) to implicate 491 disease genes including 192 novel, 152 known and 147 phenotypic expansion genes.

Conclusion
Each discovery has shown the diagnostic capabilities in using WES and has taught lessons in disease mechanisms that continue to drive investigation into those cases that remain unsolved.

Disclosure of interest
None declared.

O31
Using read overlap assembly to accurately identify structural genetic differences in an Ashkenazi Jewish Trio
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Human Genomics 2016, 10(Suppl 1):O31

Objectives
Accurately identifying genetic differences between individuals or within samples taken from the same individual (tumor with normal control) is important to understanding the etiology of diseases, particularly for disease areas where large structural changes in the genome have been associated with the disease, such as neurological conditions, cardiovascular conditions and cancer. In clinical practice, identifying a previously uncharacterized de novo SV in an offspring that could be causing a condition is challenging with current methods that often have high false discovery rates.

Methods
Here, we present Biograph Anchored Assembly (BAA), an SV caller using whole read overlap assembly of reads that do match the reference exactly. In a previous study with next-generation sequencing of the reference individual HS1011 (English et al. (2015)), the method has been shown to have high sensitivity compared to other SV callers and a false discovery rate of less than 5%. The method is based upon the BioGraph data storage format (BAF). The BAF is a specialized index of the read overlap graph of a genome that can be queried up to one million times a second. Querying by both coordinate and sequence is particularly applicable to SV typing.

Results
Using BAA on a trio sequenced by the Personal Genome Project (PGP), the BAA variant caller detected a 3.4kb insertion inherited in the offspring that matched an alternate allele assembly now in GRCh38. The breakpoint and sequence of the insert were reported. The resolution of this inserted sequence allowed for five SNPs and an indel that were inherited from the father and a single SNP inherited from the mother to be distinguished in the offspring.

Conclusion
This level of resolution and accuracy in calling allows for structural variants, and even differences between structural variants, to be compared across individuals. This is important both for understanding the etiology of disease in larger studies as well as identifying de novo variants in an offspring in a clinical setting. Here, we further present results from 100 HiSeq X samples sequenced at 30x, including multiple classes of structural variants and multi-sample classification of shared breakpoints.

References

O32
Legal interoperability: a sine qua non for international data sharing
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Human Genomics 2016, 10(Suppl 1):O32

Objectives
The successful translation of genomic discovery into improvements in human health hinges on the effective and responsible sharing of genomic and health related data. The Global Alliance for Genomics and Health (GA4GH) brings together over 375 leading healthcare, research, disease advocacy, life science, and information technology organizations committed to establishing harmonized approaches to data sharing. The Regulatory and Ethics Working Group (REWG) of the GA4GH specifically promotes the legal interoperability of data.

Methods
The REWG establishes Task Teams with diverse, international membership across sectors to develop tools and solutions to facilitate data sharing. Existing Task Teams address Consent, Privacy & Security, Accountability, Paediatrics, Ageing and Dementia, Registered Access, Participant Values, and Machine Readable Consent.

Results
The REWG has developed a Framework for Responsible Sharing of Genomic and Health-Related Data, which aims to activate the human right of every citizen to benefit from the progress of science. This human rights basis gives the Framework political and legal dimensions that reach beyond the moral appeals of bioethics. Specific policies build on the Framework. The Consent policy provides guidance for balancing individual autonomy with the public good, and addresses sharing of legacy data where consent is silent on data sharing. The Privacy and Security policy emphasizes that safeguards should be proportionate to the risk of harm to patients. The new Accountability Policy provides best practices for transparency of data handling, as well as processes for monitoring and responding to non-compliance with data sharing standards.

Disclosure of interest
None declared.
Conclusion
In addition to these policies, ongoing initiatives within the REWG are tackling a host of related questions. What language is appropriate for consent to broad data sharing? Is there a middle ground between open and controlled access? Can conditions of data access be rendered machine readable, so as to enable automated data discovery? Can researchers be held accountable for not sharing data where patients have consented to broad sharing? And how do patients and the public feel about sharing their data for broad research purposes?

References

Disclosure of interest
None declared.

O33
High throughput screening platform of competent sineups: that can enhance translation activities of therapeutic target

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Human Genomics 2016, 10(Suppl 1):O33

Objectives
Short interspersed elements B2 (SINE B2) are broadly distributed transposable elements in whole mouse genome. Evolutionarily, SINE B2s share common ancestors with tRNAs. Among the various remarkable functions of recently discovered non-coding RNA functions, we discovered a new class of antisense non-coding RNAs (SINEUPs) that promote translation of partially overlapping sense coding mRNAs with no effects on RNA levels. In order to develop synthetic SINEUPs to up-regulate therapeutically interesting genes, we have determined that SINEUPs function requires two essential domains; one is SINE B2 element, also called the Effector Domain (ED), and the other is overlapping antisense RNA sequence, which provide specificity and is called the Binding Domain (BD). We have produced functional SINEUPs for a PD associate gene, PARK7 (DJ-1), as well as other genes. This synthetic SINEUP specifically enhances translation level of PARK7 mRNAs in human neuronal cell lines. Through a novel high-throughput screening (HTS) system, we aim at further optimizing the ED and BD of SINEUPs to produce very effective SINEUPs against any possible mammalian protein.

Methods
We report here our HTS, which is based on high-resolution automated fluorescent imaging of CeligoS instrument. We screened with the HTS several BDs for a hepatic transcription factor alpha (Hnf4-alpha), which is associated to maturity-onset diabetes of the young type 1.

Results
In addition, we validated that several SINEUPs targeting Hnf4-alpha are able to upregulate translation in mouse hepatoma cells and hepatocyte cells. We also validated that other EDs derived from natural SINE B2 sequences revealed target mRNA specific translation enhancement.

Conclusion
To conclude, synthetic SINEUPs are promising tools for gene/RNA therapy of haploinsufficiencies, and the HTS system is a powerful SINEUPs screening platform.

Disclosure of interest
None declared.

O34
The undiagnosed diseases network international (UDNI): clinical and laboratory research to meet patient needs

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Human Genomics 2016, 10(Suppl 1):O34

Objectives
Rare and undiagnosed disorders challenge patients, families, and clinicians. In 2008, NIH started an Undiagnosed Diseases Program (UDP), with the goals of providing answers to patients with mysterious conditions that eluded diagnosis and advancing medical knowledge about diseases. The UDP has expanded in the US, as the Undiagnosed Diseases Network, with 6 additional clinical sites, a coordinating center, 2 DNA sequencing cores, a model organisms screening center, a metabolomics core, and a biorepository. For further expansion, meetings were held in Rome and Budapest with clinician scientists from 7 nations.

Methods
The plan includes launching the UDNI (Australia, Canada, Hungary, Italy, Japan, Sweden, and the United States). The goals are to improve the level of diagnoses and care for such patients by common protocols, to facilitate research in disease etiology, and to create a collaborative research community. A comprehensive “–omics” approach would include exomic and genomic sequencing as well as metabolomics. The interim website is http://test.areasl.com/udni/home.

Results
To date, several principles are being implemented: Engaging centers of excellence, fostering a collaborative research environment, establishing a cooperative governance structure, designing a common research protocol, providing a uniform patient experience, collecting data by recognized standards, protecting patient data, observing ethical, legal, and social guidelines, devising broadly data sharing, stimulating dissemination of results, and ensuring a well-functioning network. A Board of Directors integrates working committees, including Patient Advocacy, Clinical Management, Sequencing, Databases, and Repositories.

Conclusion

Disclosure of interest
None declared.

O36
Performance of computational algorithms in pathogenicity predictions for activating variants in oncogenes versus loss of function mutations in tumor suppressor genes

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Human Genomics 2016, 10(Suppl 1):O36

Objectives
Several computational methods have been developed to predict whether amino acid substitutions result in disease. This type of analysis
is included in the ACMG/AMP guidelines for pathogenicity classification of variants and is being used by the Clinical Genomics (ClinGen) resource. These methods are generally blind to the underlying disease mechanism. Little is known about how mechanism of disease affects the predictive ability of these algorithms for variants implicated in inherited diseases. We address this by focusing on two classes of genes that differ in their molecular mechanism of action. Activating/gain-of-function mutations in oncogenes and loss-of-function mutations in tumor-suppressor genes (TSG) are pathogenic in cancer development. Moreover, unlike TSG, oncogenes are recurrently mutated at several amino acid positions.

**Methods**

We obtained 5078 missense variants in 29 oncogenes and 50 TSG, classified based on their pattern of mutations in COSMIC(1), from the ClinVar database and annotated them with 20 computational algorithms. These variants had clinical assertions provided by the submitting laboratory. We analyzed variants classified as either pathogenic or benign in oncogenes (n=321) and TSG (n=832).

**Results**

We found less concordance among the algorithms assessed for pathogenic variant prediction in either class of genes. Also the set of algorithms that were concordant in predicting benign and pathogenic variants differed on whether the variant was an oncogene or TSG. The concordant (e.g. GERP++) algorithms are primarily based on evolutionary conservation. A combination of GERP++ and the functional prediction algorithm FATHMM is more likely to produce discordant results for oncogenes. This implies that curators choosing different sets of computational algorithm are likely to result in different inferences for the same variants. We are developing statistical approaches to identify algorithms that produces maximal separation of benign and pathogenic variants for oncogenes and TSGs and applying them in a larger set of variants in the list of 56 genes recommended for reporting for incidental findings differing in their disease mechanism.

**Conclusion**

We find evidence that disease mechanism needs to be taken into consideration when deciding on algorithms for predicting pathogenicity. Our findings may aid in further classification of variants of uncertain significance.

**References**


**Disclosure of interest**

None declared.

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**O37**

**Identification and electronic health record incorporation of clinically actionable pharmacogenomic variants using prospective targeted sequencing**

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**Objectives**

The Baylor College of Medicine’s Human Genome Sequencing Center and the Mayo Clinic’s Center for Individualized Medicine are collaboratively working to sequence up to 10,000 patients from the Mayo Clinic Biobank. The objective of these efforts is to incorporate the results in patient electronic health records (EHRs) thus guiding clinical drug prescribing practices in terms of efficacy and avoidance of adverse events.

**Methods**

This study takes a prospective approach using a combination of re-agents including a targeted panel of seventy-six pharmacogenomically relevant genes developed as part of the Pharmacogenomic Research Network (PGRN). Specific targets are based primarily on a combination of community feedback and clinical guidelines published by the PGRN’s Clinical Pharmacogenetics Implementation Consortium (CPC). The targets include both gene coding regions and SNP targets aimed at characterizing both known and novel variants while keeping the costs equal to or below microarray-based genotyping approaches.

**Results**

Preliminary data was generated using 500 samples used previously as part of the Mayo Clinic’s eMERGE Network studies. Each institution developed both data generation and analysis pipelines geared toward identification of genomic variants and haplotypes influencing commonly prescribed drug efficacies and toxicities. Preliminary sequencing data quality was outstanding and demonstrated high variant correlation with the previous dataset. Improvements in haplotype calling and clinical decision support are ongoing. Moving forward, the Mayo Clinic’s Center for the Science of Health Care Delivery will be tracking outcomes to confirm the value of this approach.

**Conclusion**

In conclusion, this study will provide a large cohort blueprint for implementation of pharmacogenomics in precision individualized health care.

**Disclosure of interest**

None declared.

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**O38**

**Melanoma reprogramming state correlates with response to CTLA-4 blockade in metastatic melanoma**


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**Human Genomics 2016, 10(Suppl 1):O38**

**Objectives**

Targeting immune checkpoints has proven to be an effective strategy for the treatment of metastatic melanomas. However, less than half of patients respond to the immune checkpoint blockade. A complete understanding of molecular mechanisms underlying tumor response is lacking. In this study, we propose that the degree of melanoma cell “re-programming” may contribute to melanoma tumor resistance to immune therapy.

**Methods**

We employ RNA-sequencing (RNA-seq) and Reverse Phase Protein Array (RPPA) data from 68 early passage melanoma cell lines derived from tumor infiltrating lymphocyte harvests of 63 patients to identify biological processes and marker genes underlying the melanocyte “re-programming”. We propose a scoring system of the process based and employ it to study effects of the re-programming on outcomes of the immune therapy using a transcriptomics dataset (Van Allen et al, 2015) from pretreatment metastatic melanoma tumor samples from patients treated with ipilimumab (anti-CTLA-4).

**Results**

Melanoma cells grouped into 3 major concordant clusters by both RNA-seq and RPPA analysis (Fig. S). Examination of the genes underlying the clustering revealed profound differences in the expression of genes associated with melanocyte differentiation (including MITF) and with the Epithelial to Mesenchymal Transition (EMT) process. We determined the mean Z value for genes within each process, and designated the difference between the mean expressions as the “re-programming score” (RPS). Using the same set of marker genes for melanoma tumor samples we significantly separated responders from non-responders of the immune therapy and revealed 2 groups of non-responding tumors. Each group had a different subset of highly expressed EMT-associated genes, and opposite expression of the differentiation-associated genes. Combining a subset of genes that
are differentially expressed between responders and non-responders we markedly enhanced the prognostic value of the cytolytic score, a known prognostic feature.

Conclusion
The proposed scoring system of the melanocyte re-programming based on the RPS may hold prognostic value for immunotherapy treatments outcomes.

Disclosure of interest
None declared.

O39
Data-driven refinement of complex disease classification from integration of heterogeneous functional genomics data in GeneWeaver
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Objectives
Challenges in research, diagnosis and treatment of complex disease emerge from the poor alignment of the underlying biology of disease with a nosology defined by externally manifest signs and symptoms. Aggregate functional genomics data can enable the development of a data driven nosology in which disease characteristics, research models, diagnostic categories and drugs are more precisely aligned to specific, biologically based facets of disease.

Methods
GeneWeaver consists of a database and analysis tools for aggregation of heterogeneous functional genomics data across species, including curated pathways, ontology annotations, publication data, genetic mapping, transcriptome and proteome experiments, and other functional genomics data including user-submitted experimental results. Each is described with meta-content, enabling retrieval by disease related terms. Gene identifiers are harmonized to enable aggregation of data through homologous genes and gene products. To evaluate specificity of disease descriptors, we analyzed intersections among gene sets associated with disease-related terms. These gene sets were derived from studies of nine different organisms and consisted of genome-wide experiments, curated annotations to ontology terms and genes associated to disease-related terms through transitive association of genes, publications and MeSH terms.

Results
Analyses of term-to-term associations reveals that genes associated to co-occurring or difficult to discern diseases exhibit weak overlap across many different terms, especially those associated with psychiatric disorders which display extensive cross-disorder overlap. In contrast, more well-bounded disorders, such as degenerative ocular disorders, reveal strong and specific overlap and good matching of empirically derived data sets and annotations. By enumerating all intersecting associations of genes to disorders, we are simultaneously able to identify genes that differentiate among overlapping disorders, potentially defining the specific and unique aspects of these conditions for precise differentiation of disease.

Disclosure of interest
None declared.

O40
A general statistic framework for genome-based disease risk prediction
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Objectives
How to efficiently extract biomarkers for risk prediction and treatment selection from millions or dozens of millions of genomic variants raises a great challenge. Traditional paradigms for identifying variants of clinical validity are to test association of the variants. However, significantly associated genetic variants may or may not be efficient for diagnosis and prognosis of diseases. Alternative to association studies for finding genetic variants of predictive utility is to systematically search variants that contain sufficient information for phenotype prediction.

Methods
To achieve the goal, we introduce concepts of sufficient dimension reduction (SDR) which project the original high dimensional data to very low dimensional space while preserving all information on response phenotypes. We then formulate a clinically significant genetic variant discovery problem into the sparse SDR and optimal scoring problem and develop algorithms that can select significant genetic variants from high dimensional data. To speed up computation, we apply the alternating direction method of multipliers to solving the sparse optimal scoring problem which can easily be implemented in parallel.

Results
To illustrate its application, the proposed method is applied to a coronary artery disease (CAD) dataset from the Wellcome Trust Case Control Consortium (WTCCC) study, Rheumatoid Arthritis (RA) dataset from the GWAS of North American Rheumatoid Arthritis Consortium (NARAC) and the early-onset myocardial infarction (EOMI) exome sequence datasets which have European origin from the NHLBI’s Exome Sequencing Project. To evaluate the performance of the SDR for disease risk prediction, we present Table 1 that lists AUC of our SDR and other 10 existing methods. Table 1 clearly demonstrated that the proposed SDR method has much larger AUC than other 10 existing methods.

Conclusion
We shift the paradigm of feature selection from P-value and risk score ranking to optimal genome-wide searching. The SDR-based optimal genome-wide searching methods substantially outperform other existing methods for disease risk prediction. Our results strongly demonstrate that the rich genetic variation information provides powerful resources for disease risk prediction.

Disclosure of interest
None declared.
Integrative large-scale causal network analysis of imaging and genomic data and its application in schizophrenia studies

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Human Genomics 2016, 10(Suppl 1):O41

Objectives

Next generation genomic and image technologies produce a deluge of DNA sequencing, transcriptomes, metabolic, image, physiological phenotypes with millions of features. Analysis of increasingly larger and more complex data gives scientists access to vast amounts of information that was previously unavailable, but also poses great methodological and computational challenges. This talk provides perspectives for paradigm changes in current public health data analysis.

Methods

We develop novel statistical methods for paradigm changes in big genomic, epigenomic and imaging data analysis from low dimensional data to high dimensional data analysis. We develop novel functional structural equations with integer programming as a new framework for inferring large-scale causal networks of genomic-images and detecting pleiotropic effects of genetic variants on imaging. In addition, we develop new causal machine learning methods for network classification and combine images and genomic data for disease risk prediction.

Results

The proposed method for large-scale genomic-imaging causal network analysis was applied to the MIND clinical imaging consortium’s schizophrenia image-genetic study with 142 diffusion tensor images (DTI) with 538265 voxels and 14,142 genes in 64 schizophrenia patients and 78 controls. Each DTI were segmented into 41 regions. The causal image-genotype networks were constructed for all the individuals. In cases, the image network consisted of 41 nodes and 68 edges, and in controls, the image network consisted of 41 nodes and 65 edges. We identified 1,035 and 1,618 genes that were significantly connected to the image regions respectively. 27 genes in cases and 40 genes in controls were in the 108 schizophrenia associated genetic loci. The developed network classification algorithm was also applied to predict schizophrenia. Using cross validation, we can achieve 100% prediction accuracy in the training data and the average prediction accuracy, sensitivity and specificity in the test data were 95.1%, 96.2% and 93.9% respectively.

Conclusion

We shift the paradigm of big genomic and imaging data analysis from association studies to causal inference and provide powerful tools for unravelling causal chain of mechanisms of psychiatric disorders, delivering new therapeutic targets and biomarkers for precision medicine.

Disclosure of interest

None declared.

Cpipe: a convergent clinical exome pipeline specialised for targeted sequencing

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Human Genomics 2016, 10(Suppl 1):O43

Objectives

Efforts to move high throughput sequencing into the clinic must confront many challenges including meeting clinical standards for cost, reproducibility, quality, ethical and privacy considerations. The Melbourne Genomics Health Alliance was formed from a diverse group of institutions with the aim of sharing the burden of these challenges through a common sequencing and bioinformatics platform. In this work, we present a shared, open source analysis pipeline that was developed to meet these needs.

Methods

To enable a single solution for many different diseases and laboratories, we employed the Bpipe pipeline platform which allows for analysis stages to be easily added, substituted, replaced or customized on a per-sample or per-disease basis. Bpipe also offers powerful...
features for parallelization, so that pipelines can easily run on different distributed clusters or on dedicated computing resources. To ensure that the solution can be freely shared, it is released under the GPLv3 open source license.

Results
We have developed Cpipe, a shared, open source sequencing pipeline designed for clinical users. Cpipe includes support for the full range of features required in real world clinical sequencing applications. It allows for customized sets of targeted regions, prioritized genes, screening of incidental findings, automatic exclusion of sequencing artefacts and population variants, PDF provenance and quality reports. Cpipe produces a clinically interpretable report available in Excel format. Results can be optionally imported into an LOVD (Leiden Open Variant Database) instance for curation. Cpipe is available at http://cpipe.org.

Conclusion
Cpipe offers an open source solution that any clinical laboratory can employ, either as their main analysis pipeline or as a benchmark for comparison to improve their results.

References

Disclosure of interest
None declared.

O44
A Bayesian classification of biomedical images using feature extraction from deep neural networks implemented on lung cancer data
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Objectives
This project aims the formulated algorithm will be implemented explicitly on lung cancer pathological images. Specifically, this project has two goals. First, it aims to apply the concept of deep neural networks to supervised learning in the classification of images, with the understanding that modifying existing machine learning methods to target specific image sets can optimize the precision and accuracy of the analysis. The algorithm that will be formulated will be able to sort a given set of data (in this case, images) into desired sets with given qualifications. Second, it aims to apply the concept of deep neural networks, supplemented by Bayesian networks to pattern analysis of lung cancer data sets. Performing Bayesian network procedures on given lung cancer data can help us determine parameter values that will characterize those data. In turn, those parameters can be used to infer whether or not new data can be classified with the given training data or otherwise.

Methods
First, we take tiles of size 512x512 from images published on the cancer genome atlas database (http://cancergenome.nih.gov). Several samples from those 512x512 will be segmented further into 32x32 subregions, which served as training data for the algorithm. The main output will be an algorithm embedded in a web application wherein the user can just input newly acquired images, and the program can provide classify which 32x32 subregions depict cancer cells. A demonstration of the application can be shown during the presentation.

Conclusion
An executable program, whose input is a lung cancer image, will be segmented into 32x32 subregions, each of whose features will be extracted using deep neural networks and will undergo a Bayes’ classification scheme, and determine whether the subregion is normal or abnormal, either because of a broken cell, a stained cell, or an actual cancer cell.

Disclosure of interest
V. A. Antonio, Grant/Research Support from: CICP Research Project (https://cicp.naist.jp/ja/node/85), N. Ono: None declared.

O45
MAV-seq: an interactive platform for the Management, Analysis, and Visualization of sequence data
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Objectives
The increasing amount of heterogeneous genomic datasets generated today necessitates a robust platform for efficient data management and analysis. To address this need, we developed a software for the Management, Analysis, and Visualization of sequence data (MAV-seq), capable of addressing the issues related to the exponential growth of genomic applications and their datasets of enormous size and diversity. Our software also integrates various genomic preprocessing pipelines with user-friendly graphical interface to enable biologist with no programming experience conduct complex data analyses.

Methods
MAV-seq (Figure) is a desktop application developed to integrate bioinformatics methods, software engineering principles, human computer interaction guidelines and big data analytics. The graphical user interface (GUI) and back end development of MAV-seq is performed in Java and Python, and schemas for backend data storage are implemented in MySQL and MariaDB relational database management systems. MAV-seq allows direct data manipulation using GUI as well as data import and export in “csv” file formats.

Results: We developed MAV-seq as an interactive, user friendly, cross platform, encrypted and multi-roles based system for the management of sample repertoires and automation of the data preprocessing of epigenomic and transcriptomic data. It supports users in performing downstream data analysis by integrating several analysis pipelines for diverse data sets including ATAC-seq, mRNA-seq, tRNA-seq, Chip-seq, WES, WGS. MAV-seq can be customized for increasingly large scaled and complex datasets of different types. Moreover, it can directly interact with multiple data clusters to locate, input and process genomics data by automatically generating and running multiple-sequential and parallel pipelines.

Conclusion
MAV-seq is a comprehensive data management and analyses platform that is newly designed, developed, tested, validated and deployed at The Jackson Laboratory for Genomic Medicine. With this platform, we aim to advance genome-wide big data management, standardization and automation, which will expedite the pace and improve the levels of efficiency in loading, handling, tracking, securing, sharing, processing, analyzing and visualizing data.

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Disclosure of interest
None declared.

O47
Allele specific enhancer in EPAS1 intronic regions may contribute to high altitude adaptation of Tibetans
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Human Genomics 2016, 10(Suppl 1):O47

Objectives
Tibetans have shown their physiological phenotypes adaptive to hypoxia environment, including relative low hemoglobin concentration and higher oxygen saturation. Among tens of candidate genes revealed by various studies that relate to living in high altitude, EPAS1 and EGLN1 showed strongest selection signals as well as solid correlation to lower hemoglobin concentration in Tibetans. However, several tens of Tibetan specific SNPs in EPAS1 were all found in introns and the mechanism of this important gene contributing to genetic adaptation in Qinghai-Tibet Plateau is still elusive. This study aims at the function of Tibetan specific SNPs in EPAS1.

Methods
1) 82 candidate genes were designed for capture and sequencing in 45 Tibetans samples to discover more SNPs for further association and functional studies. 2) Statistical tests including Chi-square tests, haplotype association, and integrative haplotype score, and Refseq and NONCODE were used for annotation. 3) Cell culture of HepG2 and HEK293 under normoxia and hypoxia (1%) for all functional assays. 4) Enhancer validation in a dual-luciferase reporter system. 5) Circular chromosome conformation capture (4C) in HepG2 followed by sequencing in Hi-Seq 2000.

Results
By re-sequencing 82 genes selected from previous studies and our recent discoveries, totally 9947 SNPs were revealed in 45 Tibetan individuals (15 from Bomi; 20 from Anduo; 10 from Dingri). Six genes were consistently identified by three statistical tests. In addition to EPAS1 and EGLN1, studies on quantitative traits identified DISC1 and EDAR in association with diameter of Main Pulmonary Artery and SaO2, respectively. By integrating enhancer annotation data from FANTOM5 and ENCODE, 12 EPAS1 intronic regions were resulted. Further validation for the enhancer function demonstrated that three of them are true enhancers. Moreover, enhancer 1 showed an allele specific activity in both HepG2 and HEK293 cell lines. Especially, two Tibetan-Han divergent SNPs showed the addictive effect in enhancer activity. Further 4C-sequencing results of Enhancer 1 demonstrated the similar chromosomal conformation structures in most enhanced regions under hypoxia and normoxia treatments but a few interactions were shown to be hypoxia sensitive. Annotation with Refseq and NONCODE database revealed that targets potentially regulated by Enhancer 1 were mostly non-coding RNAs.

Conclusion
An allele-specific enhancer in EPAS1 intronic region may contribute to high altitude adaptation in Tibetans.

Disclosure of interest
None declared.

O48
Nanochannel based next-generation mapping for structural variation detection and comparison in trios and populations
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Human Genomics 2016, 10(Suppl 1):O48

Objectives
Genome structural variations (SV) have been well established to be associated with diseases and traits; however, SV analysis of human genomes has been severely limited to date by technical shortcomings. Traditionally, SVs have been detected by microarray (limited to imbalanced copy number variation (CNV) with a short dynamic range, low resolution, and relative readouts), next-generation sequencing (NGS) (primarily CNV, some balanced events but too short to span most repeats) and karyotyping and fluorescence in situ hybridization (FISH) (both are very low resolution).

Methods
Using a single-molecule genome analysis system, BioNano Genomics Irys® System, utilizing next-generation mapping (NGM) technology, it is now possible to comprehensively analyze whole genomes for SVs > 2 kilobase pairs (kbp), including balanced events in a cost-effective and high-throughput manner. This technology allows for the comparison of family pedigrees and populations, which is needed to potentially uncover genomic structural causes of Mendelian and complex diseases.

Results
We demonstrate the robustness of NGM for genome-wide discovery of structural variation in the CEPH trio set from the 1000 Genomes Project where the individuals were sequenced and analyzed in-depth. We generated de novo assemblies that covered at least 96% of the hg38 reference assembly. Compared to tens of large SV events detected by NGS, we uncovered hundreds of insertions, deletions, and inversions greater than 5 kbp, a large portion of which was novel, and some are located in the regions likely leading to disruption of gene function or regulation. Based on the pedigree structure, we estimated that the Mendelian concordance rate was 96%. We have also begun analysis of a trio of Ashkenazi Jewish descent from the NIST GIAB project, where we have found hundreds of inversions, insertions, and deletions, including large deletions in the UGT2B17 gene (involved in graft versus host disease, osteopathic health, and testosterone and estradiol levels) in the mother and son.

Conclusion
We show that NGM is a robust and effective method for structural variation detection in the human genome. Systematic whole genome structural variation within disease population cohorts is needed, in addition to the conventional SNP analysis, to study the effects of a full spectrum of genomic variations in human disease and complex traits.

Disclosure of interest
None declared.
Archaic introgression in indigenous populations of Malaysia revealed by whole genome sequencing

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Human Genomics 2016, 10(Suppl 1):O50

Objectives
The investigations of the indigenous populations from Malaysia and their migration history have been scarce. In this study, we performed the whole genome sequence for Malaysian indigenous (MI) genomes from Peninsular Malaysia (PM) and Northern Borneo (NB). The genomic structures of these samples were assessed, followed by identification of the archaic genomes introgression between these samples with Denisovan and Neandertal.

Methods
Seventeen genomes from indigenous populations from PM and NB were sequenced using Illumina Hi-Seq, thus unveiled the full spectrum of genetic architecture of the MI. Population genetic structure of these samples were assessed with PCA and ADMIXTURE. Inference of coalescent time and effective population size were performed using PSMC, and gene flow was estimated using TreeMix. Archaic genomes introgression was estimated using the D-stat and f-test. S-t analysis was applied to identify the introgressed genome segments.

Results
The divergence between Negrito and Austronesian occurred ~20K years ago, and were gradually replaced by the Austronesian expansion. Events of multiple gene flow into PM and NB was observed, in line with previous investigations. However no evidence of significantly higher gene flow from archaic genomes to MIs was observed, and that the archaic DNA segments found in MIs were different from those carried by the East Asia and Europeans.

Conclusion
Our analyses further strengthens the findings of the population structure of the indigenous people revealed by various earlier studies using SNP array, yet suggests that the history of these populations are far more complex than expected. The archaic genome introgression provided evidence of no significantly higher archaic genome component in our samples. This preliminary study complements the gaps of various speculations about archaic genomes introgression in the regions of SEA and Oceanic.

Disclosure of interest
None declared.

Breast and ovarian cancer prevention: is it time for population-based mutation screening of high risk genes?

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Human Genomics 2016, 10(Suppl 1):O50

Objectives
Germline mutations in BRCA1 and BRCA2 confer high lifetime risk of breast and ovarian cancer but importantly these risks are not irreversible. Identification of asymptomatic carriers could significantly reduce the incidence of these diseases. As a first step toward population based BRCA gene screening, we are sequencing the entire coding region of 20 known and proposed HBOC genes in 4,000 cancer-free Australian women.

Methods
Cancer-free women were selected from the LifePool study (www.life pool.org) which is a cohort of women attending the Australian population, based mammographic screening program. All exons of the target genes were enriched using the Haloplex system (Agilent) and sequenced on a HiSeq2500 instrument (Illumina). The data were filtered for known pathogenic or novel loss of function mutations.

Results
To date, data from 1,997 women has identified 17 with actionable mutations in BRCA1 (4 mutations), BRCA2 (9 mutations) or PALB2 (4 mutations). All 17 women subsequently accepted an invitation to attend a Familial Cancer Centre and then proceeded to formal clinical genetic testing. In addition 4 women had pathogenic mutations in BRIP1.

Conclusion
Our unique pilot data directly demonstrates a population carrier frequency of ~1% for pathogenic mutations in these recognized high risk breast and/or ovarian cancer genes and that such testing is well accepted by the screened population.

Disclosure of interest
None declared.
method enables single base resolution of methylation status through-out the genome, even from limiting DNA input quantities.

Conclusion
We have demonstrated the utility of increasing the efficiency of library preparation as a means of improving sequencing results obtained through both WGS and WGBS. This innovative technology enables sequencing of sample types that have been previously unavailable due to input or quality limitations.

Disclosure of interest

O54 Methods for large scale construction of robust PCR-free libraries for sequencing on Illumina HiSeqX platform
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Human Genomics 2016, 10(Suppl 1):O54

Objectives
With drastic drop in the cost of whole genome sequencing (WGS) on Illumina’s HiSeqX systems, it’s now economically feasible to plan large scale sequencing projects in research as well as clinical settings. However, successful execution of such projects will require design of robust sample preparation (library) workflows that can work with a spectrum of DNA quality and quantities. This is especially true for the highly desirable PCR-Free libraries that are known to provide improved gene and genome representation compared to PCR amplified libraries. Here we discuss the use of multiple PCR-Free protocols for use on HiSeqX.

Methods
Previously we automated Illumina TruSeq PCR-Free protocol, which is recommended to use with 1 ug of good quality DNA to prepared size selected libraries. However, to broaden the scope of samples that can be used for preparing PCR-Free libraries, we evaluated two additional library construction methods 1. Swift Biosciences 25 library kit and 2. Kapa Biosystems, hyper prep kit. PCR-Free libraries were prepared using 200 ng - 1 ug DNA of HGSC internal human control sample (HS1011) with these kits and sequenced on HiSeqX to generate 34-38X genome coverage data.

Results
Exome representation as measured by complete coverage of Online Mendelian Inheritance in Man (OMIM) genes at 20X read depth was lowest for the TruSeq PCR-Free libraries (2687 genes) when compared to the 25 libraries (2750 – 3020 genes) and the Kapa hyper libraries (2800 genes). GC representation was better in PCR-Free libraries when compared to the TruSeq Nano libraries. Kapa Hyper protocol was optimized on Beckman Coulter’s Biomek FXP liquid handler using 500 ng DNA and can prepare 96 libraries in ~ 6 hours. This protocol also works well with DNA of different integrities. Enhancements have also been made for precise quantification of PCR-Free libraries by qPCR and to eliminate unused adapter molecules in libraries that can impact sequencing.

Conclusion
Availability of such robust and automated protocols has positioned us to efficiently work with large sample sets to fully exploit the use of HiSeqX platforms for population level genomic studies and to drive its use in clinical setting.

Disclosure of interest
None declared.

O55 Rapid capture methods for clinical sequencing
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Human Genomics 2016, 10(Suppl 1):O55

Objectives
Advancement of next generation sequencing in clinical settings has required methods for rapid, robust delivery of high-quality sequencing data. Effective and timely diagnoses or prediction of risk of genetic diseases are important for medical intervention.

Methods
We have developed a ‘lightning capture’ process to deliver variant calls in 5–7 days after sample intake. This process includes: quick enrichment library preparation (5–6 hours), capture enrichment (about 8 hours), rapid sequencing (Illumina HiSeq2500) and data analysis via the HGSC-developed Mercury pipeline. The lightning capture protocol has been deployed in BMGL clinic lab for whole exome sequencing (WES) and recent carrier screening with a novel 500kb carrier mutation gene capture panel. Our WES design contains 3643 clinically relevant genes primarily from GeneTests and OMIM, and the carrier panel includes 168 complete genes that contain at least 850 known common genetic variants of clinical relevance. We also employ the genotyping with SNPTrace panel by Fluidigm in order to ensure reliable sample identification, and to test for sample cross-contamination.

Results
This method has been validated with ~500 WES and more than 5000 carrier samples. WES samples are processed in single capture or 3-plex co-capture, while carrier samples are in a cost-effective 47plex co-capture format for hybridization followed by sequencing of 94 samples (2 capture pools) per HiSeq 2500 lane. High enrichment efficiency was observed (72-80% reads on target and buffer) and superior coverage metrics across the design with 11 Gb sequencing yield for WES and ~400 Mb for Carrier samples. A detailed analysis of the carrier design performance using 140 de-identified samples found that known carrier mutations were correctly identified with high confidence (98.5%), including large/complex indel mutations.

Conclusion
Sample turnaround time for tests is often considered one of the most significant measures of performance for a clinical lab. The lightning capture process we developed enables data delivery in 5–7 days without sacrifice of data quality. This novel method may impact applications in prenatal, neonatal intensive care and other critical settings for clinical and research samples.

Disclosure of interest
None declared.

O56 A diploid personal human genome model for better genomes from diverse sequence data
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Human Genomics 2016, 10(Suppl 1):O56

Objectives
With drastic drop in the cost of whole genome sequencing (WGS) on Illumina’s HiSeqX systems, it’s now economically feasible to plan large scale sequencing projects in research as well as clinical settings. However, successful execution of such projects will require design of robust sample preparation (library) workflows that can work with a spectrum of DNA quality and quantities. This is especially true for the highly desirable PCR-Free libraries that are known to provide improved gene and genome representation compared to PCR amplified libraries. Here we discuss the use of multiple PCR-Free protocols for use on HiSeqX.

Methods
Previously we automated Illumina TruSeq PCR-Free protocol, which is recommended to use with 1 ug of good quality DNA to prepared size selected libraries. However, to broaden the scope of samples that can be used for preparing PCR-Free libraries, we evaluated two additional library construction methods 1. Swift Biosciences 25 library kit and 2. Kapa Biosystems, hyper prep kit. PCR-Free libraries were prepared using 200 ng - 1 ug DNA of HGSC internal human control sample (HS1011) with these kits and sequenced on HiSeqX to generate 34-38X genome coverage data.

Results
Exome representation as measured by complete coverage of Online Mendelian Inheritance in Man (OMIM) genes at 20X read depth was lowest for the TruSeq PCR-Free libraries (2687 genes) when compared to the 25 libraries (2750 – 3020 genes) and the Kapa hyper libraries (2800 genes). GC representation was better in PCR-Free libraries when compared to the TruSeq Nano libraries. Kapa Hyper protocol was optimized on Beckman Coulter’s Biomek FXP liquid handler using 500 ng DNA and can prepare 96 libraries in ~ 6 hours. This protocol also works well with DNA of different integrities. Enhancements have also been made for precise quantification of PCR-Free libraries by qPCR and to eliminate unused adapter molecules in libraries that can impact sequencing.

Conclusion
Availability of such robust and automated protocols has positioned us to efficiently work with large sample sets to fully exploit the use of HiSeqX platforms for population level genomic studies and to drive its use in clinical setting.

Disclosure of interest
None declared.
Objectives

Generate a high quality reference genome from a single individual to avoid limitations of the haploid mosaic human reference and explore the contribution of different data types to understanding the comprehensive clinical genome.

Methods

Illumina data from a variety of libraries (180 bp, 300 bp and 500 bp paired end data; 3kb, 6.5 kb and 8 kb mate-pair data), as well as Illumina Hi-C data, 20x PacBio RS long read data and BioNano optical mapping data were produced and assembled de novo. Structural variants that are difficult to characterize with exome sequencing or short sequence reads from small fragments were identified using two methods. We identified putative novel insertions in reads that did not map to the GRCh38 reference with calls supported by 6.5 kb mate-pair data, and confirmed in the de novo WGS assembly contigs and 3 kb mate-pair data. We identified tandem duplications with combined signatures of inverted 300 bp to 500 bp read pairs identifying the boundaries and larger read pairs confirming the size of the duplicated region.

Results

We report here the assembly of data from a single individual (HS1011). The data are available at NCBI under BioProject 203659. The released assembly is highly contiguous with a 394 kb Contig N50 and 148 Mb Scaffold N50 with full chromosome scaffolds. Data from the parents of HS1011 and long-read data allow us to phase variants within this genome. Putative novel insertions (78) and tandem duplications (70) were identified.

Conclusion

The assembly and underlying data reported here allow us to optimize methods to combine these data types and explore the utility of the different data types to identify structural variation and define which heterozygous variants are located on the same haplotype for haplotype-aware downstream analyses.

Disclosure of interest

None declared.

OS57

Development of PacBio long range capture for detection of pathogenic structural variants

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Human Genomics 2016, 10(Suppl 1):OS57

Objectives

Discoveries of chromosomal alterations ranging from single nucleotide variants (SNVs) to large structural variants (SVs), via next-generation sequencing (NGS) techniques, have drastically impacted the delineation of genomic architecture and disease mechanisms. However, in contrast to being able to identify SNVs efficiently and accurately, current short-read NGS platforms have been inefficient for discovery of complex chromosomal rearrangements. To detect chromosomal pathogenic SVs, we have developed a novel method, Pacific Biosystems large insert targeted capture-sequencing (PB-LITS). We further tested PB-LITS protocol for detection of clinical relevant complex gene structural mutations with a set of 6 breast cancer samples.

Methods

Genomic DNA was fragmented to 6kb and ligated with illumina adaptors, followed by size-selection at 4.5kb-9kb range. Long range PCR was performed on size-selected inserts. Pre-Capture PCR products were hybridized with a set of NimbleGen SeqCap probes targeting 94 genes that represent a spectrum of diseases with low, and thus unsatisfactory, sequencing access by current short-read NGS methods. Captured products were processed by standard PacBio large insert library preparation procedure. The final product, a 6kb insert capture library with single molecule, real-time (SMRT) bell adaptor, was sequenced on PacBio RSII platform.

Results

Initial PB-LITS protocol has shown its value of discerning complex structural rearrangements with 1.5ug input DNA. Further development of PB-LITS enabled successful construction of 6kb-insert capture library with minimal 150ng input, by optimizing adaptor ligation and long range PCR reaction. Sequencing with 2~3 SMRT cells of HapMap DNA libraries constructed by current protocol yielded 20x coverage of 93% of 14mb target size with an average insert size of 3.7kb-7.1kb. Sequencing and further analysis of the libraries constructed from 6 cases with strong family histories of breast cancer who previously tested negative for mutations in BRCA1 and BRCA2 are undergoing.

Conclusion

Optimized PB-LITS protocol provided a reliable research tool for studying complex chromosomal SVs. Development of library construction procedure of PB-LITS has laid solid foundation to employ the method in clinical settings where identification of pathogenic complex SVs with limited sample amount is challenging.

References


Disclosure of interest

None declared.

OS58

Rhesus macaques exhibit more non-synonymous variation but greater impact of purifying selection than humans

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Human Genomics 2016, 10(Suppl 1):OS58

Objectives

We generated whole genome sequence data for 133 rhesus macaques (Macaca mulatta), the primary nonhuman primate in biomedical research.

Methods

Using the intersection of GATK and SNPTools SNV calls, we identified 43 million high-quality SNVs, including >126,000 missense and >148,000 synonymous coding variants.

Results

Comparisons with equivalent whole genome data from the Human 1000 Genomes project shows that macaques have 2.5-fold higher levels of overall variation and 20% higher levels of nonsynonymous variation per individual. Comparing the ratio of nonsynonymous to synonymous variants between species shows a lower ratio of NS:Syn
in macaques, indicating more effective purifying selection, which can be explained by higher effective population size. Looking more specifically at 740 genes from the SFARI autism genetic association database, the ratio of NS:Syn variants among the macaques is lower in the SFARI gene set than in the complete macaque gene set.

**Conclusion**

These data suggest that large effective size in macaques leads to higher levels of both total and nonsynonymous variation than humans, and that purifying selection in macaques is more efficient in restricting mildly deleterious mutations.

**Disclosure of interest**

None declared.

O59

Assessing RNA structure disruption induced by single-nucleotide variation

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**Human Genomics 2016, 10(Suppl 1):O59**

**Objectives**

It has been challenging to interpret noncoding variants in complex traits and human diseases. RiboSNIitches, single-nucleotide variants (SNVs) that alter the structures of RNAs, have recently found in many human diseases. RNA structure change mediated by riboSNIitches has emerged as a plausible mechanism of pathogenic consequences of mutations. Thus, it is desirable to automatically predict riboSNIitches from millions of SNVs of the human genome. However, current computational methods based on *in silico* RNA-folding algorithms suffer from limited accuracy. We seek to develop a new method for improved riboSNitch detection.

**Methods**

We introduce a new measurement to quantify the structural difference between the wild-type and mutant RNAs in which the two sequences are differed by an SNV. The new measurement is based on assessing the consistency of RNA structure change of individual bases. Using this new measurement, our method automatically selects a region that maximizes the effect of RNA structure disruption induced by an SNV.

**Results**

We applied our method to analyze a genome-scale dataset of riboSNIitches and non-riboSNIitches determined from the parallel analysis of RNA structure experiments on a family trio of human lymphoblastoid cell lines. The dataset contains rigorously validated subsets of 11 “probed”, 63 “validated”, and 223 “symmetric” riboSNIitches. We found that our method consistently outperforms other existing methods on these rigorously validated subsets of riboSNIitches.

**Conclusion**

Our new method improves the accuracy of computational prediction of riboSNIitches. It facilitates the prioritization of noncoding variants for interpreting personal genomes. It also holds the promise to identify disease-causing variants potentially through RNA structure disruption.

**Disclosure of interest**

None declared.

P2

Missense polymorphic genetic combinations underlying down syndrome susceptibility

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**Human Genomics 2016, 10(Suppl 1):P2**

**Objectives**

Single nucleotide polymorphisms (SNPs) draw much attention as prospective biomolecular markers for human health management and disease therapies. Moreover SNPs have been surmised to constitute unique genetic makeup characteristic to each human individual, which can be utilized to cater therapeutic approaches to personalized medical care. Down syndrome (DS) or trisomy 21 is the most frequently occurring birth-related defects affecting live-born children. Molecular mechanisms that regulate and/or result in the formation of trisomy 21 in DS mothers, remain hitherto unknown. We posit a genetic basis for disposition of DS occurrence. We therefore performed bioinformatics studies of published nonsynonymous SNPs in conjunction with structural information of the proteins encoded by DS risk genes in the attempt to identify novel governing principles of DS risk.

**Methods**

We surveyed all SNPs in the published literature focusing on missense mutations, and superimpose on bioinformatic reconstruction of secondary structural motifs of proteins encoded by DS genes.

**Results**

In our survey, we observed that even the most penetrant SNP implicated in DS is not completely associated with the disease. On the other hand, a combination of co-occurrence of SNPs is important, suggesting a synthetic cooperation between missense mutations to underlie occurrence of DS phenotype. Superimposing documented SNPs from several public databases showed a preferential localization of these SNPs with specific structural motifs within the proteins. Interestingly, we noticed several closely situated SNPs that have not been reported to be associated with DS risk within regulators of the one carbon folate metabolism that included reduced folate carrier 1 (RFC1) and methionine synthase. These may represent
novel SNPs that can be assessed experimentally in a targeted manner in future population studies.

**Conclusion:** Taken together, our analyses showed that SNPs that result in change of protein sequences act synergistically to impact DS phenotype and suggest that SNP combinations to be a more reliable criterion than single SNPs for Ascertaining DS risk, at least in the case of missense SNPs. Our study also identified probable secondary structural motifs implicated in DS risk-associating factors. These results will form the basis for future experiments that may hold potential for translation into personalized diagnosis or therapeutic management of DS.

**Competing interests**
None declared.

**P8**
Differentiating inflammatory bowel diseases by using genomic data: dimension of the problem and network organization

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**Human Genomics 2016, 10(Suppl 1):P8**

**Objectives:** To determine the minimum number of genes (dimension of the problem) and their network organization in the distinction between Crohn’s Disease (CD), Ulcerative Colitis (UC) and normal patients (N).

**Methods:** We relied on the results published by Burczynski et al. where transcriptional profiles in peripheral blood mononuclear cells (PBMC) from 42 healthy individuals, 59 CD patients, and 26 UC patients were assessed. We applied a newly proposed gene selection method, based on statistical and machine-learning principles, which finally delivered a set of models which best predicted the disease class. These models were inserted in a network where the biomarkers were placed in specific positions according to their relevance in discriminating between the diseases.

**Results:** We found that a set of models, containing only two RNA’s from the PBMC, were sufficient to discriminate CD from UC patients and normal individuals. These RNA’s were organized in networks where the gene in first position could well classify when placed in a model with any of those in the corresponding second position. A summary of these networks is as follows.

**Conclusion:** Our statistical method is a new powerful tool that gives (1) the dimension of the statistical model, (2) the network organization of the selected genomic biomarkers, (3) a set of interchangeable models giving the same information.

Moreover, all the RNA’s in position 1 of the selected networks are known to have a clinical significance in Inflammatory Bowel Diseases. Amyloidosis is a well-known complication of CD and UC (Amyloid beta A4 precursor protein). Chemotaxis and neutrophil activation are fundamental pathways in the pathogenesis of IBD (Chemokine C-X-C motif ligand 5). RAB31 enhances FcγR-mediated phagocytosis through P38K/Akt signaling in macrophages and plays a role in the maturation of phagosomes. Some clinical evidence links the menstrual cycle to the IBD activity (Progestrone receptor membrane component 1).

**References**

**Competing interests**
None declared.

**P9**
Vulnerability of genetic variants to the risk of autism among Saudi children

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**Human Genomics 2016, 10(Suppl 1):P9**

**Objectives:** Single nucleotide polymorphisms (SNPs) have been reported in different autistic populations. Here we present the first association study investigating SNPs of some genes; serotonin receptor (HTR2A IVS2A>G rs7997012; HTR2C 68G>C rs6318), serotonin transporter (SLC6A4 rs8313034), ankyrin repeat and kinase domain containing 1 (ANKK1 rs1800497), methylentetrahydrofolate reductase (MTHFR rs1801394), and BDNF rs6265 in Saudi autistic children. Epidemiologic, clinical and psychometric aspects were used to examine the possible risk factors of autism.

**Methods:** We used TaqMan SNP genotyping to examine 68 Saudi children (48 Males and 20 females) diagnosed with autism according to DSM IV criteria & ICD-10 criteria, including deficits in reciprocal social interaction, impaired verbal, and non-verbal communication as well as restricted, repetitive, and stereotyped patterns of behaviors. Healthy controls (n= 78) with no history of mental illnesses, behavioral disorders or substance abuse were used. The severity of behavioral symptoms in cases was assessed at admission using the Childhood Autism Rating Scale (CARS). Hardy-Weinberg equilibria of the genetic variants were assessed using online software (http://www.zege.org/software/hwe-mr-calc.shtml). Chi-square tests were used to compare sociodemographic and clinical characteristics. Odds ratios and confidence intervals were calculated.

**Results:** Overall, our data provide strong evidence of associations between these two SNPs and risk of autism in this population. Compared to healthy subjects, children with autism showed significant overexpression of the mutant alleles in the SNPs rs7997012, rs6318, rs8313034, rs1800497, and rs1801394, but not in the rs6265 SNP. Data on linkages or associations between these genetic loci and the disease vary among different ethnicities.

**Conclusion:** Our findings presented associations between autism and some genetic variants under investigation, and showed that the potential influence of just one copy of the mutant alleles in the Saudi patients. Replicating our findings in other ethnic population may support our data before making any firm generalizations. Ongoing analyses of genetic variants associated with autism are being extended using next-generation sequencing and some fruitful data are going to be published.

**Competing interests**
None declared.

**P10**
Chromatim profiles from ex vivo purified dopaminergic neurons establish a promising model to support studies of neurologica function and dysfunction

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**Human Genomics 2016, 10(Suppl 1):P10**

**Objectives:** The majority of variants associated with common disease reside in non-coding sequence. Efforts to understand complex disease risk thus focus heavily on regulatory variation. How to seek biologically relevant sequences in which variation may reside and how to make in vivo predictions of the impact and function of regulatory variation remains a major challenge. We recently demonstrated how GWAS results can be informed by enhancer analyses in homogenous,
phenotype-appropriate cell populations. Further, such enhanced datasets make feasible the prediction of functional variants. Presently, we are leveraging our past experiences to generate enhancer catalogues using homogenous populations of dopaminergic (DA) neurons to better understand the role of regulatory variation in Parkinson Disease (PD) and related disorders.

Methods: We assayed open chromatin regions using ATAC-seq, producing multiple enhancer catalogues. Each catalogue is generated from 50,000 FAC-sorted DA neurons from either the forebrain (n=3 libraries) or midbrain (n=3 libraries) of E15.5 transgenic mice. Each ATAC-seq library is of high quality, yields over 30 million reads sequenced, with over 28 million reads mapping, and following filtering of reads, MACS2 calls ~50,000 peaks, indicating intervals of open chromatin.

Results: Our preliminary analyses indicate that peaks from all libraries show evidence of functional sequence constraint (PhastCons scores>0.3) and significant enrichment for processes and functions appropriate to neuronal function/dysfunction by GO/GREAT. The high quality of these open chromatin signatures also facilitate development of a computational classifier (regulatory vocabulary) of DA neurons. Consequently, we have begun to gain further insight into the transcription factors active in DA neurons and the nature of variation that might influence the function of identified DA enhancers.

Conclusion: We are currently validating the DA neuron enhancer catalogues in vivo and are evaluating the shared and unique content of the catalogues and their pertinence to disease. Additionally, we have generated single cell and bulk RNA-seq from these isolated populations to corroborate and inform our chromatin-based findings. With this data in hand, we are able to assay the impact of PD and related movement disorders’ GWAS-implicated variation on DA neuron function and disease pathogenesis.

Competing interests
None declared.

P11
Utilization of a sensitized chemical mutagenesis screen to identify genetic modifiers of retinal dysplasia in homozygous Nr2e3rd7 mice

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Human Genomics 2016, 10(Suppl 1):P11

Objectives: To identify and characterize genetic modifiers capable of altering the retinal dysplasia observed in Nr2e3rd7 mutants, a model for human Enhanced S-Cone Syndrome (ESCSD). Methods: Nr2e3rd7 mice were chemically mutagenized, and mated to generate a G3 population. The G3 mice were screened by indirect ophthalmoscopy to establish lines bearing genetic modifiers that altered the pan retinal fundus spotting phenotype, characteristic of homozygous Nr2e3rd7 mice. Quantitative trait locus analysis combined with high-throughput sequencing of an exome capture libraries was used to identify the molecular basis of modifiers of the retinal dysplasia in Nr2e3rd7 mutants. Apart from fundus imaging, longitudinal histological studies were carried out to characterize the progression of altered retinal phenotypes. Finally, immunoblotting and marker analyses were performed to reveal the defects that underlie the rd7 retinopathy and how the defects were affected by the modifier(s).

Results: Seven heritable modifier mouse lines with an altered retinal phenotype have been established so far. Among them three potential genetic modifiers have been identified, which directly or indirectly are associated with growth and development of neuroretinal cells. For example, the Tvrn222 line bears a nonsense mutation of the Rab6 gene, which leads to a vitreal dysplastic phenotype that is more severe in the presence of the Nr2e3rd7 mutation. While a reduced retinal spotting phenotype and suppression of formation of rosette-like structures associated with the rd7 retinopathy in the Frmd4b gene. The modifying effect of the Frmd4b.Tvrn222 allele is achieved through its effect on cell-cell junctions, revealed by immunoblotting and marker analysis.

Conclusion: As an animal model of ESCS, retinal dysplasia in Nr2e3rd7 mouse can be phenotypically altered by multiple genetic modifiers via different pathways. The modifying effects on rd7-associated retinopathies by these particular genetic modifiers, to our knowledge, are the first to be described. They provide novel insights into the pathogenesis of retinal dysplasia as well as degeneration in Nr2e3rd7-associated disease, and may become potential interfering targets for clinical applications against ESCS and related retinopathies.

Competing interests
None declared.
Competing interests
None declared.

P13
Digital expression profiling of Purkinje neurons and dendrites in different subcellular compartments
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Methods: In previous work, we used a technology called translating ribosome affinity purification (TRAP) to isolate the ribosome-associated transcriptome — the translatome. We modified it to target any cell type that can be specifically infected by a modified adeno-associated virus, and applied it to Purkinje cells (PCs) in the rat cerebellum. We obtained quantitative expression data in single-base-pair resolution by profiling the ribosome-associated, isolated RNA using the nanoCAGE protocol.

Results: Subsequent data analysis revealed the landscape of ribosome-associated RNA of PCs in different subcellular compartments: cytoplasm and rough endoplasmic reticulum and dendrites.

Conclusion: In neurons, protein translation occurs not only in the soma but also distally in dendrites near or within the dendritic spines. This distal translation is thought to be regulated in response to external stimuli including long-term depression and memory formation. We have applied TRAP to Purkinje dendrites and sequenced the isolated RNA with an improved nanoCAGE protocol including a tagmentation step, to address the increased difficulty of sequencing from dendrites, which contain even less RNA than Purkinje cell soma.

References

Competing interests
None declared.

P14
The evolution of imperfection and imperfection of evolution: the functional and functionless fractions of the human genome
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Human Genomics 2016, 10(Suppl 1):P14

Objectives: Genomes are products of natural processes. Hence all genomes contain functional and nonfunctional parts. Here, I present a functional classification of genomic elements and estimate the functional fraction within the human genome.

Methods: The classification into different categories of functionality were based on the concept of selected-effect function. Intraspecific and interspecific genomic comparisons and standard evolutionary methodology were used to infer the functional fraction within the human genome.

Results: According to their selected-effect function, the genome is divided into functional and rubbish DNA. Functional DNA is further subdivided into literal and indifferent DNA. In literal DNA, the order of nucleotides is under selection; in indifferent DNA, only the presence or absence of the sequence is under selection. Rubbish DNA is further subdivided into junk and garbage DNA. Junk DNA neither contributes to nor detracts from the fitness and, hence, evolves under selective neutrality. Garbage DNA, on the other hand, decreases the fitness of its carriers; it exists in the genome because natural selection is neither omnipotent nor instantaneous. Each of these four functional categories can be transcribed and translated, transcribed but not translated, or not transcribed. The affiliation to a particular functional category may change during evolution: Functional DNA may become junk DNA, junk DNA may become garbage DNA, and so on; however, in the absence of prophetic powers determining the functionality or nonfunctionality of a genomic sequence must be based on its present status rather than on its potential to change in the future. Changes in functional affiliation are categorized into pseudogenes, Lazarus DNA, zombie DNA, and Jekyll-to-Hyde DNA. Intraspecific and interspecific genomic comparisons indicate that the functional fraction in the human genome ranges from 8% to 15%.

Conclusion: A common misconception exists according to which evolutionary processes can produce a genome that is wholly functional. Actually, evolution can only produce such a genome if and only if the effective population size is infinite; the deleterious effects of increasing genome size by even one nucleotide are considerable, and the generation time is short. Not even in the commonest of bacterial species are these conditions met. In species with small effective population sizes and long generation time, such as humans, a genome that is ~100% functional is contrary to reason.

Competing interests
None declared.

P16
Species-independent identification of known and novel recurrent genomic entities in multiple cancer patients
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Human Genomics 2016, 10(Suppl 1):P16

Objectives: Here we present a new method for the identification of recurrent genomic entities that play a causative role in the onset of disease. Our approach is particularly amenable for the analyses high-throughput sequencing data.

Existing approaches often follow a bottom-up approach where taxonomic determination necessarily takes place before associations to disease can be determined; naturally failing to establish the causality of novel pathogens not present in reference databases.

Methods: To overcome this intrinsic limitation, we have developed a species-agnostic top-bottom approach that clusters sequences and identifies co-occurrence in multiple patients, associates recurrent sequences to disease and, finally, determines the taxonomic content where existing knowledge permits.

Results: We analysed 686 sequencing libraries from 252 cancer specimens and 56 controls. Recurrent sequences were statistically associated to biological, methodological and technical features to identify novel pathogens and contaminants stemming from laboratory reagents.

Conclusion: We provide examples of identified inhabitants of the healthy tissue flora, known experimental contaminants and
uncharacterised sequences that co-occur with high statistical significance with disease. The latter represent a category that can only be addressed by a species-independent approach. Thus, our method helps to chart the unknown sequence-space where novel pathogens can be identified.

Competing interests
None declared.

P18
Discovery of active gene modules which are densely conserved across multiple cancer types reveal their prognostic power and mutually exclusive mutation patterns
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Human Genomics 2016, 10(Suppl 1):P18

Objectives: An active module which is densely conserved across multiple gene networks exhibit strong and conserved interactions among its active member genes across all the networks. Identification of such modules across multiple species using gene networks in stem cell differentiation has provided new insights into evolutionary conserved developmental pathways and conserved pioneer factors. Similarly in cancer, such modules (if exists) will represent core common carcinogenesis-driving pathways in multiple types of cancer. These common pathways can have the same progeny in cancers stemming from different tissues. Even though, a comprehensive amount of gene expression data for multiple types of cancer has been acquired, there is no study on comparison of several cancer gene networks. In this study, we perform local alignment of 8 cancer gene networks to identify active gene modules which are densely conserved across the cancer types. We further study the prognostic and mutation patterns within these modules.

Methods: We developed a computational framework to identify active modules of genes which are densely conserved across 8 different types of cancer by extending a previous tool called neXus to allow it to work on more than 2 cancer gene networks. We identified 174 modules of genes satisfying these three strict criteria – 1) member genes were differentially expressed between normal and cancer tissues (FDR < 0.05) in at least 7 cancer types of cancer of interest, 2) average fold change of 2 in the member genes across all 8 cancer types and 3) the clustering coefficient of at least 0.5 in all the 8 cancer gene networks.

Results: The identified modules represented some known pathways associated with multiple cancers such as the RAS pathway, but several new genes which may play important in triggering cancer in multiple types of tissues were discovered. We found that these conserved modules of genes have high prognostic power in all the 8 types of cancer of interest. We also found that these active and conserved modules exhibit similar mutually exclusive mutation patterns among the gene members and these patterns were also conserved across the majority of the types of cancer.

Conclusion: These findings reveal a new set of previously unknown pathways, which most likely run as a common thread during carcinogenesis in many cancer types and hence are worth investigating further for experimental validation.

Competing interests
None declared.

P19
Whole exome sequencing of dysplastic leukoplakia tissue indicates sequential accumulation of somatic mutations from oral precancer to cancer
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Human Genomics 2016, 10(Suppl 1):P19

Objectives: Oral leukoplakia (OL) is the most common precancerous lesion in the oral cavity. The percentage of individuals with dysplastic OL in whom there is malignant transformation to oral squamous cell carcinoma (OSCC) is high, up to 36%. Germline and somatic copy number variations in mitochondrial DNA of OL patients have earlier been noted. We sought to test the hypothesis that about 36% of patients with the pre-cancerous lesion (OL) will possess somatic mutations in genes that are recurrently mutated in OSCC.

Methods: Whole exome sequencing of DNA isolated from the affected oral tissue and from peripheral blood of twelve OL patients with dysplasia, was used to profile the landscape of autosomal somatic recurrent mutations in OL and to investigate whether mutations in the genes that drive OSCC are present in OL patients or whether the mutational landscapes of OL and OSCC are largely disjoint.

Results: We have detected mutations in some genes that drive both oral leukoplakia and oral cancer. TGFBR2 is recurrently mutated in OL as well as in head and neck squamous cell carcinoma (HNSCC). Some significantly mutated genes in OSCC or HNSCC, viz., FAT1, NOTCH1 and CDKN2A are also found to be mutated in OL patients. Further, we have identified that MAPK signalling and oxidative phosphorylation (OXPHOS) pathways are significantly altered in OL patients.

Conclusion: We have found that the proportion of OL patients with epithelial dysplasia among whom mutations were found in the set of genes that is also recurrently mutated in OSCC/HNSCC, closely corresponds to the fraction (~36%) of patients with dysplastic leukoplakia who develop oral cancer. The leukoplakia patients recruited in this study were free of malignancy in the oral cavity; our results are, therefore, not influenced by field cancerization.

Competing interests
None declared.

P21
Epigenetic mechanisms of carcinogenesis by hereditary breast cancer genes
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Human Genomics 2016, 10(Suppl 1):P21

Objectives: BRCA2-induced breast cancers share a predominant histologic and molecular phenotype (ER+, luminal B) that distinguishes them from most sporadic breast cancers and breast cancers arising in other inherited disorders. This suggests that breast cancers arising in BRCA2-mutation carriers have essential shared properties that drive carcinogenesis and can be targeted for intervention. To investigate this observation, we performed a cell biological screen in non-transformed breast epithelial cells for phenotypes specific to BRCA2 loss-of-function.

Methods: Non-transformed breast epithelial cells were treated with siRNAs targeting hereditary breast cancer genes. Growth curves were obtained in complete growth media and growth factor-withdrawal media. Whole genome sequencing and transcriptomics were performed. Functional cell biological assays including treatment with recombinant cytokines and inhibitors were performed.

Results: Despite the role of BRCA2 in homologous recombination-mediated DNA repair, no recurrent de novo mutations were recovered. Instead, we discovered a novel pathway whereby BRCA2 depletion induces strong, persistent transcriptional activation of the chemokines on chromosome 4q13 (CXCL1, CXCL3, CXCL5, CXCL8). Surprisingly, these chemokines were sufficient to stimulate EGF-independent growth of non-transformed breast cells. Furthermore, inhibitors of the receptors of the chemokines impaired cell proliferation after BRCA2 depletion.

Conclusion: Altogether, these findings indicate that transcriptional activation of the 4q13 chemokine locus induces an early cell-autonomous autocrine signaling pathway in BRCA2-mediated carcinogenesis that could be exploited to prevent cancer onset.

Competing interests
None declared.
RNA direct: a novel RNA enrichment strategy applied to transcripts associated with solid tumors

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Objectives: RNA sequencing (RNA-seq) is a powerful tool used for the interrogation of transcripts that enables the analysis of gene expression and the identification of nucleotide or structural variations. RNA-seq, however, can be cost prohibitive due to the size and complexity of most transcriptomes, which require deep sequencing coverage to achieve the necessary sensitivity for a reliable analysis. By combining targeted enrichment strategies with next-generation sequencing (NGS), a subset of transcript regions or sequences can be enriched and analyzed with the resolution required for both research and clinical applications. Here, we applied a novel RNA enrichment strategy, RNA DIRECT, to the capture of targets commonly associated with solid tumors.

Methods: RNA DIRECT target enrichment utilizes cDNA and probe-based hybridization to capture only desired cDNA sequences with removal of off-target regions via enzymatic digestion. Targeted sequences are then ligated to NGS platform specific adapters and amplified by PCR.

Results: RNA converted to cDNA was used for targeted enrichment with RNA DIRECT. Analysis of sequenced reads showed at least 97% alignment to the transcriptome with greater than 90% aligning to targeted regions. We also report the identification and detection of variants and gene fusions associated with solid tumors with high specificity and sensitivity.

Conclusion: The RNA DIRECT strategy provides a robust and cost-effective method for the enrichment of targeted transcript sequences for the identification and detection of known or novel variants and gene fusions.

Competing interests
None declared.

RNA sequencing identifies gene mutations for neuroblastoma

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Human Genomics 2016, 10(Suppl 1):P23

Objectives: Neuroblastomas are the most common of all malignancies in infants and the most common extracranial malignancy of childhood. With 650 new cases each year it accounts for over 15% of childhood cancer mortality. This neuroectodermally derived malignancy is most often found in the adrenal glands but can be found throughout the body in sympathetic ganglia. Though a small number of cases are familial (1-2%) with known genetic causes, most cases are sporadic with little known about what causes them or what causes the diverse outcomes of this disease.

Methods: The current study used transcriptome sequencing to locate possible genetic mutations from a small cohort of neuroblastoma samples. The mutations were used to construct a phylogenetic tree that demonstrated the tumor progression and predicted the outcome.

Results: Using this method a handful of associated mutations have been found, but few effect disease outcome. In this study we used next generation RNA sequencing to fully sequence 249 neuroblastoma samples. Using these samples and focusing on indel mutations we located 1247 mutated genes that affect the favorable or unfavorable outcome of the disease. Due to the large number of mutations, online databases were used to identify genes associated with neuroblastomas. Comparing these genes to our sample we found a subset of genes that when mutated affected the survival rate of patients.

Conclusion: This study suggests the importance of a few genes that drive the progression of neuroblastomas and determine the clinical outcomes.

Competing interests
None declared.

Targeted Methylation Sequencing of Prostate Cancer

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Human Genomics 2016, 10(Suppl 1):P25

Objectives: Aggressive and potentially life-threatening prostate cancer often requires radical treatment. The many side effects that are associated with such treatments underscore the importance of accurate differentiation between these aggressive tumors and indolent prostate cancer in order to reduce overtreatment. Thus, we focus on the discovery of prognostic signatures based on changes in DNA methylation, stratifying by disease aggressiveness.

Methods: Whole-genome bisulfite sequencing (WGBS) is cost-prohibitive for analyzing large numbers of human cancer specimens. Further, in order to detect subtle methylation differences between cancer samples and in samples with low tumor purity, higher
sensitivity is achieved through a greater read depth, further increasing sequencing costs. Therefore, we utilize a targeted bisulfite sequencing method for the comprehensive analysis of genomic regions relevant to cancer, comprising 8 megabases (~3% of the human genome), covering 3.7 million CpGs, including most RefSeq and GENCODE gene promoters, all known cancer genes, CpG islands, and their shores. Up to 100-fold read depth can be achieved on average by pooling four human prostate samples on one Illumina HiSeq4000 lane.

Results: Of the differentially methylated regions (DMRs) we detected, the majority is hypermethylated in the aggressive versus the indolent prostate tumors. Most DMRs overlap a CpG island and are predominantly found in or close to gene promoters, as well as the promoter regions of long non-coding RNAs, and to a lesser extent within gene bodies or intergenic. Often, we find strong hypermethylation of the up- and downstream CpG islands surrounding the transcription start site (TSS), while the TSS itself stays unmethylated. Furthermore, ~70% of the hypermethylated DMRs overlap regions reported to be a bivalent promoter in various cell types from the ENCODE project. The DMRs that can be associated with RefSeq genes are enriched for transcription factors, and more than half of those contain a homeodomain, such as different members of the HOX, SOX and FOX gene families.

Conclusion: The overarching theme for DMRs called in this set of prostate cancers is that the targeted deep Methylation Sequencing approach is hypermethylation of regions within bivalent CpG islands, presenting a prognostic methylation signature that warrants further investigation.

Competing interests
None declared.

P26

Mutant TPMT alleles in children with acute lymphoblastic leukemia from México City and Yucatán, Mexico
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Human Genomics 2016, 10(Suppl 1):P26

Objectives: To know the frequency of the TPMT deficient alleles in children with acute lymphoblastic leukemia and healthy subjects from two Mexican populations
Methods: We included 813 unrelated subjects, 392 were children with ALL and 421 were healthy subjects. Genotyping of the rs1800462, rs1800460 and rs1142345 SNPs was performed by TaqMan assays. To assess the differences of the genotype and allele frequencies among groups we used the Chi-square test. Written informed consent was obtained from both ALL children’s parents and healthy participants.

Results: The mutant TPMT alleles were carried by 5% of the 1636 samples obtained from both ALL children and healthy participants.

Conclusion: The mutant TPMT alleles performed in ALL Mexican pediatric patients [1]. Because ALL is the leading cause of childhood cancer in México [2] and homozygotes TPMT deficient alleles subjects have high risk to develop severe and potentially fatal hematopoietic toxicity after treatment with standard doses of thiopurines [3]; TPMT alleles genotyping should be performed in Mexican ALL patients. Furthermore, large-scale genotype and phenotype correlation studies are needed to assess the contribution of this variants in other Mexican-American populations.

References

Competing interests
None declared.

P28

Genetic modifiers of Alström syndrome
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Human Genomics 2016, 10(Suppl 1):P28

Objectives: Alström syndrome (AS) is a progressive multi-systemic disorder caused by recessive mutations in the ciliary protein ALMS1. Amongst patients with AS, we observe variability in onset and/or severity of disease phenotypes including hearing and vision loss, obesity, hyperinsulinemia, hepatosteatitis, and cardiomyopathy, likely due to the presence of modifier genes. Like in patients, disease phenotypes in murine models of AS can be modified by the genetic background. The goal of the study was to map genetic modifier loci using an AS mouse model.

Methods: A genetrap in intron 13 of the mouse Alms1 gene was placed onto the C57BL/Ei and BALB/c inbred background strains, respectively. To genetically dissect modifier loci of AS, we performed a phenotypic screen of 135 Alms1Gt/Gt backcross progeny from two backcrosses: ((C57BL/Ei X Balb/c)F1-Alms1+/Gt X C57BL/Ei-Alms1+Gt) and ((C57BL/Ei X Balb/c)F1-Alms1+/Gt X Balb/c/c-Alms1+/Gt). DNA of the backcross progeny was typed using evenly spaced microsatellite markers throughout the genome and quantitative trait locus (QTL) analysis was performed. Recombinational fine mapping and characterization of subcongenic lines was used to refine a retinal degeneration QTL on Chr. 2.

Results: QTL for body weight, plasma insulin and triglyceride levels, alanine aminotransferase levels, hepatic steatosis, hepatic fibrosis, and retinal degeneration were mapped to regions on five chromosomes. The location of a major modifier locus on Chr. 2 in which the B6 allele protects Alms1GtGt retinas from rapid photoreceptor degeneration was refined to a 12 Mb region. A candidate mutation in the glutamylase TTLL9 was identified and is associated with reduced glutationylation of tubulin in Balb/c/Alms1GtGt retinas.

Conclusion: Elucidation of the genetic networks of Alms1 may lead to a better understanding of the role of Alms1 in metabolic and neurosenory disease and may provide novel targets for therapeutic intervention.

Competing interests
None declared.

P31

Association of genomic variants with the occurrence of angiotensin-converting enzyme inhibitor (ACEI)-induced coughing among Filipinos
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Human Genomics 2016, 10(Suppl 1):P31
Methods: A case-control study involving 186 unrelated patients who were taking ACEI for at least 6 months was done (101 males, 85 females; 62 cases, 124 controls). DNA from blood samples were extracted and were genotyped using customized Illumina Goldengate microarray chips for 384 gene variants.

Results: Results show that allelic variants of the genes ZPR1, ADAMTS7, CTB-129P6.7 and LOC157273 are significantly associated with ACEI-induced coughing (OR: 2.34, 3.120, 2.49 and 2.64, respectively, p<0.01). Further, an intergenic loci in chromosome 4 is also significant. Interestingly, using logistic regression, in addition to ZPR1, ADAMTS7 and CTB-129P6.7, eight variants located proximal to each other in the X chromosome have been associated with the ACEI-induced coughing.

Conclusion: The study presents possible pharmacogenetic markers for Filipinos, as well as genomic regions of interest that may further shed light on the mechanisms of ACEI-induced coughing.

Competing interests
None declared.

P34
Molecular regulation of chondrogenic human induced pluripotent stem cells
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Objectives: Human induced pluripotent stem cells (hiPSC) are a promising source for chondrogenic stem cells. Sequential differentiation of hiPSC provides a platform for dissecting the molecular pathways associated with chondrogenesis in vivo and could reveal targets for better control of chondrocyte fate for cartilage repair applications. The aim of this study was to use next generation sequencing (NGS) to investigate the transcriptome and the methylome of chondrogenic hiPSCs.

Methods: The hiPSC line (C19) was derived through reprogramming of human dermal fibroblasts using viral vectors expressing Oct4, Sox2 and Klf4. A protocol of sequential growth factors including Activin A, FGF2 and BMP4 was used to drive the formation of chondrogenic progenitors directly from hiPSC colonies. The chondrogenic hiPSCS were characterised exhaustively by tissue engineering, histochimical analysis and biochemical analysis. The transcriptome of undifferentiated hiPSCs, hiPSC derived chondrocytes and native chondrocytes was interrogated utilizing RNAseq on the SOLID S500 XL platform where the polyA fraction was sequenced at a coverage level of at least 25 million reads.

Results: Differential gene expression revealed the induction of several collagen genes including type 1 to type 12, type 14 and type 18 during transition from the pluripotent state to the chondrogenic state (Fig. 7). Collagen regulatory genes such as PCOLCE which drives the endopeptidase cleavage of procollagen as well as regulators of collagen glycosylation were upregulated. The expression of various fibroblasts growth factors (FGFs) including FGF11, FGF22 and insulin growth factor2 (IGF2) was upregulated as well as the Wnt induced secreted protein, WISP2. Mitotic genes were downregulated in derived and differentiated chondrocytes. Integrated methylome and transcriptome analysis revealed the step-wise differentiation process.

Conclusion: NGS analysis demonstrated the recapitulation of early events in cartilage development during hiPSC chondrogenesis. The upregulation of many members of the collagen family indicate the intricate nature of collagen expression during chondrogenesis. Further analysis of nonchondrogenic targets may reveal novel pathways for controlling the fate of chondrogenic hiPSCs.

Competing interests
None declared.

References
Molecular profiling of hematologic malignancies: implementation of a variant assessment algorithm for next generation sequencing data analysis and clinical reporting

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Objectives: Next generation sequencing (NGS) can be used to identify clinically relevant variants for accurate diagnosis, prognostic risk stratification, and identification of therapeutic targets for genotype-matched trials in cancer. Our objective was to design methods for efficient prioritization of variants for interpretation and reporting. We describe a triaging algorithm to identify clinically relevant variants from tumour-only analysis of NGS in hematological malignancies.

Methods: Blood or bone marrow DNA from 260 patients recruited in the Princess Margaret Cancer Centre’s Advanced Genomics in Leukemia AGILE trial were profiled using the 54 gene TruSight Myeloid Sequencing Panel (Illumina). Variants were called using the Illumina MiSeq Reporter (MSR) software. Data were uploaded into a commercially available tool, Cartagenia Bench NGS, for filtering and analysis.

Results: Of all variants detected by NGS (median 427, range 338–643 variants/case), 35% (median 150, range 125–172 variants/case) passed all MSR quality criteria. Applying a variant allele frequency threshold refined the data to 7.4% of the original dataset (median 30, range 16–48 variants/case). Reporting was restricted to well-covered, exonic nonsynonymous, intronic splice site, and known pathogenic synonymous variants, resulting in a median of 4 variants/case for manual review (range 0–13). When combined with our dataset of >600 interpretations across 8 hematological malignancies, this approach enabled rapid review and interpretation of previously known variants, and an effective system to prioritize novel variants in order of clinical actionability (Sukhai et al., 2015). We excluded variants with high germline population frequencies (median 50, range 42–62 variants/case) through the use of multiple reference population databases.

Conclusion: We describe our approach to prioritize NGS derived variants, based on data quality, functional effects, allele frequency, coverage depth, and coding effects. This approach iteratively utilizes our lab-developed variant knowledge base, and enables us organize and use variant interpretations to generate clinical reports.

P36
Accessing genomic evidence for clinical variants at NCBI

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Objectives: NCBI provides many resources for evaluating and declaring evidence of pathogenicity for an increasing number of human sequence variants. Primary details about a genetic test’s analytical validity and clinical utility are reported in the Genetic Test Repository, and evidence for inferences of variant pathogenicity are summarized in ClinVar. NCBI has developed several tools that integrate these high level records with the more basic factual data describing them as population level variants (dbSNP/dbVar), and deeper still as individual-level observations with called genotypes (the NCBI Genotype Server), phenotypes (dbGaP), whole genome/exome sequences (SRA) and finally as positions on the reference genome (GRChuman reference assemblies).

Methods: Traversing the connections between variant-level records, e.g. GTR, ClinVar and dbSNP, and individual-level data (e.g. genotypes, sequences, samples, and phenotypes) is a computationally intensive activity, and NCBI has developed several new services to pre-compute these relationships and permit users to quickly move from summary records to individual level data.

Results: This presentation will introduce several of these services including allele registry, the ClinVar beacon search, and the dbGaP genome browser with particular emphasis on how users can access and review individual level data for clinical variants or ad hoc genomic positions of particular interest.

Conclusion: Examples of use include the review of ClinVar records by expert panels, identify research into the existence of specific sequence alleles, automatic notification when new data for specific potential alleles of interest are submitted to NCBI, confirmation of variant properties during manuscript review, and research in general questions of human genetic architecture.

P37
NGS-SWIFT: a cloud-based variant analysis framework using control-accessed sequencing data from DBGAP/SRA

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Objectives: Genetic variation analysis plays an important role in elucidating the causes of various human diseases. The drastically reduced costs of genome sequencing driven by next generation sequence technologies now make it possible to analyze genetic variations with hundreds or thousands of samples simultaneously, but with the cost of ever increasing local storage requirements. The tera- and peta-byte scale footprint for sequence data imposes significant technical challenges for data management and analysis, including
the tasks of collection, storage, transfer, sharing, and privacy protection. Currently, each analysis group must download all the relevant sequence data into a local file system before variation analysis is initiated. This heavy-weight transaction not only slows down the pace of the analysis, but also creates financial burdens for researchers due to the cost of hardware and time required to transfer the data over typical academic internet connections.

Methods: To overcome such limitations and explore the feasibility of analyzing control-accessed sequencing data in cloud environment while maintaining data privacy and security, here we introduce a cloud-based analysis framework that facilitates variation analysis using direct access to the NCBI Sequence Read Archive through SRA Toolkit, which allows the users to programmatically access data housed within SRA with encryption and decryption capabilities and converts it from the SRA format to the desired format for data analysis.

Results: A customized machine image (ngs-swift) with preconfigured tools, including SRA Toolkit and NGS Software Development Kit, and resources essential for variant analysis has been created for instantiating an EC2 instance or instance cluster on Amazon cloud. Performance of this framework has been evaluated using dbGaP study phs000710.v1.p1, and compared with that from traditional analysis pipeline, and security handling in cloud environment when dealing with control-accessed sequence data has been addressed. We demonstrate that with this framework, it is cost effective to make variant calls without first transferring the entire set of aligned sequence data into a local storage environment.

Conclusion: This direct data access approach using NCBI SRA Toolkit from cloud for next generation sequencing analysis is more cost-effective in terms of time and disc spaces being used for the analysis, and thus will accelerate variation discovery using control-accessed sequencing data.

Competing interests
None declared.

P40
Flowr: robust and efficient pipelines using a simple language-agnostic approach; ultraseq: fast modular pipeline for somatic variation calling using flow
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Human Genomics 2016, 10(Suppl 1):P40

Objectives: Bioinformatics analyses have increasingly become compute intensive processes, with lowering costs of data production and increasing numbers of samples. Each laboratory spends time creating and maintaining a set of pipelines, which may not be robust, scalable, or efficient. Further, the existence of different computing environments across institutions hinders both collaboration and the portability of analysis pipelines.

Methods: Flowr is a robust and scalable framework for designing and deploying computing pipelines in an easy-to-use fashion. It implements a scatter–gather approach using computing clusters, simplifying the concept to the use of five simple terms (in submission and dependency types). Most importantly, it is flexible, such that customizing existing pipelines is easy, and since it works across several computing environments (LSF, SGE, Torque, and SLURM), it is portable.

Results: Ultraseq: Using Flowr's framework we have created a flexible pipeline for somatic variant calling. This aligns raw reads, splitting the processing across multiple nodes of a computing cluster. Results are gathered, creating a merged bam; which is then split according to the contigs in the genome. Further steps including de-duplication, indel realignment and base-quality recalibration and variant calling are performed at the contig level, using a scatter–gather approach. This enables complete processing of a typical whole-exome in about two hours, and whole-genome in a few hours (depending on the library size).

Conclusion: Flowr follows the “design once” principle, enabling the user to develop robust, portable pipelines that can be run on a host of computing platforms. Further, the same pipeline can be run on a local machine, computing cluster, or cloud-based environment. With automatic logging of each step and the preservation of the exact commands run to produce the output, the system allows users to generate an easy-to-use, efficient, and reproducible analysis pipeline. Ultraseq provides a fast and robust and modular pipeline for somatic variation calling. This enables users to easily add new tools and methods are they are released, and possibly share with other users in a format, which is robust, efficient and portable.

Availability: http://docs.flowr.space

Competing interests
None declared.
P41
Applying “Big data” technologies to the rapid analysis of heterogenous large cohort data
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Human Genomics 2016, 10(Suppl 1):P41

Objectives: Analysis of large cohort genomic data is a complex task, typical bioinformatics approaches use collections of scripts and R code, this becomes difficult to manage as sample numbers grow to 10s of thousands and file sizes grow to terabytes. We applied “Big data” technologies to the analysis of 578 Hi-seq X whole genomes (WGS) and 10,913 exomes (WES), in both cases we QC’d and integrated billions of variant calls from Atlas and GATK to provide concordant and consensus genotypes.

Methods: Raw variant data was pre-processed to produce key value pairs; keys representing variants and values representing sample specific data stored as JSON. Using a modest Hadoop test cluster (88 cores 512 Gb memory 7tb disk), we imported the data into Hbase tables. WGS data was stored as 578 columns, one column per sample. WES data was stored as a single column due to the large number of samples.

QC and analysis were performed by first building normalised summary tables in Hive, this process required a full Hbase table scan and took several hours to complete. However, once the data was processed, we were able to perform rapid analysis (minutes) on both the WGS and WES data sets using identical SQL queries regardless of how the underlying data was structured.

In this way we were able to perform QC filtering such as Ti/Tv, mappability, monomorphic sites, missingness rates and Hardy-Weinberg equilibrium with GATK. This became practical to iteratively tune parameters, in this way we were able to modify the Atlas genotype boundaries so as to maximise the numbers of concordant genotypes with GATK.

Results: Our queries were verified by comparison to 2 independent sets of QC statistics for the 578 WGS samples, once we had confirmed that we could exactly reproduce these numbers we were then able to quickly apply the same QC steps to the much larger 10,913 WES data set. Because the tests could be performed rapidly it became practical to iteratively tune parameters, in this way we were able to modify the Atlas genotype boundaries so as to maximise the numbers of concordant genotypes with GATK.

Conclusion: By leveraging the power of map-reduce and the rich ecosystem of tools around Hadoop we were able to streamline the analysis of these large genomic data sets, rapidly processing billions of genotypes using simple SQL queries to provide accurate, reproducible QC statistics without the need to track and manage thousands of files.

Competing interests
None declared.

Table 2 (abstract P41). Example of summary statistics comparing Atlas genotype calls with differing allele fraction thresholds with GATK calls.

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<th>Genotype</th>
<th>GATK</th>
<th>ATLAS (ORIGINALS)</th>
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<th>ATLASDP &gt;= 100.1/c/ 1&gt;0.9</th>
<th>ATLASDP &gt;= ATLAS (0.1/c-0) 1&gt;0.75</th>
<th>ATLASDP &gt;= ATLAS (0.1/c-0) 1&gt;0.75</th>
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</thead>
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<td>1,778,668,240</td>
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<td>1,778,668,240</td>
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<tr>
<td>0/1</td>
<td>138,744,203</td>
<td>69,153,666</td>
<td>112,165,710</td>
<td>147,763,797</td>
<td>124,056,267</td>
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<tr>
<td>1/1</td>
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<td>73,417,546</td>
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<td>1.14</td>
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<td>2.97</td>
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<td>1,964,251,496</td>
<td>1,976,142,053</td>
<td>1,976,147,802</td>
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</tr>
</tbody>
</table>

Table 2 (abstract P41). Example of summary statistics comparing Atlas genotype calls with differing allele fraction thresholds with GATK calls.

P42
FANTOM5 web resource for the large-scale genome-wide transcription start site activity profiles of wide-range of mammalian cells
T. Kasukawa1, M. Lizio1, J. Harshbarger1, S. Hisashi1,2, J. Severin1, A. Imad1, S. Sahin1, T. C. Freeman3, K. Baillie3, A. Sandelin4, P. Carninci1, A. R. R. Forrest1, H. Kawaji1,2,5, The FANTOM Consortium 1
1Center for Life Science Technologies, RIKEN, Yokohama, Japan; 2Preventive Medicine and Diagnosis Innovation Program, RIKEN, Wako, Japan; 3The Roslin Institute and Royal (Dick) School of Veterinary Studies, University of Edinburgh, Edinburgh, United Kingdom; 5Department of Biology & Biotech Research and Innovation Centre, University of Copenhagen, Copenhagen, Denmark

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Human Genomics 2016, 10(Suppl 1):P42

Objectives: To identify genome-wide transcription start site activities in various mammalian cells, a worldwide collaborative project, FANTOM5 (Functional Annotation of Mammalian Cells 5) was organized. In this project, diverse range of mammalian samples (~2000 for human and ~1200 for mouse, including primary cells, cancer cell lines, tissues, and transiting cells in time courses) was profiled to obtain promoter-level transcriptional atlas using high-throughput sequencers. To provide this large-scale data collection to the scientific community, we developed an integrated web resource, the FANTOM5 web resource.

Methods: To support various sort of inspections and analyses, the web resource contains several tools. SSTAR (Semantic catalogue of Samples, Transcription initiation And Regulators) provides a wide range of analysis results as well as detailed information of individual samples; ZENBU, a data integration, data processing, and expression enhanced visualization system designed for big data genomics projects, provides interactive way to inspect the entire promoter activities measured in FANTOM5 interactively; and Table Extraction Tool (TET) provides an easy-to-use interface to download subsets of the overall FANTOM5 expression table in an efficient way. A BioMart instance and a track hub for UCSC genome browser enable us to access our TSS resources with widely used interfaces. Furthermore, PRESTo provides an interface to browse human enhancers identified in the FANTOM5 project, and Biolayout enable us to visualize biological states in a three dimensional expression space with interactive interface.

Results: The web resource is accessible from the FANTOM5 portal page: http://fantom.gsc.riken.jp/s.
**P43**

**Rapid and scalable typing of structural variants for disease cohorts**

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**Human Genomics 2016, 10(Suppl 1):P43**

**Objectives:** For studies such as the Alzheimer’s Disease Sequencing Project and the CHARGE Consortium, disease complexity manifests as data heterogeneity, requiring informatics that can additively scale to thousands of samples and analytics that go beyond identifying small variants in capture data. Specifically, whole-genome sequencing (WGS) can assess large, complex genomic variation in both coding and non-coding regions. At scale, the challenge of evaluating these structural variants (SVs) becomes the “N+1” problem of incrementally adding samples without having to perpetually reevaluate petabytes of population read data stored in BAM files.

**Methods:** The BioGraph Analysis Format (BAF) is a novel method of indexing NGS data that allows rapid, at-scale queries by coordinate and genomic sequence. A BAF of HiSeq X 30x WGS data is 8.3 Gb, 95% smaller than the corresponding BAM. Generated from the BAM and genomic sequence, a BAF of HiSeq X 30x WGS data was added. The FANTOM5 resource is continuously updating, for example, the web resource is a gateway to access the large expression atlas in native and admixed Mexican populations

**Results:** To evaluate the BAF, we indexed the HS1011 reference genome sequence with 60x coverage on the HiSeq X. The HS1011 individual has previously been characterized for SV content, identifying ~10,000 high-confidence SVs from ~30,000 putative sites. These sites were BAF-queried in less than 60 seconds. SV-typing these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours. Because these sites across 10,000 HiSeq X WGS samples in BAF would require less than 100 TB, 166 CPU hours, and 2.5 total hours.

**Conclusion:** We will describe how the BAF API allows users to query SV data, how these queries inform SV typing, and how BQF typing performs on a pilot set of 100 30x HiSeq X samples from a disease cohort, providing counts and allele frequencies for multiple SV types.

**Competing interests**

None declared.

**P46**

**Genetic divergence of CYP3A5*3 pharmacogenomic marker for native and admixed Mexican populations**

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**Human Genomics 2016, 10(Suppl 1):P46**

**Objectives:** The frequency distribution of pharmacogenomics markers across the populations in the worldwide can differ in magnitude or be absent depending on the population being assessed. The CYP3A5*3 (rs776746) is the most frequent and well-studied variant allele of CYP3A5. This polymorphism have been related to the guidelines regarding the use of pharmacogenomic tests in dosing of immunosuppressive Tacrolimus drug published in Clinical Pharmacology and Therapeutics by the Clinical Pharmacogenetics Implementation Consortium (CPIC). The CYP3A5*3 genotype is polymorphic in several continental and large populations, for example the minor allele frequency (MAF) of rs776746 marker in Europeans and Africans populations is opposite, while A allele is the minor (0.036) for Europeans G allele is the minor (0.15) in Africans. However, this pharmacogenomic marker is not well-studied in the admixed and Native American populations. The objective in this study is evaluate the distribution of allele frequencies for the CYP3A5*3 variant across the admixed and Native American (Mexican) populations.

**Methods:** Genotypes for rs776746 from the Mexican Genome Diversity Project database were analyzed. Allele frequencies were calculated for 4 Native American-Mexican populations located in north, central and southwest Mexico territory plus 370 Mexican mestizos

**Conclusion:** This is the first study that addresses polymorphism of CYP3A5 and SULT1A1 genes in Sudanese. We provide a reference database of allele frequencies of the genotypes of these genes among Sudanese.

**Competing interests**

None declared.
and 60 European, 60 African, 90 East Asian from the HapMap project. The $F_{st}$ statistic was analyzed as measure of the degree of genetic differentiation pairwise populations.

**Results:** Extreme genetic differentiation among HapMap populations for was observed for $\text{CYP3A5}^*$ (rs776746) as one would expect, particularly for the comparisons of African and European ancestry populations. However, the largest $F_{st}$ statistic and allele frequencies values was found among Native Americans. For example, the allele frequency of $\text{CYP3A5}^*$ allele is more than 6-fold higher in Tepehuanes (0.43) than Zapotecas (0.07) with $F_{st}$ value of 0.1746 (largest divergence).

**Conclusion:** We identified novel CASR variants that have a potential to be related to serum calcium levels in Korean population. Inter-ethnic differences were suggested in some associated SNPs. Given the significant role played by calcium in many diseases and cell signaling, further studies with more East Asian subjects or meta-analyses on them may enable validation of our results and identification of novel genetic loci associated with serum calcium levels.

### Competing interests
None declared.

**Table 3 (abstract P46), Genetic differentiation among Native and admixed Mexican populations**

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<tr>
<th>POP1 Name</th>
<th>POP2 Name</th>
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**P47**

**Whole exome sequence meta-analysis of 13 white blood cell, red blood cell, and platelet traits**

L. Polfus, CHARGE and NHLBI Exome Sequence Project Working Groups
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**Correspondence:** L. Polfus – Human Genetics Center, University of Texas Health Science Center, Houston, USA

**Objective:** Blood cell counts and erythrocyte indices are clinically important indicators of a variety of disease states. The ability to detect and include novel rare variants in gene-based tests is best accomplished via sequencing.

**Methods:** In order to identify novel genes influencing hematologic traits, we analyzed whole exome sequencing data in 15,459 participants of 6 population-based cohort studies for 13 traits in European descent (EU) and African Americans (AA), analyzed together and separately, in fixed-effects meta-analyses. We analyzed variants individually and aggregated functional and loss-of-function (LOF) variants with MAF<5% within genes using SKAT and T5 burden tests. CRISPR/Cas9 genome editing was performed to functionally follow up a replicated novel finding.

**Results:** Discovery phase associations identified 4 novel gene-based results and 3 novel single variants meeting exome-wide Bonferroni corrected thresholds ($P<2.6\times 10^{-6}$ and $P<1.6\times 10^{-7}$). Rare functional variants in $\text{MYOM2}$ were associated with mean corpuscular hemoglobin concentration ($\text{SKAT}_{\text{EU,AA}}=2.2\times 10^{-5}$). In EUs, rare variants in $\text{MRPL43}$ were associated with hemoglobin ($\text{SKAT}_{\text{EU}}=1.2\times 10^{-5}$). LOF variants in $\text{MMACHC}$ were associated with hemoglobin ($\text{SKAT}_{\text{EU}}=1.3\times 10^{-6}$). $\text{MMACHC}$ regulates vitamin B12 metabolism and known variants cause a Mendelian syndrome including anemia. In AAs, rare functional variants in $\text{ACTN4}$ were associated with mean corpuscular volume ($\text{SKAT}_{\text{EU}}=1.5\times 10^{-7}$). $\text{ACTN4}$ variants were previously associated with focal segmental glomerulosclerosis of the kidney. Among the single variant tests, we found low-frequency variant rs9656464 (MAF=0.03, $\text{P}_{\text{EU,AA}}=1.5\times 10^{-5}$) in $\text{AGBL3}$ to be associated with basophil count. A common variant in $\text{CPS1}$ (rs1047891, MAF=0.33, $\text{P}_{\text{EU,AA}}=5.7\times 10^{-8}$) was associated with platelet count. We also identified an association of platelet count with a low-frequency (MAF=0.009) synonymous variant (rs150813342, MAF=0.328, $\text{P}_{\text{EU,AA}}=1.2\times 10^{-6}$) in $\text{GFIB}$, a gene known to cause gray platelet syndrome. In silico algorithms predict rs150813342 to affect exon 5 splicing. Replication in up to 52,024 individuals were significantly associated with lower platelet count for CPS1 rs1047891 (MAYAS+AA=0.328, $\text{P}=1.02\times 10^{-4}$) and $\text{GFIB}$ rs150813342 (P=5.71x10^{-21}).

**Conclusion:** In this exome sequencing study, we identified a novel low frequency locus. We demonstrate an alternative splicing mechanism by which the $\text{GFIB}$ rs150813342 variant suppresses formation of a GFIB isoform that preferentially promotes megakaryocyte differentiation and platelet production.

### Competing interests
None declared.

**P48**

**Association of adipon gene with type 2 diabetes and related phenotypes in african american men and women:**

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**Human Genomics 2016, 10(Suppl 1):P48**
Objectives: African Americans experience disproportionately higher prevalence of type 2 diabetes and related risk factors. Little research has been done on the association of ADIPOQ gene on type 2 diabetes, plasma adiponectin, blood glucose, HOMA-IR and body mass index (BMI) in African Americans. The objective of our research was to assess such associations with selected SNPs. The study included a sample of 3,020 men and women from the Jackson Heart Study who had ADIPOQ genotyping information.

Methods: Unadjusted and adjusted regression models with covariates were used with type 2 diabetes and related phenotypes as the outcome stratified by sex. There was no association between selected ADIPOQ SNPs with type 2 diabetes, blood glucose, or BMI in men or women.

Results: There was a significant association between variant rs16861205 and lower adiponectin in women with minor allele A in the fully adjusted model (βSE) p = -0.13(0.05), 0.003. There was also a significant association with variant rs7627128 and lower HOMA-IR among men with minor allele A in the fully adjusted model (βSE) p = -0.74(0.20), 0.0002.

Conclusion: These findings represent new insights regarding the association of ADIPOQ gene and type 2 diabetes and related phenotypes in African American men and women.

Competing interests
None declared.

P49
Common variants in casr gene are associated with serum calcium levels in Koreans
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Human Genomics 2016, 10(Suppl 1):P49

Objectives: Calcium is a universal intracellular messenger that has an important role in controlling various cellular processes. In this study, we attempt to evaluate the genetic polymorphisms that affect serum calcium levels in Korean population through a two-stage genome-wide association study with the sample of 8642 unrelated Koreans (4558 for discovery and 4093 for replication).

Methods: Study subjects were selected from an ongoing population-based study known as the Korean Genome and Epidemiology Study (KOGES) and genotyped using the Affymetrix Genome-Wide Human SNP Array 5.0. We applied standard quality control parameters such as SNP call rate >95%, minor allele frequency >5% and HWE P>0.001. After this quality control process, genotypes of 4558 individuals for 1219546 autosomal SNPs were used for stage 1 association analysis.

Results: Using SNP arrays, we discovered 963 associated SNPs in stage 1, and replicated 105 SNPs among them in stage 2. We examined them in a combined set of stage 1 and 2 samples and observed that 65 SNPs were significantly associated with serum calcium levels. Among them, rs13068893 in the CASR gene showed the strongest significance (P=3.85x10^-8). Considering the high allele frequency and significance level of the rs13068893 C>G in the CASR gene, this SNP may have a key role in regulating the serum calcium level. We also successfully replicated the four loci (CASR, CSTA, DGKD and GCKR) using our data set that have been previously reported to be significantly associated with calcium levels in Europeans and Indians.

Conclusion: In this exome sequencing study, we identified a novel low frequency loci. We demonstrate an alternative splicing mechanism by which the GFI1B rs150813342 variant suppresses formation of a GFI1B isoform that preferentially promotes megakaryocyte differentiation and platelet production.

References

Competing interests
None declared.

P50
Inference of multiple-wave population admixture by modeling decay of linkage disequilibrium with multiple exponential functions
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Objectives: To infer the histories of complex admixture, two important challenges stand out with methods based on the admixture-introduced linkage disequilibrium (LD): getting rid of the effect of confounding LD (CLD) brought by source populations and fitting LD decay induced by admixture. In previous studies, the decay curve of weighted LD between pairs of sites whose genetic distance were bigger than a certain starting distance was fitted by single or multiple exponential functions, for the inference of single- or multiple-wave of admixture.

Methods: In this study, we developed a new LD based algorithm, named MALDmef, to date the multiple-wave admixtures. Different from previous software, MALDmef takes advantage of derived source populations to reduce the effect of CLD and fits the remaining weighted LD decay curves with hundreds of exponential functions.

Results: The performance of MALDmef was evaluated by simulation and it was shown to be more accurate than MALDER, a state-of-the-art LD based method that was recently developed for similar purposes, under various admixture models. We further applied MALDmef to analyzing genome-wide data from the Human Genome Diversity Project (HGDP) and the HapMap Project. Interestingly, we were able to identify more than one admixture events in several populations, which have yet to be reported. For example, two major admixture events were identified in Xinjiang Uyghur, occurring around 27–30 generations ago and 182–195 generations ago, respectively. In an African population (MKK), three recent major admixture occurring 13–16, 50–67, and 107–139 generations ago were detected.

Conclusion: Our method is a considerable improvement over other current methods and further facilitates the inference of the histories of complex population admixtures.

References
1. Ying Zhou, Kai Yuan, Yaoliang Yu, Xumin Ni, Pengtao Xie, Eric P Xing, Shuhua Xu*. Inference of multiple-wave population admixture by modeling decay of linkage disequilibrium with multiple exponential functions. doi: http://dx.doi.org/10.1101/026757

Competing interests
None declared.

P51
A Bayesian framework for generalized linear mixed models in genome-wide association studies
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Human Genomics 2016, 10(Suppl 1):P51

Objectives: Recent technical and methodological advances have greatly expanded genome-wide association studies (GWAS). The advent of low-cost whole-genome sequencing facilitates high-resolution variant identification, and the development of linear mixed models (LMM) allows improved identification of putatively causal variants. While essential for correcting false positive associations due to population
stratification, LMMs have been restricted to numerical variables. However, phenotypic traits in association studies are often categorical, coded as binary case–control or ordered variables describing disease stages. Furthermore, optimally integrating the results of prior studies remains a methodological challenge.

Methods: To address these issues, we have devised a method for genomie association studies that implements a generalized linear mixed model (GLMM) in a Bayesian framework, called Bayes-GLMM. Bayes-GLMM has four major features: support of categorical variables; cohesive integration of previous GWAS results for related traits by Bayesian modeling; correction for sample relatedness by mixed modeling; and model estimation by both MCMC sampling and maximal likelihood estimation.

Results: To demonstrate our method, we applied Bayes-GLMM to the whole-genome sequencing cohort in the Alzheimer’s Disease Sequencing Project (ADSP). This study contains 576 individuals distributed across 111 families, each with Alzheimer’s disease diagnosed at four confidence levels. The profound population structure in these data required a mixed model approach, and the categorical trait necessitated a generalized model.

Conclusion: In summary, this work provides the first implementation of a flexible, generalized mixed model approach in a Bayesian framework.

Competing interests
None declared.

PS2
Targeted sequencing approach for the identification of the genetic causes of hereditary hearing impairment
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Human Genomics 2016, 10(Suppl 1):PS2

Objectives: Hearing loss is one of the most common afflictions in the world affecting about one in every 1000 newborns (Petersen and Willems 2006). Genetic factors are estimated to be the underlining cause of more than half of the hearing loss cases. The majority of hereditary hearing loss cases is not associated with syndromes (nonsyndromic hearing loss; NSHL) which can be transmitted in an autosomal recessive, autosomal dominant or X-linked modes of inheritance (Petersen and Willems 2006). At least half of the hereditary NSHL cases are caused by mutations in the GJB2/CONNEXIN26 gene (Hereditary Hearing Loss Home Page, http://webh01.ua.ac.be/ hhhl/). However, the contribution of GJB2 mutation to NSHL in the Kingdom of Saudi Arabia is minimal (Al-Qahtani et al. 2009) where the rate of children affected with sensineural hearing loss was estimated to be approximately 26 children out of 1000 (Bafaeche et al. 1994). NSHL can be caused by mutations affecting any one of over 80 deafness loci identified so far making NSHL a very heterogenous trait and complicates diagnosis and genetic counseling (Hilgert et al. 2009).

Methods: The recent advancements in targeted sequencing technologies have made it feasible to sequence multiple genes at a reasonably low cost. Therefore we have designed a targeted sequencing panel using the Ampliseq technology to amplify and sequence 84 genes known to cause NSHL. Genes were selected and custom primers were designed and manufactured through the Ampliseq portal (http://ampliseq.com).

Results: The design resulted in a coverage of 97.42% generating 2697 amplicons with a size range of 125–275 bp in two pools and generating 500.44 kb of DNA sequence.

Conclusion: This panel, which we termed OtoScan will be a useful front line genetic screening tool that will speed up the identification of many genetic causes of hereditary deafness in the Kingdom of Saudi Arabia.

Competing interests
None declared.

PS3
Identification of enhancer sequences by ATAC-seq open chromatin profiling
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Human Genomics 2016, 10(Suppl 1):PS3

Objectives: Enhancers are cis-regulatory elements that regulate gene expression in the control of cell type-specific functions, developmental fate and evolution. Recent studies revealed the importance of enhancer misregulation for the pathogenesis of certain diseases including cancer through point mutations in either regulatory elements or factors modulating enhancer-promoter communication. Enhancers are often located far from their target gene promoters within the noncoding genome, which makes their identification challenging. Using statistical models, such as ChromHMM, we can infer enhancer locations by studying multiple histone modifications obtained by the ChIP-seq technology. However, ChIP-seq requires large cell numbers (~10M), which is not compatible with clinical samples. The Assay for Transposase Accessible Chromatin (ATAC-seq) is an alternative assay to generate chromatin accessibility maps from 500–5,000 cells, making this an ideal approach to study enhancers in clinical samples. Here we developed a machine-learning model to infer the entire repertoire of enhancers from ATAC-seq open chromatin profiles.

Methods: We conduct our analysis on ATAC-seq data generated from GM12878 human lymphoblastoid cell line and CD4+ T-cells. From these samples, we extracted i) ATAC-seq features, e.g., peak strength; ii) DNA sequence based features, e.g., conservation scores; and iii) TF binding features using footprinting algorithms and compiled a comprehensive dataset. We applied Random Forest algorithm on this data to predict reference enhancers obtained from ChromHMM models.

Results: Our final data include ~60K ATAC-seq peaks and 490 features and a class indicating chromatin state obtained via ChromHMM. Random Forest based model predicted enhancer sequences with 74.8% sensitivity and 73.8% specificity. We also showed that the most important features for accurately predicting enhancer sequences include ATAC-seq based features and certain TF binding profiles.

Conclusion: The present study demonstrated that prediction of regulatory elements using ATAC-seq data alone is comparable to ChromHMM predictions. It is noteworthy that ChromHMM states are also predictions and the actual performance of ATAC-seq based model needs to be tested on experimentally validated enhancers. Our future work includes comparing our predictions with known enhancers from public databases such as FANTOMS and Vista enhancers.

Competing interests
None declared.

PS4
Direct enrichment for the rapid preparation of targeted NGS libraries
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Human Genomics 2016, 10(Suppl 1):PS4

Objectives: Target enrichment, coupled with next-generation sequencing, enables the interrogation of specific targets of interest at a level of sensitivity that is typically cost-prohibitive with whole genome sequencing. However, the drawbacks to target enrichment generally include longer and more complex library preparation workflows or a loss of the ability to detect PCR duplicates. Here we present DIRECT target
enrichment, a unique and highly specific method for hybridization-based target enrichment that incorporates the use of unique molecular indexes in a simple, one-day protocol.

Methods: DNA was isolated from fresh frozen tissue, FFPE samples, and liquid biopsies followed by enrichment of 36 kb of cancer-related targets with the DIRECT enrichment method. Combined with unique molecule barcoding and Illumina sequencing, variants were detected and frequencies reported using common variant callers.

Results: Application of DIRECT target enrichment across different DNA sample types show similar results, with no loss in specificity or sensitivity across DNA isolated from different sources. Somatic variations can be detected with high sensitivity using as little as 50 ng of input DNA, and libraries can be made with as little as 10 ng of starting material for the detection of genetic variations.

Conclusion: DIRECT target enrichment enables the highly specific capture of regions of interest within 7 hours and with low amounts of starting material. As a result, this method can easily be applied to a variety of applications for which time and sample amount may be limiting.

Competing interests
None declared.

P56
Performance of the Agilent D5000 and High Sensitivity D5000 ScreenTape assays for the Agilent 4200 TapeStation System
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Objectives: This Poster focuses on the performance of both D5000 ScreenTape assays with respect to the accuracy and precision of quantification and sizing, as well as the sensitivity of these assays. Data analysis for quantification and molarity determination was compared against the corresponding assay for the Agilent 2100 Bioanalyzer system. Additionally, performance of both the D5000 and High Sensitivity D5000 assays on the 4200 TapeStation was compared to the 2200 TapeStation system.

Methods: The 4200 and 2200 TapeStation systems were used with the D5000 and High Sensitivity D5000 ScreenTape assays.

Results: Data is presented that prove the specifications of sensitivity, reproducibility, accuracy and linearity of the new D5000 assay for the Agilent 4200 TapeStation system.

Conclusion: This Poster shows that the Agilent D5000 ScreenTape and Agilent High Sensitivity D5000 ScreenTape assays for the Agilent 4200 TapeStation system provide highly accurate and reproducible sizing and quantification of DNA fragments ranging from 100 to 5,000 bp. Furthermore, it demonstrates that the assays can also be applied to determine the DNA average region size, molarity, and concentration of distributed DNA smears. The sizing, concentration, and molarity results highly correlate with the data obtained from equivalent assays on the Agilent 2100 Bioanalyzer system. In addition, D5000 and High Sensitivity D5000 ScreenTape assays show equivalent performance in the 4200 and Agilent 2200 TapeStation systems.

Competing interests

P57
ClinVar: a multi-source archive for variant interpretation
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Objectives: ClinVar archives submitted interpretations of the medical importance of human variants, standardizes content, and aggregates data from multiple sources to facilitate peer review and track revisions.

Methods: Interpretations are provided by clinical testing laboratories, research laboratories, OMIM®, GeneReviews®, locus-specific databases, expert panels and organizations that provide practice guidelines. ClinVar integrates those data with information from other NCBI databases such as dbSNP, dbVar, Gene, and MedGen.

Results: The ClinVar website can be searched with gene symbols, variant names including HGVS expressions, rs numbers, and diseases or phenotypes. Searches with many results may be narrowed with filters, such as restricting to a clinical significance, type of variant, and more. Searching by location is facilitated by Variation Viewer, a browser to view all NCBI variation data in a region of interest, such as a region of structural variation. Each row of the search results table in ClinVar links to a page for that variant(s), which has two major sections: aggregate data and submitter-specific data. The aggregate data section provides a summary of information about the interpreted variant(s). The summary includes the review status for the variant(s), represented with a description and a number of gold stars, and a variation identifier, which is a unique identifier that represents the variant(s) that were interpreted. The summary also displays an overall clinical significance, which is calculated based on interpretations provided by submitters; ClinVar staff do not curate or review clinical significance. A list of conditions for which the variant was interpreted is provided, as well as general information about the variant including links to other public databases. The submitter-specific section includes details of the assertions of clinical significance and evidence provided by each submitter. Evidence may include citations, counts of individuals with the variant, free text descriptions of the evidence, and more. Data is updated only by the submitter; thus maintenance of an interpretation and the supporting evidence is the responsibility of each submitter.

Conclusion: Those with new evidence and interpretations are encouraged to contribute their own data to improve content for all. An online Submission Portal is available to upload data to ClinVar, as a single submission or in batch. ClinVar provides monthly releases available via FTP, and API to access specific records.

Competing interests
None declared.

P59
Association of functional variants and protein physical interactions of human MUTY homolog linked with familial adenomatous polyposis and colorectal cancer syndrome
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Human Genomics 2016, 10(Suppl 1):P59

Objectives: To the best of our knowledge, this is the first report confirming the structural and functional association of MUTYH mutations (Y165C; G382D; P54S; A22V; Q63R; G45D; S136P and N43S) through in-silico approach.

Methods: We retrieved clinical records of MUTYH for multiple adenomas, test unit code 84304 from the Mayo Medical Laboratory, USA. The nonsynonymous single nucleotide polymorphisms (nsSNPs) may alter the structure or function of expressed proteins, and therefore have an effect on disease outcome. In order to evaluate the phenotypic effects of nsSNPs in human DNA repair genes, we studied the polymorphism in terms of various functional properties. The molecular origin of disease-related phenotypes was caused by six mutations in MUTYH, and that were associated with hereditary predisposition of colon and stomach cancer syndrome of FAP.

Results: We confirmed that MAP was inherited in an autosomal recessive pattern due to the mutations in MUTYH (P54S, A22V, Q63R, G45D, S136P and N43S), hence both copies of the gene become inactivated during disease occurrence. The parents of an individual with an
autosomal recessive condition may serve as carriers, each harboring one copy of the mutated gene without showing signs and symptoms of MAP. The MUTYH protein interacts with six partners, but only four of these proteins showed direct physical interactions in our study. These proteins were mMSH6, hPCNA, hRPA1, and hAPEX1. We also for the first time examined specific interactions of these protein partners with MAP associated MUTYH mutants using molecular dynamics simulations. These approaches provided tools for exploration of the conformational energy landscape accessible to these protein partners. The study also determined the impact before and after energy minimization of protein-protein interaction (PPI) and binding affinities of MUTYH wild type and mutant forms, including interactions with other proteins. Taken together, this study has provided innovative insights into the role of MUTYH and its interacting proteins in MAP.

Conclusion: This study provides interesting information and will open up a fresh avenue for the FAP researchers. We also strongly believe that the identified important features of the HMUTYH mutations through our In-silico study will support further in vitro studies in future.

Competing interests
None declared.

P61

Compound heterozygous mutation in the LDLR gene in Saudi patients suffering severe hypercholesterolemia

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Human Genomics 2016, 10(Suppl 1):P61

Objectives: Familial hypercholesterolemia (FH) is most commonly caused by variants in the LDL receptor (LDLR) gene. Herein, we describe a severely affected FH proband and their first-degree blood relatives, the proband was resistant to statin therapy and was managed on an apheresis program.

Methods: Genetic screening for the LDLR variants was performed by exon sequencing analysis. Using a bioinformatics approach we theoretically explored the putative structure of the wild type and mutant LDLR protein and the implication of structural changes on the possibility of LDLR dimer formation.

Results: We identified a compound heterozygous mutation with missense c.1731G>T, p.(W577C) and frameshift c.2027delG, p.(G676Af*33) variants at exons 12 and 14 of the LDLR gene respectively in a proband of a Saudi family. DNA sequencing of LDLR from the parents demonstrated that the missense variant was inherited from the mother and frameshift variant was inherited from the father. The frameshift variant resulted in a stop signal 33 codons downstream of the deletion, which most likely lead to a truncated protein that lacks important functional domains, including the transmembrane domain and the cytoplasmic tail domain. The missense variant is also predicted to be likely pathogenic and affect EGF-precursor homology domain of the LDLR protein. The segregation pattern of the variants is consistent with the lipid profile, suggesting a more severe FH phenotype when the variants are in the compound heterozygous state. We also describe a three dimensional homology model of LDLR structure and examine the consequence of the missense and frameshift variants, as this could affect the LDLR structure in a region involved in dimer formation, and protein stability.

Conclusion: The finding of a compound heterozygous mutation causing severe FH phenotype is important for the genotype-phenotype correlation. Furthermore, provided bioinformatics based structural analysis of the mutant protein and its molecular consequences on the physiology of FH that could facilitate developing mechanistic models of LDLR function. These results also enlarge the spectrum of FH-causative LDLR variants in the Arab population, including the Saudi population.

Disclosure of interest
None declared.