Abstracts

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Guest Editors
Jaana Hartiala, Los Angeles, CA
J. Alfredo Martinez, Navarra
Zhaoping Li, Los Angeles, CA
Hooman Allayee, Los Angeles, CA

Contents

Oral Presentations
Abstracts O1–O5

Poster Presentations
Abstracts P5–P66

Author Index

94

96

124
Oral Presentations

O1

Metabolic Responses to Dietary Patterns Differ with Genetic Backgrounds in Mice

William Barrington1, Carolina Mantilla Rojas2, Selene Howe1, Brian Bennett3, Daniel Pomp4, David Threadgill2

1University of California Los Angeles, Lusis Lab, Medicine-Cardiology, Los Angeles, CA; 2Texas A&M University, Molecular and Cellular Medicine, College Station, TX; 3United States Department of Agriculture, Agricultural Research Service, Davis, CA; 4University of North Carolina Chapel Hill, Genetics, Chapel Hill, NC, USA

Dietary patterns have been shown to have profound effects on health when studied at the population level. For example, Japanese and Mediterranean diets are associated with longevity and low rates of chronic disease. The American diet is associated with increased risk of heart disease and certain types of cancer. However, studies evaluating dietary interventions in individuals find significant variations in responses. Much of this variation is likely due to underlying genetic differences among individuals. To determine how diet affects cardiometabolic health in different genetic backgrounds, we compared the health effects of an American diet to a standard mouse diet in each sex of four inbred mouse strains (A/J, C57BL/6J, FVB/NJ, and NOD/ShiltJ) selected for their known susceptibilities to cancer, diabetes, and other diseases. Twenty mice per strain were fed each diet for six months while undergoing a range of clinical analyses. All strains had negative health effects when fed the American diet, however the severity of effects varied widely by strain with the most severe effects in C57BL/6J and the mildest effects in A/J. We then investigated the health responses in the same strains fed alternative human-relevant diets including Mediterranean, Japanese, and ketogenic diets. Many diet responses were dependent upon genetic background including: adiposity, glucose tolerance, blood chemistry profiles, liver triglyceride storage, and metabolic rate. The severity and directionality of many of these diet responses differed depending on the genetic background. For example, C57BL/6J maintained good health on a ketogenic diet; FVB/NJ mice became obese and showed signs of cardiometabolic distress on the same diet. Most strains had beneficial effects from the Japanese diet, although abnormal liver effects were identified in A/J mice. The Mediterranean diet failed to improve body composition in most strains but improved blood lipids. Mouse strains differed as to which diets were optimal or suboptimal, indicating individuals likely have a specific diet for optimal health based upon their unique genetic makeup. The results emphasize the need to evaluate dietary efficacy at an individual level.

O2

Fasting Serum Lipid and Dehydroepiandrosterone Sulfate as Important Metabolites for Detecting Isolated Postchallenge Diabetes: Serum Metabolomics via Ultra-High-Performance LC-MS

Changhao Sun1, Ying Li1, Xiaoyan Wu1

1Department of Nutrition and Food Hygiene, Public Health College, Harbin Medical University, Harbin, P R China

Background: Isolated postchallenge diabetes (IPD), a subtype of type 2 diabetes mellitus (T2DM) defined as 2-h postprandial plasma glucose ≥200 mg/dL (≥11.1 mmol/L) and fasting plasma glucose (FPG) <108 mg/dL (<6.0 mmol/L), is often overlooked during screening for diabetes on the basis of FPG concentrations. A key challenge is early identification of IPD by the use of fasting serum, which is critical for large-scale diabetes screening.

Methods: We applied a nontargeted metabolomic approach using ultra-high-performance liquid chromatography–quadrupole TOF–mass spectrometry (UPLC-QTOF-MS) to analyze serum samples from 51 patients with IPD, 52 with newly diagnosed T2DM, and 49 healthy individuals. We processed metabolite profiles by multivariate analysis to identify potential metabolites, which were further confirmed by tandem MS (MS/MS). We also used GC-MS and ELISA methods to detect potentially important metabolites. A number of independent samples were selected to validate the identified candidates.

Results: We selected 15 metabolites with a view to distinguishing patients with IPD, whereas 11 were identified with an authentic standard. The selected metabolites included linoleic acid, oleic acid, phospholipids, and dehydroepiandrosterone sulfate (DHEA-S). In IPD samples, significantly higher linoleic and oleic acid (P < 0.001) and lower DHEA-S (P < 0.001) concentrations were observed, compared with controls. The area under the curve from a combination of linoleic acid, oleic acid, and DHEA-S in the validation study was 0.849 for the IPD group.
Conclusions: The current study provides useful information to bridge the gaps in our understanding of the metabolic alterations associated with IPD and might facilitate the characterization of patients with IPD by the use of fasting serum.

O3 Evidence for a Genetic Association between High Milk Consumption and Cardio-Metabolic Traits

Vimal Karani,1, Alana Cavadino,2,3, Ang Zhou,4 Julie A. Lovegrove,3 GIANT consortium, Global Lipids Consortium, MAGIC consortium, DIAGRAM consortium, CARDIOGRAM consortium and ICBP consortium, Elna Hypponen2,4

1Hugh Sinclair Unit of Human Nutrition, University of Reading, UK; 2Population, Policy and Practice, UCL Institute of Child Health, UK; 3Wolfson Institute of Preventive Medicine, Queen Mary University of London, UK; 4Centre for Population Health Research, University of South Australia, Adelaide, Australia

Studies have shown associations between high milk intake and cardio-metabolic traits. We conducted a genetic study to obtain evidence for the causal relationship between milk consumption and cardio-metabolic traits using the lactase persistence (LCT-13910 C>T, rs4988235) variant as an instrumental variable in a Mendelian Randomization analysis. We tested the association of LCT genotype with milk consumption (for validation) and with cardio-metabolic traits (for a possible causal association) in three large-scale population studies, with up to 87,790 participants (the 1958 British Birth Cohort (N=5,231), the Health and Retirement study (N=8,520) and the UK Biobank (N=73,590)). The findings from our analyses were confirmed using summary statistics from consortia meta-analyses on intermediate traits [Genetic Investigation of Anthropometric Traits (GIANT, N=327,665), Global Lipids Consortium (N=188,577), the Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC, N=229,233) and International consortium for Blood Pressure (N=146,581)] and extended to cover disease end-points [DIAGRAM (34,840 cases and 114,981 controls) and CARDIOGRAM (22,233 cases and 64,762 controls)]. Carriers of the ‘T’ allele of the LCT variant were more likely to consume milk (P=3.0x10^{-6}). In meta-analysis including all three studies, under a recessive model, ‘T’ allele was associated with higher body mass index (BMI), lower low-density lipoprotein (LDL-C) and total (TC) cholesterol levels. Large intervention trials are needed to establish the causal link between high milk consumption and cardio-metabolic phenotypes before changes in dairy consumption could be recommended for the prevention of cardio-metabolic diseases.

O4 Effects of Prebiotic Fiber Xylo-oligosaccharides in Adenine Induced Nephropathy in Mice

Jieping Yang1, Qing Li1, Susanne M. Henning3, Zhong Jin4, Mark Hsu1, Rupe Lee1, Jianfeng Long1, David Heber1, Zhaoping Li1,*

1Center for Human Nutrition, David Geffen School of Medicine at UCLA, Los Angeles, CA90095, USA; 2Department of Pathology & Laboratory Medicine, David Geffen School of Medicine at UCLA, Los Angeles, CA 90095, USA

Background: There is increasing evidence from epidemiological studies showing that high intakes of dietary fiber are associated with decreased risk of chronic kidney disease (CKD), and that patients with CKD have an altered gut microbiota.

Objective: This study evaluated the effect of prebiotic fiber Xylo-oligosaccharides (XOS) supplementation in kidney function and gut microbiome of mice with adenine-induced CKD.

Methods: Male 7-week old C57BL/6 mice were fed with AIN93M control diet or control diet containing 0.2% adenine (W/W) for 3 weeks to induced CKD. Four mice from control and adenine group were sacrificed after 3 weeks. Control and CKD mice were then placed on control diet, 3% or 10% XOS (W/W) for additional 3 weeks (n=8/group). Mice were sacrificed after 3 weeks, blood urea nitrogen (BUN), creatinine, kidney pathology, gene expression for fibrosis and inflammatory markers were measured. Cecum content was used for microbiota and short chain fatty acids (SCFAs) analysis.

Results: CKD mice consuming the control diet exhibited increased BUN, creatinine, interstitial fibrosis, inflammation and tubular damages. Supplementation with XOS for 3 weeks significantly attenuated the damages and preserved renal function. 16S rRNA gene sequencing of cecum microbiota revealed that alpha diversity was comparable between CKD mice and control mice, and XOS induced dose dependent decreases of alpha diversity in both CKD and control mice. Beta-diversity analysis (unweighted UniFrac) showed that samples clustered according to experimental groups. Linear discriminant analysis identified 9 bacterial genera enriched in CKD mice; 6 of these including Dorea, Clostridium, Lactobacillus, Erysipelotrichaceae, Staphylococcus and Allobaculum were significantly reduced with XOS supplementation. XOS increased cecal SCFAs production. Furthermore, cecal SCFAs were negatively correlated with blood levels of BUN.

Conclusion: Our results showed that XOS supplementation improved kidney function in mice with adenine-induced CKD, which was associated with profound changes in gut microbiome composition and the production SCFAs.
The adverse health effects of obesity and accompanying complications remain a major concern due to the scarcity of relevant progress and suitable tools for overweight prevention and treatment. Precision nutrition is an emerging and innovative therapeutic approach that takes into account not only the individual’s genetic, epigenetic, metagenomic and metabolomic information, but also age, gender, ethnicity, clinical and family history, as well as pathophysiological conditions and treatments or physical activity patterns. In recent years, scientific advances in “omics” sciences are contributing to a better understanding of the role of genetic variants (Nutrigenetics), epigenetic signatures (Nutriepigenetics), gut microbiota composition (Metagenomics), and the metabolome (Metabolomics) in the development of obesity and accompanying complications, and their potential application for therapeutic purposes. For example, interactions between polymorphisms and diet are implicated in disease risks, and also in differential responses to diets restricted in energy or modified in the macronutrient distribution. In this context, studies conducted in several populations have investigated the effects of various SNPs on weight loss, weight regain and reductions in blood lipid levels and insulin resistance. These investigations include SNPs in genes involved in food intake regulation, lipid metabolism, insulin signaling, glucose homeostasis, inflammatory response, and circadian cycle. In addition, experimental studies have investigated the epigenetic mechanisms underlying the beneficial effects of some nutrients and bioactive food components. Based on these evidences, it has been proposed that the introduction of these dietary compounds into an “epigenetic diet” could serve as an effective strategy to reduce the incidence of obesity and associated comorbidities. Also, it has been reported that some epigenetic signatures may be used as biomarkers to follow-up the effect of nutritional treatments on weight loss and improvements in metabolic profiles. Meanwhile, the use of sequencing technologies and metagenomic studies are contributing to the characterization of gut microbiota and the subsequent impact on health. Of particular interest are the effects of dietary carbohydrates and polyphenols on attenuating microbial dysbiosis in the obese through the promotion of “healthy” intestinal bacterial communities for the control of body weight and other metabolic abnormalities related to excessive adiposity. On the other hand, metabolomics focuses on the analysis of changes in the concentration of metabolites in biological samples in order to assess the compliance/adherence to dietary regimens, as well as the identification of new biomarkers related to food intake or disease progression. In this sense, metabolomic analyses are allowing to metabolically categorize individuals depending on dietary intake or nutritional responses. Overall, this scientific knowledge based on “nutri-omics” technologies is enabling the design of personalized strategies for obesity management in the era of precision nutrition.

References:
regular live webinars. Affiliations with food and pharmaceutical industry (e.g. genetic testing companies), professional organisations and research/education institutes were observed.

**Conclusions and Implications:** The training courses predominantly delivered online enable nutrition professionals worldwide to upskill in nutritional genomics and personalised nutrition. As courses do not appear equivalent, those seeking training should scrutinise and compare cost, duration, mode, content, and affiliations of course providers to ensure learning needs are met.

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**P6**  
Potential Role of Abscisic Acid (ABA) in Protecting Beta Cell Function Under Glucolipotoxic Conditions

Fadi Alkayal1, Hossein Arefanian1, Eman Al Shawal1, Sulaiman Marafe1, Ali Tiss1  
1Research Division, Dasman Diabetes Institute, Kuwait

**Background:** In type 2 diabetes, hyperglycemia in combination with high levels of free fatty acids, a condition known as glucolipotoxicity, affects the mass viability and function of beta cells. The phytohormone abscisic acid (ABA) reportedly induces insulin secretion and might help prevent insulin resistance. In plants, ABA prevents cell apoptosis and DNA degradation.

**Aims:** Here, we hypothesized that ABA might have similar protective effects in mammalian beta cells under glucolipotoxic conditions. Thus, we have investigated the effects of ABA on the viability and function of mammalian beta cells treated with high levels of glucose and palmitic acid (PA).

**Methods:** A rat beta cell line (INS-1) was exposed for 14 d to 16 mM glucose with or without 400 μM PA in the presence or absence of 10 nM ABA. Label-free proteomic quantification approaches were used to investigate the abundance and differential expression of proteins under these treatment conditions. Proteomics results were validated using RT-PCR and western blotting. Pathway analysis was carried out using Ingenuity Pathways Analysis (IPA) software. Cell viability under each treatment condition was also assessed.

**Results:** The abundance of key proteins involved in insulin signaling was modulated by ABA in the presence of high levels of glucose and PA. The expression of the transcription factors PDX1, RICTOR, and MAP4K4 was increased under treatment with ABA, while the expression of pathways downstream of ANGPT2 and of pirinixic acid was inhibited under these conditions. Interestingly, canonical signalling pathways such as those involving EIF2, PKA, and the NRF2-mediated Oxidative Stress Response were also clearly modulated by ABA.

**Conclusion:** Specific pathways and proteins modulated by ABA that were identified in the present study will provide insight into the potential beneficial effects of ABA in protecting beta cells against glucolipotoxic conditions and might promote the use of ABA in the prevention and management of diabetes.

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**P7**  
The FTO Gene’s Relationship to Obesity and Type 2 Diabetes

Amylee Amos1  
1Amos Institute, University of Southern California, CA, USA

FTO is the obesity and fat mass associated gene in humans and is colloquially referred to as the “Fat Gene”. Located on chromosome sixteen, FTO contains a series of single nucleotide polymorphisms (SNPs) that have been studied to determine their relationship to adiposity and type two diabetes. Specifically SNP rs9939609, located within the first intron of FTO, has been studied extensively. Research shows that genotype variants of rs9939609 are correlated with increased BMI, increased waist circumference, altered hunger and satiety related hormones, altered eating behaviors, and greater risk for type two diabetes. While the exact mechanism of this SNP is still unknown, research shows that A allele homozygotes are consistently at higher risk for greater adiposity than are heterozygotes or T/T wild types. Many studies have proven that this relationship is independent of glucose levels, hemoglobin A1c, lipid levels, or blood pressure. Additionally, heterozygotes are at increased risk for type two diabetes than are T/T wild types. Much research has been performed to explore potential mediators to these relationships and investigate the mechanism behind the increased disease risk. This literature review analyzed the research of ten studies and the relationship of FTO and obesity and type two diabetes.

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**P8**  
Omega-3 Index Predicts Cognitive Impairment Score in an MTHFR-C677T Dependent Manner

Emma L. Beckett1, Charlotte Martin1, Katrina King2, Suzanne Niblett1, Jessica Ferguson1, Kyle Abbott1, Mark Luccock1, Manohar Garg1, Martin Veysey1,3  
1The University of Newcastle, Australia; 2Hull York Medical School, UK

**Background:** B vitamins, plasma homocysteine and omega-3 polyunsaturated fatty acids (Omega-3PUFA) have independently been found to modulate risk for cognitive decline, however results have been mixed. MTHFR-C677T polymorphisms, which alter B vitamin related one carbon metabolism, have also been independently linked to risk for cognitive impairment. Therefore, we investigated the interactions between omega-3 PUFA and MTHFR-C677T genotype in predicting risk for cognitive impairment.

**Methods:** This study is a secondary analysis of the Retirement Health and Lifestyle Study cohort (an elderly cohort from NSW, Australia). Participants completed the Mini–Mental State Examination (MMSE), and gave blood for assessment of serum folate, B12, homocysteine, omega-3 index (O3I) and MTHFR-C677T genotype (RFLP-PCR). Age, sex, education, income and smoking history were also recorded. Complete data sets were available for 555 individuals (55.8% female).
Results: 12.1% of participants had MMSE scores indicative of cognitive impairment. 44.3% were homozygous for the MTHFR-C677T ancestral allele (CC). No parameters varied by allelic carriage, except plasma homocysteine, with higher levels in the subgroup carrying the T allele.

When all subjects were analysed together (adjusted for serum folate and B_{12}, sex, socioeconomic variables and smoking history) O3I was a significant predictor of a normal MMSE result (β = 0.18, p = 0.02, R^2 = 0.12, \textit{p}_{\text{interaction}} = 0.0006). Odds ratio analysis revealed an increased likelihood of a normal MMSE result with each unit rise in O3I [1.19 (1.03-1.38)]

When analysis was stratified by MTHFR-C677T polymorphic allele carriage (CC vs CT/TT), results were only significant in those carrying the T variant. There was a significant interaction between MTHFR-C677T allelic carriage and O3I in predicting a normal MMSE score (\textit{p}_{\text{interaction}} = 0.02). A significant interaction was also detected for O3I and serum folate (\textit{p}_{\text{interaction}} = 0.04). There was no interaction between O3I and, sex (\textit{p}_{\text{interaction}} = 0.9), age (\textit{p}_{\text{interaction}} = 0.2) or serum B_{12} (\textit{p}_{\text{interaction}} = 0.7). Results were similar when adjusted for plasma homocysteine.

Conclusions: O3I predicts cognitive impairment only in those possessing the MTHFR-C677T polymorphic variant. This variant reduces MTHFR enzyme activity, and may impair B vitamin related one carbon metabolism. Omega-3PUFA may therefore only be beneficial in preventing cognitive impairment in those who have impaired B vitamin metabolism or low folate status.

P9

Rs7903146 and Its Association with Type 2 Diabetes Mellitus

Dina Ben-Nissan

1 USC Leonard Davis School of Gerontology, CA, USA

TCF7L2 is among the highly replicated genes presenting a strong association with type 2 diabetes mellitus (T2DM). The single nucleotide polymorphism (SNP) rs7903146 (C/T) on the intron 4 of the TCF7L2 gene has been studied extensively for its link with T2DM risk. In this literature review, ten studies were evaluated in order to understand the relationship between rs7903146 and T2DM. Many studies suggested that the T-allele is indeed related to increased diabetes risk, where people with the homozygous TT allele genotype combination have highest genetic risk for developing this chronic disease. Some of the studies attribute the diabetes prevalence to less incretin secretions, while others attribute it to the beta-cell functionality or decreased insulin secretion. Nonetheless, the true mechanism remains uncertain. This SNP was assessed across varying populations, and was found to have consistent results cross-culturally. Additionally, it was noted that it is one of the strongest SNP’s in regard to T2DM risk. As expressed by many of the studies, the T-allele resulted in a disadvantageous trait for those individuals.

P10

Chemical Stress Induces Up-Regulation of Nurr1: Role of Antioxidants

Laura Bordoni1, Donatella Fedeli2, Cinzia Nasuti2, Dennis Fiorini3, Luana Quassinti2, Massimo Bramucci2, Rosita Gabbianelli1

1 School of Advanced Studies, 2 School of Pharmacy and 3 School of Science and Technology, University of Camerino, Italy

Objectives: The nuclear receptor NR4A2 (Nurr1) has been demonstrated to be able to modulate cellular homeostasis, energy metabolism, inflammatory response and dopaminergic system. Nurr1 is induced by stress and injury in brain, and it is necessary for maintenance of midbrain dopaminergic neurons because it regulates the dopaminergic synthesis. Down-regulation of Nurr1 has been observed together with a reduced expression of tyrosine hydroxylase (TH) and an increased level of α-synuclein in Parkinson’s disease. Finally, a key role in the protection against oxidative and inflammatory stress associated to neurodegeneration has been associated to Nurr1.

The aim of this study was to determine the role of antioxidants in modulating the up-regulation of Nurr1 due to chemical stressor residues in food.

Methods: PC12 dopaminergic cell line has been incubated with the lipophilic chemical stressor permethrin (Perm) in corn oil, for 72h with various antioxidants or antioxidants vehicles, like glutathione (GSH), tocotrienols, Extra Virgin Olive oil (EVO) and Electrolyzed Reduced Water (ERW). qPCR of Nurr1 was used to screen the protective effect of the analyzed antioxidants. Corn oil and EVO were analyzed as vehicle for their impact on Nurr1, Nrf2 and NFkB gene expression. Western Blotting analysis of Nurr1 and TH was performed as confirm where data showed interesting evidences. Trypan blue was used to count cells and determine cell viability.

Results: Cell viability, reduced in dopaminergic cell line exposed to Perm as chemical stressor, resulted to be improved only by EVO due to its redox and anti-inflammatory capacity, in agreement with its ability to promote Nrf2 and to inhibit NF-kB gene expression; moreover, the two oils, EVO and corn oil, modulated Nurr1 and TH similarly. GSH, tocotrienols and ERW could reduce the up-regulation of Nurr1 due to chemical stress, although ERW was more efficient.

Conclusion: EVO inhibits NF-kB and protects PC12 improving cell viability reduced by Perm. ERW was able to counterbalance the up-regulation of Nurr1 and TH, thus they could be helpful as a dietary tool in the prevention of stressor-induced-dopaminergic neurodegeneration.
**P11**  
Vitamin E intake and Pro12Ala Polymorphism of PPARγ Gene Interact with Adiponectin Levels  
Wendy Campos-Pérez1, Livier Torres-Castillo1, Mariana Pérez-Robles1, Erika Martínez-López2  
1Medical Molecular Biology Service, “Fray Antonio Alcalde” Civil Hospital of Guadalajara, Department of Molecular Biology and Genomics, University Center of Health Sciences, University of Guadalajara, Guadalajara, Jalisco, México  

**Background/Aim:** The importance of dietary micronutrients intake relies on their many biological functions and high impact on health; unfortunately, micronutrients intake, like vitamin E, is deficient in many populations including countries of southern Asia, United States and México. One of the beneficial effects associated with vitamin E intake is the enhancing of PPARγ activity and its consequently upregulation of adiponectin expression, nonetheless, there is no reported data about the role of the vitamin E intake on adiponectin levels between PPARγ genotypes. The aim of this study was to analyze the adiponectin levels in subjects with the Pro12Ala polymorphism of PPARγ according to vitamin E intake.  

**Methods:** A total of 283 subjects were enrolled. Habitual dietary intake was estimated by a validated three-day food consumption record. Total vitamin E intake, either from diet or supplements, was analyzed using the Nutritionist Pro™ Diet Analysis software. Genomic DNA was extracted from peripheral leukocytes. The Pro12Ala polymorphism (rs1801282) of PPARγ was determined by allelic discrimination. Analysis of covariance with Bonferroni post-hoc multiple comparison tests were used to analyzed the interaction between genotypes and vitamin E intake with adiponectin levels.  

**Results:** The 68.9% were woman, the mean age and BMI of studied population was 37.6 ± 11.4 years and 27.4 ± 7.2 kg/m², respectively. Vitamin E intake was deficient in all subjects with a mean of 1.50 ± 1.78 mg/day. When serum adiponectin levels were analyzed based on vitamin E intake among genotypes, it was observed that subjects with higher vitamin E intake and Pro12Ala/Ala12Ala genotype, had statistically significant higher levels of serum adiponectin than subjects with the Pro12Pro genotype (4.4 [3.2-5.7] versus 2.7 [2.0-3.5] μg/mL, p=0.027).  

**Conclusions:** Our results suggest that may be necessary to promote a greater consumption of vitamin E to achieve the intake recommendations that are at least 10 mg/day, especially in subjects with Pro12Pro genotype because they, with lower vitamin E consumption, might be at higher risk to develop metabolic complications.

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**P12**  
Blackberry (rubus spp) and Grumixama (eugenia brasiliensis lam) Ellagitannins and Anthocyanins-Enriched Extracts Effects on Growth and mRNA and miRNA Expression in MDA-MB-231 Human Breast Cancer Cells  
Gabriela Rezende Costa1,2,*, Tatiana N. Pedrosa3, Renata C. Albuquerque2, Silvia Stuchi Maria-Engler3, Franco Maria Cajolo1,2, Neuza Maria Aymato Hassimotto1,2, Thomas Prates Ong1,2  
1Food Research Center (FoRC); 2Department of Food and Experimental Nutrition, and 3Department of Clinical and Toxicological Analyses, Faculty of Pharmaceutical Sciences, University of São Paulo, Brazil  

**Introduction:** Breast cancer is the most prevalent cancer in women. Anthocyanins and ellagitannins found in berries represent promising breast carcinogenesis inhibitory agents. Grumixama (Eugenia brasiliensis Lam), a cherry species from Brazil, and blackberry (Rubus spp) are important sources of anthocyanins and ellagitannins. The aim of this study was to evaluate the effects of grumixama and blackberry ellagitannins and anthocyanins-enriched extracts on growth and RNAs and miRNAs expression in human breast cancer cells.  

**Methods:** MDA-MB-231 (hormone receptors negative) cells were treated with grumixama or blackberry ellagitannins or anthocyanins-enriched extracts. Cell growth analysis (cell cycle, apoptosis and proliferation) was performed. Breast cancer- associated RNAs (84 genes) and miRNAs (84 genes) expression was further evaluated by real time PCR.  

**Results:** Grumixama anthocyanins-enriched extract induced cell cycle arrest at G0/G1 in MDA-MB-231 cells. Grumixama and blackberry ellagitannins-enriched extracts also induced cell cycle arrest in G0/G1 and increased cells in subG0. Grumixama and blackberry ellagitannins but not anthocyanins-enriched extracts induced apoptosis. All extracts inhibited cell proliferation. 5 genes (ESR2, FOXA1, JUN, PTGS2, VEGFA) were up-regulated and 10 genes (ADAM23, ATM, BCL2, CDH1, EGF, GLI1, ID1, MKI67, SNAI2 and THBS1) were down-regulated by these extracts. In addition, these extracts increased expression of miR-210 and blackberry ellagitannins-enriched extract reduced the expression of miR-19a/b.  

**Conclusion:** Collectively these results suggest that blackberry and grumixama anthocyanins and ellagitannins represent promising agents for inhibition of breast cancer development. Their protective actions involved induction of cell death (apoptosis), inhibition of cell proliferation and modulation of the expression of genes and miRNAs related to apoptosis, cell proliferation and differentiation, aggressiveness, invasion and metastasis.  

**Financing:** FAPESP/Food Research Center (Proc.2013/07914-8); CNPq (Proc. 142275/2013-1)
P13
Using Targeted Next-Generation Sequencing as a Cost-Efficient Method to Characterize the Genetics of Salt Sensitivity
Olivia Dong1,2, Oscar Suzuki1,2, Mildred Pointer3, Martin Kohlmeier4, Tim Wiltshire1,2
1 Division of Pharmacotherapy and Experimental Therapeutics, UNC Eshelman School of Pharmacy, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA; 2 Center for Pharmacogenomics and Individualized Therapy, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA; 3 North Carolina Central University, NC, USA; 4 UNC Nutrition Research Institute, Kannapolis, NC, USA

Background: Genetic variation is associated with inter-individual variability in blood pressure in response to dietary sodium intake (salt-sensitivity, SS). Extensive time is required to assess this phenotype in clinical practice; however, identifying key predictive genetic markers may be a faster and less expensive screening approach. A cost-efficient method allowing for extensive and unbiased interrogation of the genome in large sample sizes is needed to identify key haplotypes and single nucleotide polymorphisms (SNPs) to elucidate the genetics associated with SS.

Objectives: Develop a cost-efficient genetic assay that interrogates extensive genomic regions to identify the genetic markers associated with SS.

Methods: A genetic assay was developed to sequence approximately 124 kb of the genome consisting of 37 genes (all coding exons and promoter regions for 18 genes; SNPs in 19 genes) associated with SS. The evidence linking these genes to SS varies in strength (i.e. highly correlated or inconclusively correlated). The assay utilizes a custom-capture method using molecular inversion probes to generate a library of DNA fragments for next-generation sequencing (NGS) on the Illumina MiSeq platform. The cost of processing one sample includes DNA extraction ($11), library preparation and NGS, and personnel time ($17).

Results: A MiSeq run multiplexing 62 DNA samples from the 1000 Genomes Project was completed using the genetic assay. Average coverage across all genes for the 62 samples was 200x (range: 8x-663x). Four genes had an average of less than 20x average coverage. The cost of processing one sample was $63 or $1.70 per gene. This cost includes DNA extraction ($11), library preparation and NGS costs ($35), and personnel time ($17).

Conclusions: The preliminary genetic sequencing results indicate further optimization of the assay is needed to ensure all genes have a minimum sequencing depth of 20x for proper genotyping and haplotype calling. The results also suggest it is a promising method for interrogating extensive genomic regions for haplotype and SNPs associated with SS, and the economical cost makes sequencing of large cohorts possible.

P15
Associations between Nutrient Intake, Visceral Fat and ECHDC3 mRNA Expression in Patients with Different Extents of Coronary Lesion
Mychelle Kytchia Rodrigues Nunes Duarte1, Jéssica Nayara Góes de Araújo2, Victor Hugo Rezende Duarte2, Jéssica Bastos Pimentel2, Carolinne Thaisa de Oliveira Fernandes2, Angélica Luiza de Sales Souza3, Thayane Oliveira Souza4, Juliana Marinho de Oliveira4, Ananília Medeiros Gomes da Silva2, Diego Marques da Costa Santos2, Marina Sampaio de Menezes Cruz5, Renata Caroline Costa de Freitas4, Aline Tuane Oliveira da Cunha1, Raquel Suzana Matos de Miranda Torrinhas3, Severina Carla Vieira Cunha Lima3, André Ducati Luchessi2, Vivian Nogueira Silbiger2
1 Postgraduate Program of Nutrition, Federal University of Rio Grande do Norte, Natal, Brazil; 2 Department of Clinical and Toxicological Analysis, Federal University of Rio Grande do Norte, Natal, Brazil; 3 Faculty of Nutrition, Federal University of Rio Grande do Norte, Brazil; 4 Hospital Universitário Onofre Lopes, Federal University of Rio Grande do Norte, Natal, Brazil; 5 Postgraduate Program of Renorbio, Federal University of Rio Grande do Norte, Natal, Brazil; 6 Postgraduate Program of Science of Health, Federal University of Rio Grande do Norte, Natal, Brazil; 7 Department of Gastroenterology (LIM-35), Faculty of Medicine, University of São Paulo, São Paulo, Brazil

This study aimed to assess the relationships among nutrient intake, visceral fat and ECHDC3 mRNA expression in blood of patients with differing extents of coronary lesions. Blood samples were collected from male and female participants (n = 41) aged 30-74 years old, who underwent coronary catheterization for serum biochemistry and peripheral blood mRNA expression of ECHDC3 assessments. A 24-hour recall for dietary intake calculation, and anthropometric and bioimpedance analysis for visceral fat mass estimation was also completed for each participant. The extent of coronary lesion was classified according to the Friesinger index (FI). Correlations between the variables studied were tested by Spearman analysis. According to FI, patients were classified: without (n = 8), low (n = 6), intermediate (n = 15), or major (n = 12) lesion. We found a positive correlation of lesion extension with carbohydrate (r = 0.574, p = 0.002), thiamin (r = 0.507, p = 0.001), riboflavin (r = 0.388, p = 0.012), folate (r = 0.599, p < 0.001) and magnesium (r = 0.528, p = 0.003) intakes, and with ECHDC3 expression (r = 0.327, p = 0.037). ECHDC3 expression was 1.93-fold higher in patients with intermediate than those with low lesions (p = 0.011), 1.91-fold higher in patients with major than those with low lesions (p = 0.013), and showed a positive correlation with FI (r = 0.327, p = 0.037) and visceral fat percentage (r = 0.416, p = 0.009). Our data suggested that the consumption of high amounts of carbohydrates and micronutrients (thiamin, riboflavin, folate and magnesium) may be associated with the development of cardiovascular disease. Moreover, the increase of visceral fat and ECHDC3 upregulation may contribute to disease progression.

P14 – WITHDRAWN
A Vegan Diet May Lead to Substantial Changes in Gene Regulation by Modifying DNA Methylation Patterns
Valery Filippov¹, Karen Jaceldo-Siegl², Alexey Eroshkin², Vasiliy Loskutov¹, Charles Wang¹, Maria Filippova¹, Gary E. Fraser³, Penelope Duerksen-Hughes¹
¹Department of Basic Sciences, Loma Linda University School of Medicine, Loma Linda, CA 92354; ²Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA 92037; ³School of Public Health, Loma Linda University, Loma Linda, CA 92354, USA

Vegan diets, which avoid all animal based foods, have attracted increasing levels of interest and acceptance in recent years. Vegan diets are believed to confer numerous health benefits that include a decreased rate of aging and mortality, thereby increasing longevity. However, the molecular basis underlying these observed effects are not well-understood. To address this gap, we asked whether there were epigenomic differences between vegans and omnivores by interrogating methylation levels across genomic DNA derived from blood cells in vegans, then compared those findings to similar data obtained in omnivores. Primary analysis identified a significant number of differentially methylated CpG sites, indicating that the presence or absence of animal derived food can indeed cause epigenetic changes that may affect the regulation of gene expression. Moreover, detailed analysis of genes that are differentially methylated also revealed that many are clustered within a few highly enriched groups of transcriptional factors, each of which has the potential to participate in the regulation of multiple other genes. This finding implies that epigenetic regulation is only one layer of gene expression control, and that the number of nutrition-sensitive genes may be much higher than the number of genes with altered methylomes. Another cluster includes genes that regulate several signal transduction cascades. Interestingly, we found that genes involved in the trafficking of two major opposing neurotransmitters, GABA and glutamate, were differentially methylated. Since the DNA analyzed in this study was derived from blood cells, these findings raise the possibility of novel and unknown functions for these two important biologically active molecules outside of the central nervous system. These studies demonstrate for the first time that differences in food consumption in humans are reflected in extensive epigenetic alterations in the DNA methylation of many genes that play important and sometimes key roles in a variety of biological processes that regulate crucial cell life functions.

Perilipin (PLIN) Gene Polymorphism (rs894160) and Anthropometric Characteristics in a Population Group in the Northeast Region of Brazil
Naia Ivo Ferreira¹, Cecília Rios Mateus Lopes³, Vanessa Rosa Oliveira¹, Leticia Borges Batinga¹
¹Clinic DNA Nutrition, Brazil

Background: Polymorphisms in the perilipin (PLIN) gene have been investigated, in different populations, due to their potential relation with obesity, insulin sensitivity and markers of metabolic syndrome. Moreover, these genes are part of a family of proteins that can be found on lipid droplet surfaces in adipocytes and steroidogenic cells, and plays a central role on lipid storage and breakdown. Objectives: The aim of the study was to investigate the effect of the PLIN (rs894160) polymorphism on anthropometric variables, such as body mass index (BMI), body fat mass and waist circumference in a population group in the Northeast region of Brazil. Methods: Cross-sectional study, with 43 individuals of both genders (aged 27-64 y), inhabitants of Northeast region of Brazil, selected by convenience. Body Mass Index (BMI), body fat mass, waist circumference and the PLIN polymorphism (rs894160) were determined. Factorial analysis of variance (ANOVA) was used to evaluate the differences between the genotypes. Results: Among the participants of the sample, the allele A in rs894160 was carried by 41.9%. This polymorphism was significantly associated with an increase of waist circumference (by WHO criteria of waist circumference higher than > 94 cm for men and > 80 cm for women) (p<0.05). The presence of the A allele led to 10 cm increase in waist circumference. In contrast, the other anthropometric measures, such as BMI and body fat mass, did not present significant difference (p>0.05). Conclusion: In this sample, the PLIN polymorphism (rs894160) showed significant association with an increase in waist circumference, what seems to be a genetic influencer over growth of waist circumference. The anthropometric variables BMI and fat mass were not related to the genetic variation, so these interactions require further attention.
DNA methylation is one of the major mechanisms through which epigenetic control of gene expression is achieved in developmental and homeostasis processes. To better understand the role of nutrition in regulating gene expression, we performed DNA methylation genome-wide profiling of blood cell DNA samples isolated from 106 individuals, 38 vegans and 68 omnivores, using the Illumina Infinium HumanMethylation450 platform. Comparative analysis using Partek Genomic software revealed that of the differentially methylated CpG sites identified, the majority are located within known genes. Using two separate analysis engines, IPA and DAVID, we identified several significantly enriched gene clusters and pathways that control vital cellular processes. This includes homeobox and forkhead box transcriptional factors that participate in variety of cell functions, as well as several signaling pathways such as Ras and Wnt. These findings indicate that there is a specific and significant response of the methylation landscape in humans. Analysis of these alterations indicate the potential of diet and dietary changes to specifically impact the regulation of genes that participate in important and vital cellular processes.

**Objective:**
This study aimed to investigate the interaction between FTO single nucleotide polymorphisms (SNP), rs8050136, rs9930506, and plasma polyunsaturated omega 3 fatty acid (n-3) profile on an inflammatory score in adults of a cross-sectional population-based study conducted in São Paulo, Brazil.

**Methods:** All adults (>19 y and <60 y) participants of the Health Survey-São Paulo who matched the inclusion criteria were selected to participate in the study (n=225). Smoking habits and physical activity information were obtained using structured questionnaires. Plasma fatty acid profile, eleven plasma inflammatory biomarkers concentration and genotyping of FTO SNP rs8050136, rs9930506, rs9939609 were analyzed from blood samples. Anthropometric measurements were performed in standardized methods. Hardy-Weinberg equilibrium was accessed using a Chi-squared test with continuity correction and the association between inflammatory score and genotypes was conducted using multiple linear regression model adjusted for age, body mass index (BMI), smoking status, gender and physical activity level and the same model was used to test for genotype-plasma fatty acid profile interaction.

**Results:** All SNP analyzed were in Hardy-Weinberg equilibrium. Inflammation score was positively correlated with waist circumference (p<0.001), BMI (p<0.001), age (p<0.001) and plasma triacylglycerol concentration (p=0.041), and smoking subjects had statistically higher mean inflammation score than non-smokers (p=0.045). Statistically significant interaction was found between FTO SNP and n-3, so that homozygous for the major alleles for all the FTO SNP presented lower inflammation score when plasma n-3 level was higher (β95%CI) for rs8050136CC, rs9930506AA and rs9939609TT are, respectively: -1.45(-2.28; -0.63) and p inter=0.005; -1.06(-1.76; -0.36) and p inter=0.032; -1.51(-2.38; -0,64) and p inter=0.006).

**Conclusion:** FTO SNP genotypes are effect modifiers of the relationship between plasma n-3 levels and inflammation score in adults from Sao Paulo city (Brazil). The replication of these results in other populations can confirm the potential of genomie-personalized nutrition for the prevention of low-grade chronic inflammation-related diseases.

**Objective:** Investigate the frequency of apolipoprotein E (APOE) single nucleotide polymorphisms (SNP) in a Brazilian population sample, evaluated by a predictive commercial test, named Nutrigenetica®.
Methods: 295 people mean aged 27 years old participated (196 women and 99 men). We selected patients who did the predictive commercial test since March, 2015 from May, 2017. Briefly, 1 mL of saliva samples was collected in an Oragene DNA (OG-510) saliva collection device and the allelic discrimination experiment was performed in the Taqman OpenArray® Genotyping system in Centro de Genomas® Laboratory, located in Sao Paulo, Brazil. We genotyped APOE SNP (rs429358 and rs7412). We calculated de minor allele frequency (MAF) for each SNP and then compared with scientific literature. The reference used was the data from 1000 genomes Project Phase 3, based on CEU population (Utah residents with Northern and Western European Ancestry from the CEPH collection).

Results: All the genetics variants were in Hardy-Weinberg equilibrium. In relation to 1000 genomes Project data, we observed similar nutrition. We encourage further intervention studies with the application of personalized nutrition.

Conclusion: The presence of ε4 allele is associated with the increasing risk for late onset Alzheimer’s disease (LOAD) and for cardiovascular disease. Alterations in lipid profile can influence both metabolic risks and some studies show the relationship between genotypes and western diet. Genetic variants described in this study may contribute to future research in precision medicine field and may also encourage further intervention studies with the application of personalized nutrition.

Abstracts

P22

Effect of the Diet on the Transcriptional Expression in Ewes Mammary Gland with Single or Twin Gestation

M.A. Gallardo1, C. Geoffroy1, L. Arias1, S. Hernandez1, J.G. Cárcamo1

1Instituto de Bioquímica y Microbiología, Facultad de Ciencias, Universidad Austral de Chile, PO Box 567, Valdivia, Chile

This experiment was carried out to evaluate the effect of the diet on the transcriptional expression levels of some genes related to mamogenesis / angiogenesis processes in the mammary gland in ewes gestating either a single lamb or twins. Eighteen Ile de France ewes, third birth and similar BCS (3.0), 9 gestating a single lamb and 9 gestating twins, were separated in two groups according to the diet consumed: naturalized pasture (NP) or red clover (RC) from day -45 pre-partum until day +70 post-partum. Samples from mammary gland were obtained by biopsy procedure at days -10 pre partum (time 1), +30 (time 2) and +70 post partum (time 3). Samples were analyzed by qRT-PCR and data processed using REST statistical software.

Results showed that while ANGPT2 (P<0.03) and TBAX (P<0.01) were underexpressed at time 2 in single gestating ewes fed RC, the ewes gestating twins fed RC overexpressed VEGF (P<0.01), VEGFR1 (P<0.02), VEGFR2 (P<0.00), ANGPT1 (P<0.00), ANGPT2 (P<0.00), MK167 (P<0.00), TBAX1 (P<0.01), BCL2 (P<0.00), IGFBP3 (P<0.02) and underexpressed TGFBR1 (P<0.04) and TGFBR1R1 (P<0.02) at time 1. Although at time 2 the ewes only underexpressed CAIV (P<0.04) and TGFBR1 (P<0.03), overexpressing TGFBR1R1 (P<0.00), at time 3 underexpressed VEGFR1 (P<0.04), VEGFR2 (P<0.04), ANGPT1 (P<0.04), ANGPT2 (P<0.03), MK167 (P<0.04), TBAX1 (P<0.02), CAIV (P<0.05), BAX (P<0.01), BCL2 (P<0.04), CCND1 (P<0.04), IGFB1 (P<0.05), LPT (P<0.01), LPTR (P<0.04) and IGFB1 (P<0.01).

It is noticed a transcriptional up-regulation of some genes related to angiogenesis, such as VEGFR1, VEGFR2, ANGPT1, ANGPT2, MK167 and TBAX1 and other genes related to cellular proliferation/turnover, such as BCL2, IGFB1 and IGFB1 at time 1, followed by a downregulation of the same genes at time 3, suggesting the diet type.
should be a factor to consider in ewes gestating twins but not in single pregnancy.

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**P23**

**Identification of miR-612 and miR-1976 as Prospective Biomarkers of Response to Specific Weight-loss Diets**

M. Garcia-Lacarte†, F.I. Milagro1,2, M.L. Mansego1,2, M.A. Zulet1,2,3, J.A. Martinez1,2,3,4

1Centre for Nutrition Research/Department of Nutrition, Food Sciences and Physiology, University of Navarra, Pamplona, Spain; 2CIBERObn, Centro de Investigación Biomédica en Red de la Fisiopatología de la Obesidad y Nutrición, Carlos III Health Institute, Madrid, Spain; 3IdiSNIA, Navarra’s Health Research Institute, Pamplona, Spain; 4Madrid Institute of Advanced Studies (IMDEA Food), Madrid, Spain

**Background:** Non-coding RNAs (i.e., miRNAs) play a role in the development of obesity and related comorbidities by regulating gene expression. The aim of this study was to identify candidate miRNA biomarkers throughout –omic approaches in order to predict the success of specific weight-loss dietary treatments.

**Materials and Methods:** Genomic DNA isolated from white blood cells of a subpopulation from the RESMENA nutritional intervention study was hybridized in Infinium Human Methylation450 BeadChip arrays (31 Low-responders (LR) vs 16 High-responders (HR)), and in Illumina Human HT-12 v4 gene expression BeadChips (31 Low-responders (LR) vs 16 High-responders (HR)). A bioinformatic prediction of putative target sites of selected miRNAs was performed by applying miRBase algorithms. A total of 134 miRNAs were differentially methylated between HR and LR in the methylation array (87 hypomethylated and 47 hypermethylated), whereas 44 miRNAs were differentially expressed between both groups in the expression array (10 downregulated and 34 upregulated). Specifically, miR-1237, miR-1976, miR-642, miR-636, miR-612 and miR-193B were simultaneously hypermethylated and downregulated in HR. miR-612 and miR-1976 showed the greatest changes in methylation and expression, respectively; thus, these miRNAs were selected for further evaluation. The bioinformatic prediction revealed that TP53 was a putative target gene of miR-612 and CD40 of miR-1976. Moreover, TP53 was downregulated in the expression array when comparing HR vs LR expression levels adjusted by sex, diet, age and baseline weight (p = 0.024) and CD40 showed a trend (p = 0.069). Gene expression levels of TP53 and CD40 were also negatively correlated with the expression of miR612 and miR1976, respectively (p = 0.290; R = -0.249 TP53-miR-612; and p = 0.023; R = -0.505 CD40 miR-1976).

**Conclusion:** miR-612 and miR-1976 could be prospective biomarkers of response to specific weight-loss diets and might regulate the expression of TP53 and CD40.

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**P24**

**Determination of Body Fat: A Pilot Study Measuring Body Composition Among College Athletes at a Historically Black University**

Pratibha Gupta1, Aditya Phadke2

1Central State University, Wilberforce OH; 2Wright State University, Dayton OH, USA

**Background:** Type 2 diabetes and obesity are major public health priorities because of their high prevalence and incidence. African Americans have the highest prevalence to obesity (29%) and diabetes (11%) among racial groups. Obesity and diabetes are strongly associated with other diseases with high mortality rates such as Hypertension, Coronary artery disease, Cancer. The Non-drug treatments included were physical activity and diet lifestyle modification.

**Objective:** The purpose of this investigation was threefold: 1. Present a profile of body composition assessment using air displacement plethysmography on a population of college athletes at Central State University, 2. Evaluate Body Composition, 3. Determine body fat in College Athletes.

**Specific Aims:** To determine Body composition, to evaluate and assess specific composition, to Identify risk associated with based on body composition.

**Research Methods:** We recruited student athletes involved in any sports activities such as volleyball, basketball or any other athletic sport to participate in the study. 51 college athletes male and female, (average age 21) participated in the study. Body Composition measures included %Fat and %Fat Free Mass.

**Conclusions:** A major finding of this study was that a significant number of college athletes had normal and ideal FFMI and FMI. This body composition result could be due to an optimal healthy life style and a particular exercise regimen. Regular assessment of body composition is of importance as changes in body composition have been linked to several health risks. Health studies have shown that a combination of physical activity and a well maintained diet produce the best effect of a decrease in fat mass and a maintenance or increase in muscle mass. Scientists have identified a small number of single nucleotide polymorphisms (SNPs) that respond to diet and exercise.
Zinc Transporter ZIP13 and Endoplasmic Reticulum (ER) Stress: A Gene-Nutrient Interaction Which May Affect Nervous System Function

Matthew D. Hart1, Morgan D. Strong1, Tony Z. Tang1, Kaitlyn N. Jerome1, Kylin G. Wang1, Brenda J. Smith1, Winyoo Chownanadisa1

1Department of Nutritional Sciences, Oklahoma State University, Stillwater, OK, USA

Implementation of precision nutrition based upon genomic information requires understanding of specific gene-nutrient interactions for distinct health relationships. ZIP13 is an intracellular zinc transport protein which transports zinc into the cytoplasm from internal compartments. Loss of function mutations in ZIP13 cause the spondylochondrodysplastic form of Ehlers Danlos syndrome, a connective tissue disorder characterized by improper collagen function. However, ZIP13 is widely expressed in nearly all body tissues, including the brain. Accordingly, ZIP13 may play a significant physiological role across multiple body tissues, and mutations or polymorphisms in ZIP13 may be a risk factor for many diseases, including disorders of the nervous system. It has previously been suggested that ZIP13 loss of function induces significant ER stress and the Unfolded Protein Response (UPR). The work presented here supports the hypothesis that ZIP13 is necessary in Neuro-2a neuroblastoma cells for preventing the induction of the ER stress response and promotion of neurite outgrowth, a critical element of neuronal differentiation. Small hairpin RNA mediated ZIP13 knockdown (KD) results in decreased ZIP13 mRNA expression relative to control transfected Neuro-2a cells. ZIP13 KD inhibits neurite extension in Neuro-2a cells differentiated with retinoic acid. ZIP13 KD cells display significant reductions in neurite length, number of neurites per cell, and number of branches per neurite. The observed reduction in neurite outgrowth of ZIP13 KD cells is restored by treatment with sodium tauroursodeoxycholate (TUDCA) and 4-Phenybutyric acid (4-PBA), compounds which can relieve ER stress. It is possible that genetic variations in ZIP13 leading to reduced function may lead to increased susceptibility to ER stressors from nutritional and environmental sources in the nervous system or other physiological systems.

Interaction Between the Betaine-Homocysteine Methyltransferase (BHMT) Gene Polymorphism and Carbohydrate Intake on Homocysteine Levels in Brazilian Adolescents

Shelini Surendran1, Carla C. Morais2, Julie A. Lovegrove1, Cristiane Cominetti2, Karani S. Virnal1, Maria A. Horst2

1Hugh Sinclair Unit of Human Nutrition, Department of Food and Nutritional Sciences, University of Reading, Reading RG6 6AP, UK; 2Nutritional Genomics Research Group, Faculty of Nutrition, Federal University of Goiás (UFG), Goiania, Goiás, Brazil

Background: Genetic variants can interact with environmental stimuli such as diet and physical activity, creating variable responses to metabolic traits in several populations. At present, several single nucleotide polymorphisms (SNPs) have been associated with B vitamins, folic acid and homocysteine status.

Objective: The aim of this study was to investigate the association between MTHFR, MTR, MTRR, CBS, TCN2, COMT, BHMT, FUT2 and NBPF3 gene polymorphisms with vitamins B12, homocysteine and folic acid blood levels and investigate whether dietary factors and physical activity modified the possible associations.

Design: A total of 115 adolescents (10-19 years old), from a public school in Brazil underwent anthropometric, biochemical and genotyping analyses. Dietary intake was estimated with food records and physical activity was based upon self-report. Interaction analyses were performed using linear/logistic regression model. Potential confounders such as age, body mass index, sex were adjusted in the regression model.

Results: None of the associations between the analysed SNPs and vitamin B12, homocysteine and folic acid concentrations were statistically significant after correction for multiple testing. In the interaction analysis, after Bonferroni correction, there was a significant interaction between BHMT SNP (rs689415) and carbohydrate intake (% energy) (Pinteraction=5x10^-8) on homocysteine levels, where the 'G' allele carriers had 38.04% higher circulating homocysteine concentrations than those with ‘AA’ genotype (P=0.04) among individuals in the highest tertile of carbohydrate intake (% energy, 71.56 ± 12.79).

Conclusion: Our findings suggest that the association between the BHMT SNP (rs689415) and circulating homocysteine might be influenced by carbohydrate intake in adolescents. Further understanding of how the BHMT influences circulating homocysteine through dietary interventions is imperative to advance the development of personalised nutrition, to reduce the risk of elevated homocysteine concentrations. The limitation of this study includes a self-report physical activity and no evaluation of betaine and BHMT protein levels.
Analysis of Intestinal Microbiota in Patients with Functional Constipation After Synbiotics Treatment
Lin sheng Huang, Tong yi Shen, Xuebing Yan, Huan fong Qin

Department of General Surgery, Shanghai Tenth People’s Hospital affiliated to Tongji University No.301, Yanchang Road, Shanghai 200072, China

Objectives: The aim of this study was to investigate whether clinical symptoms and intestinal microbiota is improved after synbiotics treatment.

Methods: Fecal samples from 53 patients diagnosed with chronic functional constipation according to the Rome III criteria were analyzed by 16S-rRNA sequencing. After Synbiotics treatment for one and three month, fecal samples collected from 36 patients and 15 patients respectively. The clinical outcome and intestinal microbiota of patients were then analyzed and 53 healthy community volunteers were employed as controls. To determine the most discriminative species, cross validated logistic ridge regression was performed.

Results: The microbiota composition of constipation group is different from that of treatment group and healthy group. With synbiotics treatment 1 month and 3 month, abundance of Escherichia-Shigella is decreased, that of Prevotella 9 and Lactococcus are increased. Compared with constipation and healthy group, genus of Parabacteroides,Erysipelotrichaceae_UCG-003 were abundant in the constipation group, while Prevotella 9,Megamonas, Klebsiella, Enterobacteriaceae unclassified are abundant in the healthy group. Compared microbiota in three groups (constipation group, healthy group as well as treatment group) Prevotella 9 is the characteristic bacteria that decreased in constipation group and increased in treatment group. (2.12% vs. 14.46% vs. 5.65%, P = 1.13E-07, q = 3.21E-06, FDR=7.56E-06). after synbiotics treatment, stool frequency is increased and the proportion of constipated complications (abdominal distension, loss of appetite, anxiety rate) is decreased, the proportion of laxative dependent is also decreased.

Conclusions: Synbiotics treatment can promote clinical remission and microbiota improvement in constipation group. Identification of the key bacterial genus is an important approach to reveal therapeutical mechanism of synbiotics treatment, and therefore promotes the development of microbiota-based individualized treatment in patients with functional constipation.

The Role of Vitamin E on Tight Junction Proteins of Intestinal Barrier
Yanjun Huang1,2, Sichun Zhou1,2, Caimei He1,2, Ting Tao1,2, Huansheng Yang1,2, Yulong Yin3,4, Xiaoping Yang1,2,3

1Department of Pharmacy, School of Medicine, Hunan Normal University, Changsha, Hunan, PR. China 410013; 2Key Laboratory of Small Targeted Molecules’ Discovery, Hunan Normal University, Changsha, Hunan, PR. China 410013; 3Animal Nutrition and Human Health Laboratory, School of Life Sciences, Hunan Normal University, Changsha, Hunan 410125, China; 4Chinese Academy of Sciences, Institute of Subtropical Agriculture, Key Laboratory of Agroecological Processes in Subtropical Region, Changsha 410125, China

Although application of vitamin E has been long time used in both human and animal nutrition worldwide, the detailed mechanisms are largely unknown. Tight junction (TJ) proteins including occludin, claudin-1 and ZO-1 play an important role in intestinal nutrition. We aim to examine the regulatory role of vitamin E on TJ proteins. Caco-2 and IPEC-J2 cell lines are treated with vitamin E alone or LPS and the alterations of these TJ protein expressions were measured by either western blot or RT-PCR. The results demonstrate that vitamin E dramatically increases the expression of TJ proteins and it relieves activation of IL-6 and NF-κB after LPS stimulation. The data collected from intestine tissues from vitamin E-treated weaning suckling pigs consistently reveals the similar findings. Altogether, the present study shows a novel role of vitamin E on protecting the intestinal barrier and improving the inflammation via affecting tight junction proteins in intestine. These results build a solid foundation for appropriate utilization of vitamin E with suitable dose on both human and animal nutrition.

Genetic Variation in Mitochondrial Complex I Activity Among Inbred Strains of Mice
Simon T. Hui1, Telisha M. Swain2, Calvin Pan1, Maria De Luca3, Shannon M. Bailey2, Aldons J. Lusis1

1Department of Medicine, Division of Cardiology, David Geffen School of Medicine, University of California at Los Angeles, Los Angeles, CA 90095, USA; 2Department of Pathology, University of Alabama at Birmingham, Birmingham, AL 35233, USA; 3Department of Nutrition Sciences, University of Alabama at Birmingham, Birmingham, AL 35233, USA

Mitochondrial dysfunction has been shown to link to metabolic diseases, such as obesity, NAFLD and diabetes. To dissect the genetic control of mitochondrial functions and its physiological role, we employed a panel of >100 strains of inbred mice, termed Hybrid Mouse Diversity Panel (HMDP), which allows for high resolution mapping. Previously, we have shown that when fed a high fat/high
sucrose obesogenic diet, HMDP mice exhibited large variations in obesity, insulin resistance and hepatic steatosis which are dependent on their genetic background. We isolated mitochondria from 287 livers from 102 strains of male mice. Complex I activity was measured and normalized to citrate synthase activity. A 10-fold difference was observed between the extreme strains. We found that complex I activity was negatively correlated with obesity traits: body weight, body fat percentage, fat mass response to diet. Parallel to the obesity connection, complex I activity was inversely correlated with food intake and HOMA-IR. These data show that complex I activity is tightly associated with body composition and insulin action. Metabolomic analysis revealed a negative correlation between complex I activity and plasma gamma-aminobutyric acid (GABA). In the liver, complex I activity showed a negative correlation with 4-aminobutyrate aminotransferase (Abat), an enzyme that catalyzes GABA catabolism and is involved in the mitochondrial nucleoside salvage pathway. Our finding is consistent with previous report that human subjects harboring homozygous ABAT missense mutation displayed increased GABA level and reduced mtDNA in the brain. These data suggest that GABA may be a useful plasma biomarker for mitochondrial function. Genome wide association mapping identified two significant loci on chromosomes 8 and 13. The chromosome 8 peak spans ~1.4Mb and contains 18 genes whereas the chromosome 13 peaks is ~1.7 Mb wide containing 4 genes. The identification and validation of causal genes in these loci will provide novel insight into the regulation of complex I function by nuclear genes.

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**P30**

**Effects of Probiotic Supplementation on Gut Microbiota and Obesity Outcomes in Obese Hispanic Adolescents: A 16-Week, Randomized, Placebo-Controlled Trial**

**Roshonda B. Jones**, Tanya L. Alderete, Ashley A. Martin, Bree A. Geary, Darryl H. Hwang, Suzanne L. Palmer, Michael I. Goran

1Department of Preventive Medicine, University of Southern California, Los Angeles, CA, USA; 2Department of Radiology, University of Southern California, Los Angeles, CA, USA

Numerous studies have shown that there are links between obesity, liver fat, and the gut microbiome. However, there are mixed results on whether probiotics could impact the gut microbiome and/or help to decrease liver fat and obesity outcomes. The primary aim of this study was to determine whether a probiotic supplement (VSL#3®) intervention altered gut microbiota and/or gut hormones associated with appetite regulation. The secondary aim of this study was to determine whether VSL#3® altered body composition and liver fat and fibrosis. We conducted a double-blind, randomized placebo-controlled trial in 19 obese Latino adolescents. Adolescents received three packets per day of either VSL#3® or a matched placebo for 16 weeks. Pre- and post-intervention measures included gut bacterial relative abundance, gut appetite-regulating hormones, anthropometrics, body composition, liver fat, liver fibrosis and dietary intake. We conducted linear models to determine whether there were any significant differences in the changes in these outcomes following VSL#3® intervention. Models adjusted for sex, change in caloric intake, and body mass index (where appropriate). Compared to placebo, adolescents that received VSL#3® had significant increases in total adiposity (1.7 ± 0.6 vs. -1.3 ± 0.5, p < 0.01) and trunk adiposity (3.3 ± 0.8 vs. -1.8 ± 0.8, p < 0.01) with no significant effects on liver fat/fibrosis, insulin/glucose, gut microbial abundances or gut hormones. We conclude that VSL#3 supplementation may lead to increased adiposity in obese Latino adolescents with no significant detectable changes in gut microbiota, gut appetite-regulating hormones, liver fat and fibrosis and dietary intake.

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**P31**

**Association between Serum Zinc Level and Body Composition: The Korean National Health and Nutrition Examination Survey**

**Ha-Na Kim**, Sang-Wook Song, Whan-Seok Cho

1Department of Family Medicine, College of Medicine, St. Vincent's Hospital, The Catholic University of Korea, Seoul, Republic of Korea; 2Department of Family Medicine, College of Medicine, Seoul St. Mary’s Hospital, The Catholic University of Korea, Seoul, Republic of Korea

**Objective:** We examined the associations between serum zinc levels and several body composition factors among Korean adults.

**Methods:** This study used data from the Korean National Health and Nutrition Examination Survey, a cross-sectional survey of Korean civilians. Data from a total of 1,896 subjects were analyzed.

**Results:** Serum zinc levels in men with elevated waist circumference were higher than in those with normal waist circumference (152.1 ± 3.7 μg/dL vs. 137.8 ± 2.2 μg/dL, P < 0.001) and serum zinc levels increased with increasing tertiles of total body fat percentage (134.2 ± 2.9 μg/dL, 142.0 ± 2.9 μg/dL, and 148.0 ± 2.7 μg/dL, P = 0.001). Among men with a normal waist circumference, the serum zinc levels of participants with the highest total body fat percentage were higher than in those with the lowest or medium total body fat percentage values (144.5 μg/dL vs. 135.2 μg/dL, P = 0.029). In contrast, in men with an elevated waist circumference, no difference in serum zinc levels according to total body fat percentage was detected. There was no relationship between serum zinc levels and body composition factors in women.

**Conclusions:** Body zinc status might be associated with the quantity and distribution of body fat in Korean men. Additional gender-specific studies are needed to determine whether the relationship of body zinc status with abdominal obesity and total body fat affects metabolic disorders and cardiovascular diseases.

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**P32 – WITHDRAWN**
Role of Sex- and Tissue-Specific Differences in Mouse Lipocalin-2 Expression in Regulating Insulin Resistance and Non-Alcoholic Fatty Liver Disease Progression

Karthickeyan Chella Krishnan1, Simon Sabir2, Calvin Pan1, Nam Chê3, Sarada Charugundla1, Hannah Qi1, Zhiqiang Zhou1, Simon Hui1, Frode Norheim1, Aldons J. Lusis1

1Department of Medicine/Division of Cardiology, David Geffen School of Medicine, University of California, Los Angeles, CA, USA; 2Department of Psychology, University of California, Los Angeles, CA, USA

Non-alcoholic fatty liver disease (NAFLD) is an umbrella term encompassing a range of liver abnormalities from simple steatosis (fat accumulation in hepatocytes) to complex non-alcoholic steatohepatitis (NASH), fibrosis and cirrhosis. The prevalence of NAFLD in the United States has increased dramatically over the past few decades. Population studies have strongly associated obesity, diabetes and insulin resistance (IR) to NAFLD. The regulation of secreted cytokines and proteins from adipose tissue, aka adipokines, and liver tissue, aka hepatokines is known to contribute to the pathogenesis of several cardiometabolic diseases including obesity, IR, NAFLD and atherosclerosis via their crosstalk between themselves, adipose and liver, and other tissues such as heart and skeletal muscle. Numerous adipokines and hepatokines have been identified so far; however, specific signalling pathways activated by most of these secreted proteins and/or mechanism(s) by which they communicate with each other (and other tissues) are currently unknown. It is therefore essential to determine the functional consequences of these proteins in regulating normal metabolic homeostasis in order to understand how they are disrupted in patients with obesity, IR and NAFLD. To determine the causal mechanism(s) mediating diet-induced obesity, IR and NAFLD, our lab subjected both sexes of over 100 inbred mice strains to high-fat, high-sucrose (HF/HS) diet for 8 weeks, collecting several transcriptomes. Extensive systems genetics analyses of these data suggest that female adipose and/or male liver derived lipocalin-2 (LCN2) may be mechanistically linked to deregulated lipid accumulation in liver, as well as enhanced inflammation in adipose and/or liver. LCN2 is a 25kDa secretory glycoprotein, first isolated from neutrophils; however, recently it has been shown to be a novel adipokine that is up regulated in adipose tissues of genetically obese animals and humans. LCN2 has also been shown to be a hepatokine associated with liver damage, inflammation and IR. Recently, LCN2 has also been implicated in recruiting neutrophils to the liver in both alcoholic and non-alcoholic steatohepatitis (ASH and NASH), a severe condition of the liver. The goal of our current study is to use systems genetics approaches to investigate the role and mechanism by which LCN2, a secreted protein from adipose and/or liver tissue, in response to diet-induced obesity, regulates IR and NAFLD in a sex-specific manner.
P35

**Xylo-Oligosaccharides Suppressed Visceral Fat Accumulation and Modulated Cecum Microbiome in Mice**

Jianfeng Long1, Jieping Yang1, Qing Li1, Susanne M. Henning1, Mark Hsu1, Brenda Chan1, Shih Lung Woo1, David Heber1, Zhaoping Li1, 2

1Center for Human Nutrition, David Geffen School of Medicine at UCLA, Los Angeles, CA 90095, USA

**Background:** Xylo-oligosaccharides (XOS) is a fiber that has been shown to improve the balance of gut microbiota and may improve insulin sensitivity and glycemic control, however, the metabolic regulatory effects of XOS remain unclear.

**Objective:** This study evaluated the metabolic regulatory effect of prebiotic fiber XOS supplementation in obese mice.

**Methods:** Male C57BL/6 mice were fed with AIN93M diet for 3 weeks, then were divided into three groups – control diet, 3%- or 10%-XOS (W/W) for additional 3 weeks (n=8/group). Body composition, liver and blood lipids, and gene expressions related to lipid homeostasis were measured. Cecum content was used for microbiota and short chain fatty acids (SCFAs) analysis.

**Result:** The body weight, liver weight, liver and blood lipids (total cholesterol and triglycerides), as well as beige adipogenic genes of E-fat did not differ among the three groups. 10% XOS supplementation significantly reduced the visceral fat, serum leptin, as well as beige adipogenic genes expression. The conditioned medium increased the binding of HIF-1a to Firmicutes when compared to non-treated cells as a negative control.

**Conclusion:** In summary, 10% XOS supplementation reduced visceral fat, and increased ratio of Bacteroidetes to Firmicutes and cecum SCFAs levels. XOS may mediate microbial changes, resulting in increased production of SCFAs, of which might modulate the metabolic environment of the host as shown by decreased mRNA expression of adipogenic genes in adipose tissue. The mechanism of how bacterial changes modulate adipogenesis in visceral adipose should be elucidated.

P36

**Identification and Validation of HIF-1α-Regulated Genes in 3T3-L1 Adipocytes Treated with Conditioned Medium**

A. López-Pascual1, S. Lorente-Cebrián1,2, J.A. Martínez1,2,3, P. González-Muniesa1,2,3

1Centre for Nutrition Research/Department of Nutrition and Food Sciences and Physiology, University of Navarra, Pamplona, Spain; 2Health Research Institute of Navarra (IdiSNAA), Pamplona, Spain; 3CIBERobn Physiopathology of Obesity and Nutrition, Centre of Biomedical Research Network, ISCIII, Madrid, Spain

**Background:** Obesity is a multifactorial chronic inflammatory disease which involves different pathways, being adipose tissue hypoxia among them. The aim of the present study was to characterize the effect of conditioned medium from LPS-activated macrophages on the regulation of HIF-1α-related genes in murine adipocytes.

**Methods:** The identification of HIF-1α-binding sites of 17 obesity-related genes was performed by applying bioinformatics predictions (PROMO web version 3.0.2). For the in vitro analyses, 3T3-L1 murine adipocytes (9 days post-differentiation) were incubated for 24h in conditioned medium (CM, composed by 75% complete medium and 25% medium of RAW 264.7 murine macrophages with 24h 500ng/ml LPS). In order to elucidate if these genes were regulated by HIF-1α, a validation with chromatin immunoprecipitation (ChIP-q-PCR) assay was carried out.

**Results:** ChIP assay showed that HIF-1α directly bound to HIF-responsive elements (HRE) in Vegfa and Ucp2 genes, regulating its expression. The conditioned medium increased the binding of HIF-1α at Vegfa (p<0.001) whereas it suppresses the enrichment at the Ucp2 gene (p<0.05). All these results were statistically significant compared to non-treated cells as a negative control.

**Conclusions:** Our results suggest that Vegfa and Ucp2 genes are direct targets of HIF-1α in 3T3-L1 adipocytes treated with conditioned medium from LPS-activated macrophages. The involvement of HIF-1α under pro-inflammatory conditions could help to clarify the role of this transcription factor in obesity and the associated comorbidities. Finally, this study propose a possible application of conditioned medium as a model of macrophage infiltration commonly found in the adipose tissue of obese subjects.
Altered Eating Disinhibition Genotype Associated with Higher Body Mass Index in Weight Loss Clinic Patients

Lori E. Arguello1, Kasuen Mauldin2, Deepika Goyal3
1The Henrried Center Weight Loss Clinic, Sacramento, CA; 2Department of Nutrition, Food Science, and Packaging, San José State University, San José, CA; 3The Valley Foundation School of Nursing, San José State University, San José, CA, USA

The purpose of this study was to (a) identify via chart review the type and frequency of six eating behavior genotypes, or the various forms of a gene, in a sample of patients in a medical weight loss clinic and (b) compare results by patients classified as overweight vs. obese (per body mass index [BMI]). This was a retrospective quantitative chart review from three medical weight loss clinics. A de-identified dataset was compiled, consisting of patient demographic data and the genotypes of six specific genes related to six eating behavior traits, or genetically determined characteristics, of eating disinhibition, food desire, hunger, satiety, snacking, and sweet tooth. Each genotype associated with these eating behavior traits was characterized as “typical” (non-weight-promoting behavior) or “altered” (weight-promoting behavior). A total of 768 charts were reviewed; 698 met inclusion criteria. Patients were separated into two groups: BMI < 30 kg/m² and BMI ≥ 30 kg/m². Of the six eating behavior traits, a statistically significant difference was noted between the two groups only for eating disinhibition (p < .001), defined as the likelihood to eat more than normal in response to a stimulus such as stress, social environment, tasty foods, etc. Based on these results, the altered genotype of the eating disinhibition gene (more likely to exhibit eating disinhibition) is more prevalent in patients with a BMI ≥ 30 kg/m² than those with BMI < 30 kg/m². Results suggested that alteration in the eating disinhibition genotype was associated with obesity. Earlier screening for this genetic alteration may help assess risk for obesity, which can lead to earlier intervention to promote optimal health outcomes and reduce obesity-related illnesses.

Preparing Animal Health Professionals with Knowledge Necessary to Understand Nutrigenetics and Nutrigenomics

Gustavo A. Mendoza-Fandino1, Mónica Duque Quintero1, Elsa María Vasquez-Trespalacios2
1Department of Animal Health, Corporación Universitaria Remington, Medellin, Colombia; 2School of Medicine, CES, Medellin, Colombia

Historically in Colombia, the role of veterinary practitioners has focused on handle and curing productive animals. Our advancing knowledge about animal genomes, now offer the opportunity to individualize dietary intervention to help prevent, mitigate or cure diseases. The development of genomics tools and collecting proof about the constituents of nutrients interacting with the genome as a mechanism to modulate the expression, open opportunities for the advancement of health and disease prevention based on the nutrition. The aim of this work is to describe strategies to introduce nutrigenomics and nutrigenetics topics for Colombian animal health professionals. The first step to achieve this goal was the adding of basic assignment topics related to understand nutrigenetic and nutrigenomics. These topics covered DNA, RNA, proteins, genomic process and regulation looking to a deeper understanding of molecular biology. The introduction of genomics and genetics concepts, in regular courses, facilitate the integration of the new concepts with the classic. Additionally, the animal science department create non-credits study groups, open to all students interested in the topic. The idea with these study groups is to promote abilities of critical thinking and reading of scientific literature related to nutrigenetic and nutrigenomics, looking to promote the animal care based in evidence. The study group has regularly organized talks, with professionals in topics related to animal nutrition and molecular biologist, to generate collaborative projects to involucrate students, as well as to expose them to both areas. Our participation in the 11th Congress of the International Society of Nutrigenetics and Nutrigenomics is our first international effort to be part of the community of scientist that are working to understand and solve the issues related to the nutrition and human health.
Methods: Eight weeks old mice were fed a High-Fat, High-Sucrose diet for 8 weeks to induce obesity, steatosis and insulin resistance. Liver ceramides from 98 strains of mice of both sexes were measured using mass spectroscopy. We also tested if gonadectomy had an effect on hepatic ceramide accumulation in both sexes of three unique strains of mice (C3H/HeJ, C57BL/6J and DBA/2J). The gonadectomized mice were compared to untreated and sham operated controls.

Results: Consistent with previous studies, ceramide levels were significantly associated with IR in male mice, and a similar significant association were observed in female mice. Surprisingly, however, female mice exhibited substantially higher hepatic C16:0 and C18:0 ceramide levels than male mice, despite being less insulin resistant. This also appears to be the case in humans based on levels of ceramides in plasma. We furthermore show that the sex difference in hepatic ceramides is mediated in part by testosterone repression of ceramide synthase 6.

Conclusions: Our data shows that most female mice strains have higher levels of liver ceramide C16:0 and C18:0 than their male counterpart. This sex difference can probably be explained by the inhibitory effect of testosterone in male mice. Future mechanistic mouse studies will show if liver ceramides oppose insulin action more potently in males than in females.

Abstracts

P40
Is Your Eating Behavior Shaped by Nature or Nurture? A Look into the Nutrigenomics of Taste

Raquel Novak1

1Leonard Davis School of Gerontology, University of Southern California, Los Angeles, CA, USA

The bitter taste receptor, known as the TAS2R38 gene, has the ability to influence one’s nutritional intake due to the perceived intensities of certain foods. A variety of children and adult studies have exposed links to increased sugar, sodium and fat intake or decreased vegetable intake based on participants’ perception of bitter taste. Individuals are born as either a taster or non-taster; this genetically inherited profile can affect preference of certain foods, potentially leading to unhealthy food choices and possible chronic diseases. Throughout numerous studies, tasters commonly disliked bitter vegetables and reported a heightened preference of sugar-sweetened items. Tasters experience increased, sometimes unpleasant, sensitivity to bitter compounds, whereas non-tasters experience decreased sensitivity or bland taste to the same bitter compounds. While non-tasters consumed a greater variety of dark vegetables, they preferred higher fat foods to provide a satisfying texture and mouth-feel to compensate for the lack of flavor non-tasters may experience. Appropriate nutrition interventions must take place to ensure a balanced, diverse, and nutritious diet for both tasters and non-tasters. The purpose of this article is to expand and increase awareness of genetic involvement with taste preferences and how to implement adequate nutrition interventions. Dietitians can work with both tasters’ and non-tasters’ genetically pre-disposed flavor-profiles and transform their taste preferences to improve overall quality of life.

P41
Identification of Plasma Metabolites Associated with NAFLD in Morbidly Obese Mexican Subjects

Elvira Ocampo-Medina1, Paola León-Mimila1, Joel Vega-Badillo1, Luis Macías-Kaufter1, Hugo Villamil-Ramírez1, Blanca López-Contreras1, Laura Reyes-Ferrín1, Francisco Campos-Pérez2, Isabel Ibarra-González2, Marcela Vela-Amieva2, Carlos A. Aguilar-Salinas3, Rogelio Hernandez-Pando3, Nahum Méndez-Sánchez4, Simon Hui5, Aldons J. Lusis6, Stanley Hazen6, Adriana Huertas-Vazquez7, Samuel Canizales-Quinteros7

1The National Institute of Genomic Medicine, Mexico City; 2Clinic for Bariatric Surgery, Hospital Dr. Rubén Leñero, Mexico City; 3The National Institute of Pediatrics, Mexico City; 4Department of Endocrinology, The National Institute of Medical Sciences and Nutrition, Mexico City; 5Department of Pathology, The National Institute of Medical Sciences and Nutrition, Mexico City; 6Médica Sur Clinic Foundation, Mexico City; 7Department of Medicine, Division of Cardiology, David Geffen School of Medicine, University of California, Los Angeles, USA; 8Cleveland Clinic, Cleveland OH, USA

Background: Non-alcoholic fatty liver disease (NAFLD) is a multifactorial disease characterized by an excess of liver fat infiltration and includes a spectrum ranging from steatosis, non-alcoholic steatohepatitis/NASH to cirrhosis and hepatocellular carcinoma. The incidence of steatosis and its progression to NASH varies among different populations, where Mexicans have the highest prevalence. The gold standard for NAFLD is the histopathological evaluation of liver biopsy, an invasive approach that is not broadly applicable to the general population. We applied targeted metabolomics and genome-wide association studies (GWAS) to discover novel biomarkers for NAFLD.

Methods and Results: A total of 46 plasma metabolites including amino acids, TMAO, choline betaine and acyl carnitines were measured using a targeted metabolomic approach. We investigate plasma levels in 425 morbidly obese Mexican subjects who underwent bariatric surgery with liver histology available (without steatosis (n=41), steatosis (n=88), NASH borderline (n=136) and NASH (n=160). We used ANCOVA to compare the differences between the groups adjusting for covariates such as age, sex and BMI. We observed that plasma levels of tyrosine (r=0.27, p=4×10^{-8}), alanine (r=0.26, p=4.7×10^{-5}), proline (r=0.22, p=1.7×10^{-4}), methionine (r=0.17, p=0.005), TMAO (r=0.14, p=0.03) and choline (r=0.13, p=0.03) were associated with NAFLD activity score (NAS) and its components including steatosis, inflammation and hepatocellular ballooning. In addition, we performed a GWAS to identify the genetic factors associated with plasma metabolite levels. We identified 3 SNPs within the proline dehydrogenase 1 (PRODH) gene to be significantly associated with proline levels (P<5×10^{-8}) and NAS score (r=0.83, p=0.0002).

Conclusions: Our findings suggested that plasma metabolites could distinguish between different grades of NAFLD, and could help to determine the mechanisms involved in NAFLD manifestation and progression.
Background and Aim: Our preceding analysis of the prevalence profile of polymorphisms in diet-related adaptive genes in West Mexico population suggested that they remained adapted to the features of the low-fat pre-Hispanic diet. Thus, a regionalized genome-based Mexican diet was designed. Herein, we aimed to evaluate its effect on biochemical and anthropometric variables in subjects with metabolic risk factors for chronic disease.

Methods: A quasi-experimental study was performed in 37 eligible mestizo subjects that underwent a 6-month dietary intervention. The diet mainly included available regional Mexican foods, known to have been the staples of the pre-Hispanic diet (e.g., beans, tortilla, tomato, avocado, chia, amaranth, etc.); and macronutrients distribution were 20% proteins, 50% carbohydrates, and 30% fat (of which saturated fat <7%, monounsaturated fat 10–15% and polyunsaturated fat 10%). Anthropometric and biochemical variables were measured at baseline and the changes after 3 and 6 months.

Results: A total of 33 subjects (age 42.6±11.7 y, 75.8% female) completed dietary intervention. Metabolic risk factors at baseline were: obesity (body mass index (BMI) of 30.1±5.6 Kg/m², body fat 10%). Anthropometric and biochemical variables were measured at baseline and the changes after 3 and 6 months.

Conclusions: A regionalized genome-based Mexican diet normalized hypertriglyceridemia, HOMA-IR and VLDL-c values, and significantly changed and normalized p < 0.05.

Conclusion: A regionalized genome-based Mexican diet normalized hypertriglyceridemia, HOMA-IR and VLDL-c values, and improved body composition. The recommendation and use of a more traditional diet based on the knowledge of Mexicans cultural history and their diet-related genetic adaptations may be efficient to prevent the development of the current non-communicable chronic diseases.
Dietary and nutritional factors have been shown to relate to a number of health-related outcomes, yet there is a need for studies that examine their relationship to molecular aging rates. Toward this end, we study the epigenetic clock and other predictors of chronological age based on site-specific DNA methylation; these biomarkers have previously been shown to be independent predictors of all-cause mortality, chronic conditions, and age-related functional decline. We analyze cross-sectional data from 4,173 postmenopausal female participants from the Women’s Health Initiative, as well as 402 male and female participants from the Italian cohort study, Invecchiare nel Chianti, examining associations between accelerated epigenetic aging and dietary, cardiometabolic, and lifestyle factors. Epigenetic age acceleration is broadly defined as the epigenetic age left unexplained by chronological age. Here we examine two distinct measures which are based on different sets of CpGs: intrinsic epigenetic age acceleration (IEAA) which is adjusted for variation in blood cell composition, and extrinsic epigenetic age acceleration (EEAA) which explicitly incorporates age-related blood cell counts. These measures can be interpreted as tracking cell-intrinsic aging and immune system aging, respectively. EEAA exhibits significant associations with reported fish intake (p=2x10^{-5}) and alcohol intake (p=3x10^{-5}), education (p=3x10^{-5}), BMI (p=2x10^{-5}), and blood carotenoid levels (p=2x10^{-5}) —an indicator of fruit and vegetable consumption, whereas IEAA is associated with reported poultry intake (p=4x10^{-5}) and BMI (p=4x10^{-5}). Both EEAA and IEAA were also found to relate to indicators of metabolic syndrome which appear to mediate their associations with BMI. Finally, longitudinal data suggests that an increase in BMI is associated with increase in both EEAA and IEAA. Overall, this analysis of epigenetic aging in blood suggests that the health benefits of fruit, vegetable, lean meat, and moderate alcohol intake, physical activity, and education, as well as the health risks of obesity and metabolic syndrome might be mediated in part by their anti- and pro-aging effects on the epigenome.

The prevalence of overweight and obese pets has increased in recent years. Excess weight is the most common medical condition in companion animals and has a number of health and wellness implications for both pets and their owners. Obese pets also increase health risks, including an increased risk of arthritis, diabetes mellitus, hepatic lipidosis and insulin resistance. Research has shown that the risk for development of diabetes increases about 2-fold in overweight pets and about 4-fold in obese pets. Altered adipokine secretion appears to be an important mechanism for the link between excess BW and many diseases. In addition, adipose tissue is an active producer of hormones, such as resistin, leptin and cytokines, including many inflammatory cytokines such as tumor necrosis factor-α, IL-1β and IL-6, and C-reactive protein. The persistent low-grade inflammation secondary to obesity is thought to play a causal role in chronic diseases such as osteoarthritis, cardiovascular disease, diabetes mellitus, and others. For example, tumor necrosis factor-α alters insulin sensitivity by blocking the activation of insulin receptors. In addition, obesity is associated with increased oxidative stress, which also may contribute to obesity related diseases. Management of obesity involves nutritional change as well as behavioral change. The emerging science of Nutrigenomics considers alternative mechanisms by which plant-derived phytochemicals may to influence signaling pathways within cells through of genes expression. Cinnamom (C. zeylanicum) could have beneficial health properties such as anti-inflammatory effects, anti-microbial activity, blood glucose control, decrease the total and LDL cholesterol levels, and reducing cardiovascular disease. Cinnamon potentiates insulin-regulated glucose utilization by enhancing insulin signaling, and it enhances cellular glucose uptake due to the stimulation of the insulin receptor activity by increasing the concentrations of the phosphorylated intracellular protein IRS-1 and the increasing of PI 3-kinase and the GLUT4 transporter. The future of nutrigenomics will be to generate natural alternatives to prevent and treat several diseases, such as dog’s obesity, because pets have a special place in the hearts of their human companions.
Apolipoprotein E Variants and Sex Modify the Action of Dairy Fat Intake on Plasma HDL-Cholesterol and Adiponectin Concentration

Ethendhar Rajendiran1, Yongbo She1, Maude Tessier-Grenier2,3, Didier Brassard2,3, Vanu Ramprasath1, Iris Gigleux2,3, Emile Levy3,4, Angelo Tremblay2,5, Patrick Couture2,6, Benoît Lamarche2,3 Peter Eck1, Peter J.H. Jones1

1Richardson Centre for Functional Foods and Nutraceuticals, University of Manitoba; 2School of Nutrition, Laval University; 3INAF, Laval University; 4CHU Sainte-Justine Research Center; 5Department of Kinesiology, Faculty of Medicine, Laval University, Canada

The health benefits of dairy saturated fat (DSF) intake remains unresolved. Apolipoprotein E (APOE) variants modify plasma lipid concentrations (PLC). However, how APOE variants and sex impact the actions of DSF consumption on PLC are not well understood. The present objective was to evaluate effects of APOE (E2E3, E3 and E3E4) variant and sex in altering the effect of DSF intake on PLC. Using a multicenter randomized crossover trial, 92 women and men consumed five isocaloric diets (35% energy from fats) for 4-wk/4-wk washout between diets that included; (i) cheese, (ii) butter, (iii) monounsaturated fatty acids (MUFA), (iv) polyunsaturated fatty acids (PUFA) and (v) a low-fat, high-carbohydrate (CHO). Results suggested the overall diet/APOE/sex interaction was significant on plasma HDL-C and adiponectin, not on plasma total cholesterol or LDL-C levels. However, HDL-C levels were increased in women as a function of diet and APOE-E3E4 variant compared to other fat intake. In summary, women-E3E4 with cheese intake showed high levels of adiponectin compared to men-E3E4 with butter intake. Our findings demonstrate the importance of evaluating APOE variants and sex effect among the diet interaction related to cardiometabolic risk (CMR). However, DSF from either butter or cheese seems to have a beneficial effect on plasma HDL-C and adiponectin levels in women with APOE-E3E4 variant compared to men with APOE-E3E4 variant compared to other fat intake. In summary, APOE variants and sex impact the role of DSF from the dairy on other CMR factors.
Association of Gly482Ser PPARGC1A Gene Variant with Differential Cholesterol Outcomes in Response to Two Energy-Restricted Diets Among Subjects with Excessive Weight

O. Ramos-López1, J.I. Riezu-Boj1,2, F.I. Milagro1,2, L. Goni1, M. Cuervo1,2, J.A. Martínez1,2,3
1Centre for Nutrition Research/Department of Nutrition, Food Sciences and Physiology, University of Navarra, Pamplona, Spain; 2Navarra Institute for Health Research (IdiSNA), Pamplona, Spain; 3CIBERObn, Centro de Investigación Biomédica en Red de la Fisiopatología de la Obesidad y Nutrición, Carlos III Health Institute, Madrid, Spain; 4Madrid Institute of Advanced Studies (IMDEA Food), Madrid, Spain

Background: Precision personalized dietary strategies based on nutrigenetics involves the understanding of individual reactions to different treatments depending on the genetic make-up.

Objective: To investigate the influence of two PPARGC1A gene polymorphisms on metabolic outcomes in response to two energy-restricted diets.

Methods: A four-month nutritional intervention was conducted involving two different hypocaloric diets (-30% of energy), based on a low-fat (LF) and on a moderately high-protein (MHP) dietary patterns. Unrelated subjects with overweight and obesity were genotyped for two PPARGC1A polymorphisms: rs8192678 (Gly482Ser) and rs3755863 (G>A). Genotyping was performed by next-generation sequencing and haplotypes were phenotypically screened. Anthropometric measurements and standard biochemical tests were assessed by validated methods.

Results: Differential cholesterol outcomes were found according to diets and Gly482Ser genotypes. Interestingly, Gly482Gly homozygotes following a LF diet had lower reductions in total cholesterol (-9 mg/dL vs. -27 mg/dL, p=0.017) and LDL-c levels (-5 mg/dL vs. -18 mg/dL, p=0.017) and LDL-c levels (-5 mg/dL vs. -18 mg/dL, p=0.017) and LDL-c levels (-5 mg/dL vs. -18 mg/dL, p=0.017) than 482Ser allele carriers. However, this finding was not evidenced in the MHP group, where Gly482Gly homozygotes underwent similar cholesterol decreases than 482Ser allele carriers. Likewise, all genotype carriers had significant reductions in the frequencies of hypercholesterolemia (total cholesterol ≥200 mg/dL), and A2/A2 (36.1%). BMI was similar regardless ANKK1/DRD2 genotypes (p=0.884). The A1/A2 genotype conferred greater risk for lower frequency consumption of legumes and vegetables than A1/A1 and A2/A2 genotypes (OR = 2.24, 95%CI 1.39–3.58), p = 0.008 respectively). Energy intake did not differ by genotypes (p=0.884). The A1/A2 genotype conferred greater risk for lower frequency consumption of legumes and vegetables than A1/A1 and A2/A2 genotypes (OR = 1.88, 95%CI 1.15–3.11), p = 0.016 (95%CI 1.15–3.11), p = 0.016; and OR = 2.65, 95%CI 1.32–5.32, p = 0.006 respectively).

Conclusions: An energy-restricted MHP diet might be more beneficial than a LF diet to reduce serum cholesterol among PPARGC1A Gly482Gly genotype carriers. The analysis of this genetic variant might be the basis for a precise nutrigenetic management of hypercholesterolemia based on the genetic make-up.

Polymorphism in ANKK1/DRD2 Gene Associates with Unhealthy Eating Behaviors and Metabolic Disturbances

Ingrid Rivera-Iñiguez1,2, Maricruz Sepulveda-Villegas1,2, Claudia Ojeda-Granados1,2, Sonia Roman1,2, Arturo Panduro1,2
1Department of Molecular Biology in Medicine, Civil Hospital of Guadalajara, “Fray Antonio Alcalde”; 2Health Sciences Centre, University of Guadalajara, Mexico

Background: Unhealthy food behaviors affect weight control management. Environmental, social and genetic factors contribute to food decision-making at the brain reward system. ANKK1/DRD2 TaqA1 (rs1800497) polymorphism affects receptor density while increasing L-DOPA activity in the brain reward system. This polymorphism has been associated with unhealthy behaviors such as gambling, alcohol abuse and substance dependence. A1 allele has been related to weight gain. However little is known about the influence of ANKK1/DRD2 TaqA1 on dietary quality.

Objective: To identify the association of ANKK1/DRD2 TaqA1 (rs1800497) with unhealthy food intake in West Mexico.

Methods: A cross-sectional study, with 549 subjects from West Mexico was analyzed for ANKK1/DRD2 TaqA1 genotypes (A1/A1, A1/A2, A2/A2) by an allelic discrimination assay. Anthropometric measurements and standard biochemical tests were performed.

Results: Subjects were 37.04 ± 13.19 years old. Genotype frequency distribution in West Mexico was A1/A1 (17.3%), A1/A2 (46.6%), and A2/A2 (36.1%). BMI was similar regardless ANKK1/DRD2 genotypes (p=0.695). Triglycerides levels were significantly higher among the A1/A1 genotype (p=0.047). Energy intake did not differ by genotypes (p=0.884). The A1/A2 genotype conferred greater risk for lower frequency consumption of legumes and vegetables than A1/A1 and A2/A2 genotypes (OR = 1.88, 95%CI 1.15–3.11), p = 0.016; and OR = 2.65, 95%CI 1.32–5.32, p = 0.006 respectively).

Conclusions: Even though energy intake and BMI did not differ by genotypes, the TaqA1 (rs1800497) polymorphism could be a risk factor for detrimental dietary quality by inducing unhealthy food behaviors that may predispose to metabolic disturbances.
Folic Acid and Other Methyl Donors Attenuate LPS-Induced Gene Expression of Proinflammatory Cytokines in Human Macrophages

M. Samblas¹, F.I. Milagro¹², M.A. Zulet¹²³, J.A. Martínez²³⁴

¹Centre for Nutrition Research/Department of Nutrition, Food Sciences and Physiology, University of Navarra, Pamplona, Spain; ²CIBERObn, Centro de Investigación Biomédica en Red de la Fisiopatología de la Obesidad y Nutrición, Carlos III Health Institute, Madrid, Spain; ³Navarra Institute for Health Research (IdiSNA), Pamplona, Spain; ⁴Madrid Institute of Advanced Studies (IMDEA Food), Madrid, Spain

Background and Objectives: Methyl donors participate in the epigenetics mechanisms by modulating the methylation levels of DNA and proteins. Moreover, methylating compounds trigger essential roles in metabolism and cellular homeostasis. Dietary supplementation with folic acid and other methyl donors has been reported to elicit beneficial effects on obesity-related manifestations and cardiovascular diseases. The aim of this study was to investigate the effects of methyl donors on the inflammatory process in the context of obesity.

Methods: White blood cells and plasma samples were obtained from 97 obese subjects (mean BMI=35.8±4.7 kg/m²) that participated in the Metabolic Syndrome Reduction in Navarra (RESMENA) controlled trial. Methyl donor levels (folic acid and vitamin B₁₂) and inflammatory markers were quantified in plasma. Expression levels of inflammation-related genes were quantified in white blood cells. THP-1 cells (human monocytes) were differentiated into macrophages with 25 ng/ml of phorbol 12-myristate 13-acetate (TPA) and incubation with 11 μM folic acid or a mix of methyl donors (folic acid 11 μM, choline 100 μM, and vitamin B₁₂ 18 nM) for 24h.

Results: Inverse correlations were found between plasma folic acid and vitamin B₁₂ concentrations, with TNFα and SERPINE1 expression in white blood cells, as well as with circulating PAI-1 levels. Incubation of THP-1 activated macrophages with folic acid or a mix of methyl donors inhibited significantly IL1B, TNFα, SERPINE1 and TLR4 gene expression. Moreover, methyl donors decreased IL1B protein levels.

Conclusions: This study reveals a negative association between methyl donor levels in blood and important inflammatory markers. Moreover, the addition of folic acid or a mix of methyl donors (folic acid, choline, and vitamin B₁₂) to human macrophages activated with LPS reduced inflammatory gene expression, suggesting a role of methyl donors in obesity-related inflammation.

Caffeine Consumption and Disease Risk Modulated by Genetic Variants in Caffeine Metabolism

Maria Schellenberger¹

¹University of Southern California, USA

There are many chronic conditions which have been linked with caffeine consumption, including positive associations with hypertension and impaired glucose control, and negative associations with Parkinson’s disease and certain cancers. Caffeine is found in a variety of commonly consumed food products including coffee, soft drinks, tea, energy drinks, caffeinated sports drinks, cocoa, chocolate, dietary supplements, and certain medications. Many enzymes are involved in caffeine metabolism, but between 90-95% is metabolized by cytochrome P450 1A2 (CYP1A2). One SNP of particular importance in influencing the inducibility of CYP1A2 is Rs762551 (aka -164A>C or -163C>A). For this review, 10 studies were consulted to investigate the impacts of caffeine consumption, genetic variation, and disease risk. Findings indicate there may be varying risks of chronic diseases associated with the SNP CYP1A2, however the risks and benefits are not consistent for each genetic variation. There appears to be a slight increased protective effect of caffeine consumption on Parkinson’s disease among slow metabolizers. However, research suggests that slow metabolizers with high caffeine consumption are at increased risk for MI and impaired fasting glucose. Fast metabolizers, on the other hand, may benefit from protective effects of caffeine in preventing hypertension and enjoy higher degrees of ergogenic effects, yet may be at higher risk for bone mineral density loss with increased caffeine consumption. The knowledge of this particular SNP may assist the patient if they are at higher risk for a specific chronic condition (ie diabetes, hypertension, osteoporosis) and are interested in reducing their risk for one specific condition. Due to the various degrees of risk and benefits between these chronic conditions for fast and slow metabolizers, it is difficult to make generalized suggestions for patients regarding their caffeine consumption.
Inter-tissue communication via secreted proteins has been established as a vital mechanism for proper physiologic homeostasis. We have developed a bioinformatics framework using a mouse reference population, the Hybrid Mouse Diversity Panel (HMDP), which integrates multi-tissue expression arrays and publicly available resources to identify and functionally annotate novel tissue-tissue communication. We validate this method by showing that we can identify known, as well as novel endocrine factors responsible for communication between tissues. We further show the utility of this approach by identifying and uncovering in vivo evidence for 7 novel inter-tissue communicatory axes. The developed methodology is generally applicable to any collection of organs, tissues, or cell types sampled from a genetically diverse population and profiled using omics platforms.
Whole Egg Consumption Does Not Alter Expression of Genes Related to Cholesterol Homeostasis or Inflammation in PBMC of Overweight, Postmenopausal Women

Allison M. Stevens¹, Lisa R. Sawrey-Kubiczek¹, Chenghao Zhu¹, Jody M. Randolph¹, Angela M. Zivkovic¹, Francene M. Steinberg¹
¹Department of Nutrition, University of California, Davis, CA, USA

Introduction: Eggs yolks are rich sources of lipids including cholesterol, ceramides, sphingolipids, and phospholipids. These bioactive compounds have been shown to impact the lipidome and proteome of lipoproteins, potentially altering cholesterol homeostasis and inflammation in the body. We investigated whether consumption of two whole eggs for four weeks altered expression of genes related to cholesterol flux and inflammation in peripheral blood mononuclear cells (PBMC) in overweight, postmenopausal women.

Methods: Using a randomized, controlled, crossover design, 20 participants (BMI 28.4 ± 2.9) were randomized to consume either a daily breakfast containing two whole eggs or the equivalent amount of egg whites for four weeks separated by a four-week washout period. PBMC were isolated from fasted blood before and after each dietary intervention for RNA extraction. Targeted gene expression analyses were assessed using qPCR.

Results: Expression of genes related to cholesterol synthesis and transport in PBMC including 3-hydroxy-3-methylglutaryl-CoA reductase (HMGCR), ATP binding cassette subfamily A member 1 (ABCA1), ATP binding cassette subfamily G member 1 (ABCG1), and ADP ribosylation factor like GTPase 4C (ARL4C) expression levels did not differ between whole egg and egg white interventions. Higher PBMC expression levels of pro-inflammatory cytokines tumor necrosis factor (TNF) and interleukin 1 beta (IL1B) were observed following the whole egg intervention compared with egg white, however these changes were not statistically significant (p = 0.06 and p = 0.12, respectively). Similarly, toll like receptor 4 (TLR4) expression levels did not differ between interventions.

Conclusions: Expression of genes related to cholesterol homeostasis in PBMC were unaffected by consumption of two eggs per day for four weeks in postmenopausal women. IL1B and TNF demonstrated more responsiveness to the dietary intervention with trends toward increased expression during the egg white intervention versus the whole egg intervention, however these changes were not statistically significant.
5-methyltetrahydrofolate. Individuals with mutations in the MTHFR gene have impaired folate metabolism, which is associated with hyperhomocysteinemia, DNA strand breaks, and aberrant DNA methylation. Imbalanced DNA-methylation is commonly associated with colorectal cancer (CRC), however, studies suggest that the MTHFR mutation may actually provide protection against the disease by increasing the availability of 5,10-methylenetetrahydrofolate, which aids in nucleotide synthesis and decreases DNA breakage. This review examines the available literature on this complicated connection.

**Methods:** Web of Science and PubMed databases were searched using the terms “MTHFR mutation and colorectal cancer” and “rs1801133 and colorectal cancer.” Ten studies were selected, including one cohort study, three case-control studies, and six meta-analyses.

**Results:** Of the six meta-analyses, four found significant inverse associations between TT/CT genotypes and CRC risk compared to the CC genotype, one found a small significant positive association between the TT genotype and CRC risk, and one found no association. All three of the case-control studies found an inverse risk of CRC for individuals with the TT mutation. However, the cohort study observed an increased risk of mortality for individuals with CRC who have a homozygous mutation (TT, HR: 2.50) as well as those with a heterozygous mutation (CT, HR: 1.85).

**Conclusion:** Folate likely plays a dose and time dependent role in the development of CRC and may modify the effects of MTHFR mutations. More research is needed on the association between mutations and CRC, taking into account folate status, along with other factors involved in folate metabolism including vitamin B2, B12, and homocysteine.

**P57 Cardiovacular Risk in CKD Patients Affected by Arterial Calcifications (AC)**

**Annalisa Terranegra**<sup>1</sup>, **Massimo Mezzavilla**<sup>1</sup>, **Sara Tomel**<sup>1</sup>, **Lee Silcock**<sup>1</sup>, **Caterina Brasacchio**<sup>2</sup>, **Laura Soldati**<sup>2</sup>, **Teresa Arcidiacono**<sup>3</sup>, **Lorenza Macrina**<sup>2</sup>, **Giuseppe Vezzoli**<sup>2</sup>

<sup>1</sup>Sidra Medical and Research Center, Qatar; <sup>2</sup>Università degli Studi di Milano, Italy; <sup>3</sup>Ospedale San Raffaele, Milan, Italy

Chronic Kidney Disease (CKD) patients at different stages suffer from high risk of cardiovascular mortality that is higher in dialysis patients than in general population. The current study recruited 208 CKD patients in hemodialysis and collected dietary 24hrs recalls, medical history, biochemical parameters and blood samples for genetic analyses. The main aim is to identify genetic and nutritional factors which could increase the risk of cardiovascular events in CKD. SNPs genotyping was performed using Illumina® MegaEX Array (2M SNPs) comparing patients with previous cardiovascular events (CV1; 135 subjects) vs patients without cardiovascular events (CV0; 73 subjects). For each marker we calculated the Fixation Index (Fst) using the CV1 and CV0 as subpopulations in order to identify the most differentiated SNPs between the two subgroups. Using the cut-off of the 99th percentile of the Fst distribution we identified 3218 SNPs as differentiated between CV1 and CV0. We interrogated two functional pathway tools (Panther and Ingenuity Pathway Analysis - IPA) to define any possible biological/pathological role of the associated SNPs. We merged the most significant results from the two tools and we identified two genes involved in both cardiovascular disease pathways (from IPA) and hearth development (Panther): Glycagon-like peptide-1 receptor (GLP1R) and Sarcoglycan delta (SGCD). Using regression tree analyses we found that GLP1R rs10305445 minor allele A is associated with higher percentages of CV1 (p=0.03) and, inversely, SGCD rs145292439 minor allele A is associated with lower percentages of CV1 (p=0.038) and higher levels of HDL (p=0.015). We did not find any significant difference in the dietary intake between the two groups as well as with the genetic data and the biochemical parameters. From this preliminary analysis, we can conclude that the genetic factors may contribute more than diet to cardiovascular disease in CKD patients and we can speculate a role of genetic risk marker for GLP1R as well as a protective marker for SGCD in CKD.

**P58 Nutrigenetic Approach: Does Vitamin D Status, CYP27B1 rs10877012 and Its mRNA Levels Influence Essential Hypertension in Indian Population?**

**Varsha Varakantham**<sup>1</sup>, **Ashok Kumar Kurakula Sailoo**<sup>2</sup>, **Balakrishna Nagalla**<sup>3</sup>, **Dinesh Kumar Bharatraj**<sup>1</sup>

<sup>1</sup>Food and Drug Toxicology Research Centre, National Institute of Nutrition, Hyderabad, India; <sup>2</sup>Former Head of Department, Department of General Medicine, Gandhi Hospital, Secunderabad, India; <sup>3</sup>Department of Biostatistics, National Institute of Nutrition, Hyderabad, India

CYP27B1 (25-OH vitamin D 1a hydroxylase) is the rate-limiting enzyme for bioactivation of 1,25(OH)2 vitamin D from 25(OH) vitamin D, and it is well known that low levels of vitamin D can lead to increased renal renin production, thus activating the renin-angiotensin-aldosterone system, thereby increasing blood pressure. Therefore, we have conducted a preliminary investigation in 292 hypertensives and 324 controls to evaluate the association of the genetic variant of CYP27B1 rs10877012, its’ mRNA expression and 25(OH) vitamin D in Indian population with essential hypertension. The percentage of rs10877012 GG, GT and TT in controls are 19.8%, 54.3%, 25.9% and in hypertensives 21.9%, 50.7%, 27.4% respectively. It is interesting to note that, we determined T (Allele frequency=0.53) as the major allele, whereas it is found to be the minor allele in all the studies done so far in various other populations. The highest allele frequency till now for T allele (0.49) was observed in Bengali in Bangladesh (BEB) population derived from 1000 genome project phase 3. In multivariate logistic analysis after adjustment for covariates including age, gender and body mass index (BMI), no association between the CYP27B1 rs10877012 gene variants and essential hypertension was seen (p>0.05). Only 6.4% and 3.7% 25(OH) Vitamin D sufficiency (≥30 ng/mL) was observed in hypertensives and controls, although India has abundant sunlight. In addition, we found no association of
P60

Cellular Energy Status Affects Carbon Metabolism Gene Expression and Has Metabolic and Possible Clinical Consequences

Alain de J. Hernandez-Vazquez¹, Josue Andres Garcia-Sanchez², Ana Salvador-Adriano¹, Noe Alberto Ramirez-Gomez¹, Elizabeth Moreno-Arriola¹, Daniel Ortega-Cuellar², Antonio Velázquez-Arellano¹

¹Nutritional Genetics Unit, Institute of Biomedical Investigations UNAM and ²Research Division, Instituto Nacional de Pediatria; Mexico City, Mexico

We found that interfering with ATP production through reduction of oxidative phosphorylation (OXPHOS) by privation of essential cofactors (biotin, thiamine), decreases the mRNAs and proteins for liver glycolysis and lipogenesis, and increases those for fatty acid oxidation and gluconeogenesis. These effects are mediated by the eukaryotic energy sensor AMPK, except in the case of hepatic gluco-

25(OH) Vitamin D status with essential hypertension (controls: 14.1±8.2 ng/mL versus cases: 13.9±7.5 ng/mL, p<0.05). Even the mRNA expression levels, failed to show any differences based on the genetic polymorphism between hypertensives and controls. In conclusion, these results clearly suggest that Vitamin D status and genetic polymorphism between hypertensives and controls. In conclusion, these results clearly suggest that Vitamin D status and genetic polymorphism between hypertensives and controls.

Further, robust nutrigenetics studies including more number of genes of Vitamin D metabolic pathway and renin-angiotensin-aldosterone system with higher sample size are warranted to draw specific conclusions.

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P61

Improvements in Long-Term Weight Loss and Clinical Parameters with the Use of Nutrigenetics in a 2-Year Prospective Study

Maria Vranceanu¹, Lorena Filip², Horatiu Albu², Ioana Ecaterina Pralea³, Marco Perricone⁴, Keith Grimaldi⁵

¹Department of Nutrigenetics and Personalized Nutrition, Eurogenetica, Milan, Italy, ²Department of Bromatology and Hygiene, Iuliu Hatieganu University of Medicine and Pharmacy, Cluj Napoca, Romania, ³Department of Molecular Nutrition, Columnna Medical Center, Bucharest, Romania, Department of Drug Analysis, ⁴Iuliu Hatieganu University of Medicine and Pharmacy, Cluj Napoca, Romania, ⁵Department of Nutrigenetics and Personalized Nutrition, Eurogenetica, Rome, Italy, ⁶Department of Nutrigenetics, Eurogenetica, London, UK

Objectives: Genetic variation is known may influence dietary requirements, giving rise to the new field of nutritional genomics and raising the possibility of individualizing nutritional intake for optimal health, disease prevention and better weight management on the basis of an individual’s genome. This study investigated whether the inclusion of genetic information to personalize a patient’s diet (nutrigenetics) could improve long-term weight management.

Methods: Two groups of patients attending a weight management clinic were prospectively studied. The ketogenic group consisted of 53 patients followed for 24 weeks a ketogenic diet plan with 1600 kcal. The nutrigenetics group consisted of 61 patients were offered a nutrigenetic test screening 26 variants in 24 genes involved in metabolism. This group followed a personalized diet with 1600 kcal too and all recommendations based on their DNA. Weight, BMI, total cholesterol, HDL cholesterol and fasting blood sugar levels were monitored.

Results: Both diets group performed well over the 24 weeks but after 2 years the nutrigenetic group fared better on the clinical values of plasma glucose, total cholesterol and HDL. Furthermore after 2 years 75% of the nutrigenetic patients had maintained weight loss compared to 21% in the non-genetic group.

Conclusions: Addition of nutrigenetically tailored diets in the weight loss phase and the general healthy eating for life phase resulted in better longer-term BMI reduction and improvements in blood glucose and cholesterol levels.

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P59

The Mexican Roots of Nutrigenetics and Nutrigenomics

Antonio Velazquez-Arellano¹, Hector Bourges-Rodriguez²

¹Nutritional Genetics Unit, Institute of Biomedical Investigations UNAM and ²National Institute of Medical Sciences and Nutrition “Salvador Zubiran”, Mexico City, Mexico

It is surprising that until just three or four decades’ ago, collaborations between nutritionists and geneticists were minimal. The developments of molecular biology and of genomics have blurred this false dichotomy. We put forward a historical perspective on this conceptual transformation, that may be considered as revolutionary, and show that some of its origins emerged in Mexico during the 1980s. This presentation focuses on the Genetic Factors in Nutrition workshop in Mexico in 1982 and the ensuing book published in 1984. Further Mexican and international developments are briefly mentioned.

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11th Congress of the International Society of Nutrigenetics/Nutrigenomics (ISNN)
Abstracts

P62

Association and Gene-Environment Interaction Effect of AGTR1 and VEGFR-2 Gene Polymorphisms with Mental Health on Metabolic Risk Factors of Non-Communicable Diseases in Chinese Malaysian Female Adults

Roseline Wai Kuan Yap1, Mei-Hua Lin2, Yoshihiro Shidoji3, Wai Sum Yap4

1School of Biosciences, Taylor’s University, Malaysia; 2Faculty of Science & Technology/Psychology, Sunway University, Malaysia; 3School of Health Science, University of Nagasaki, Japan; 4Faculty of Applied Sciences, UCSI University, Malaysia

Female adults in Malaysia have a higher risk of chronic non-communicable diseases (NCDs) and mental health problems. Associations of rs5186 in angiotensin II type 1 receptor (AGTR1) and rs2071559 of vascular endothelial growth factor receptor-2 (VEGFR-2) with metabolic risk factors of NCDs were reported in Asian populations. Hence, this study examined the association and interaction effects of AGTR1 (rs5186) and VEGFR-2 (rs2071559) gene polymorphisms with mental health on metabolic risk factors of NCDs in Chinese Malaysian female adults. Anthropometric measurements: body mass index (BMI), body fat percentage (BFP), blood pressure; and biomarkers: fasting blood glucose (FBP), glycated hemoglobin (HbA1c), total cholesterol, triglycerides, low-density lipoprotein cholesterol, high-density lipoprotein cholesterol (HDL-C), CA 15-3 and CA 125 were determined. Job Stress Scale (JSS), Depression, Anxiety, and Stress Scale (DASS-21), and Rhode Island Stress and Coping Inventory (RISCI) questionnaires were used to measure job stress, mental health (stress, anxiety and depression), and coping with perceived stress. Scores from both JSS and DASS-21 are positively correlated with the severity of stress, anxiety, and depression. Higher scores from RISCI indicate better coping with perceived stress. Genotyping of rs5186 and rs2071559 was performed by real-time PCR using Taqman probes. A total of 81 Chinese Malaysian female adults aged 30-65 years old were included in the study. The allele frequencies for rs5186 (A; C allele) and rs2071559 (C; T allele) were 0.96; 0.04 and 0.41; 0.59 respectively. The mean scores obtained from stress, anxiety and depression showed mild or normal levels. The mean score for stress, anxiety, and depression were above average. Significant genetic associations include rs5186 with CA15.3 levels (p<0.001) and rs2071559 with HbA1c levels (p=0.034) after adjusting for potential confounders. Significant correlations were obtained for stress with BMI (r=-0.022; p=0.046), depression with BFP (r=-0.242; p=0.030); and stress coping with FBP (r=0.303; p=0.006). Significant gene-environment interaction effects were obtained for rs2071559 with stress (p=0.015) and rs2071559 with depression (p=0.038) on HDL-C levels. Significant associations and/or interaction effects were obtained in a wide range of metabolic risk factors of chronic NCDs. This adds on to the complexity related to the etiology of these diseases, which warrants further investigations.

P63

MC4R Polymorphism Associated with Antipsychotic-Associated Weight Gain

Lufei Young1, Olivia Dong2,3, Tim Wiltshire2,3 Kevin Kupzyk4, Nicholas Guenzel4

1College of Nursing, Augusta University, Augusta, GA, USA; 2Division of Pharmacotherapy and Experimental Therapeutics, UNC Eshelman School of Pharmacy, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA; 3Center for Pharmacogenomics and Individualized Therapy, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA; 4College of Nursing, University of Nebraska Medical Center, Omaha, NE, USA

Background: Atypical antipsychotic medications have been linked to weight gain and other metabolic complications. Some of the variance in antipsychotic metabolic side effects may be related to genetic factors.

Objectives: Identify genetic risk markers that are predictive of antipsychotic-associated weight gain (AAWG) for patients prescribed atypical antipsychotic drugs.

Methods: A secondary analysis was completed using medical records from 447 patients at outpatient psychiatric clinics in Southeast Nebraska from 2009 to 2016. The IRB at the University Nebraska Medical Center and the clinic’s Ethics Committee approved of this study. The TaqMan single nucleotide polymorphism (SNP) genotyping assay (Life Technologies, Grand Island, New York) was used for genotyping. Medication and genetic information were required to be included in the analysis. An analysis of covariance (ANCOVA) was used to compare BMI mean values, and clinical variables were included as covariates. A Bonferroni correction was applied to adjust for multiple comparisons. The clinical variables included in the model as covariates were: age, gender, medications, diagnoses, and number of chronic conditions. The tested genetic variants included 14 SNPs in 13 candidate genes associated with obesity: SLC6A4, 5-HTTLPR, CACNA1C, ANK3, 5HT2C, MC4R, DRD2, COMT, ADRA2A, MTHFR, BCNF, OPRM1, and GRIK1.

Results: The analysis included 440 subjects (43% male, mean age: 35±19). Approximately 80% of patients had a diagnosis of refractory anxiety disorder followed by major depressive disorder (62%), ADHD (35%), bipolar disorder (17%), PTSD (11%), and schizophrenia (3%). Based on the BMI categories, 152 patients were obese (36%), 129 were overweight (31%), 136 were normal (31%) and 4 were underweight (1%). After adjusting for clinical factors, the AA genotype at rs6265 in the MC4R gene was significantly associated with a 17.61% increase in BMI (F[2, 440] = 6.027, P = 0.003).

Conclusion: The analysis suggests the rs6265 polymorphism in MC4R might be a predictive maker of AAWG. Using predictive genetic markers may help identify those at risk of weight gain to provide targeted nutrition, exercise, and pharmacological interventions. Prospective studies are needed to validate these findings.
Low dosage of Chito-Oligosaccharide (COS) Supplementation Increases the Population of Prevotella in the Cecal Contents of Weanling Pigs

Ting Yu1, Yu Wang1, Shicheng Chen1, Zhiling Wang1, Cui Zhu2, Zongyong Jiang1, Xianyong Ma1, Guozhong Wu1, Zhuang Chen2, Chuntian Zheng1

1Ministry of Agriculture Key Laboratory of Animal Nutrition and Feed Science (South China), Guangdong Public Laboratory of Animal Breeding and Nutrition, Institute of Animal Science, Guangdong Academy of Agricultural Sciences, Guangzhou, China; 2Agro-biological Gene Research Center, Guangdong Academy of Agricultural Sciences, Guangzhou, China. Department of Microbiology and Molecular Genetics, Michigan State University, East Lansing, USA; 3Shanghai institute of applied physics, Chinese Academy of Sciences. Shanghai, China.

Introduction: Chito-oligosaccharide (COS) addition promoted animal growth, increased feed digestibility, and reduced the incidence of diarrhea in weaned piglets. To investigate its influence of COS supplementation on piglets gut microbiota and compare the differences of community composition between COS and antibiotics with ZnO addition, we assessed the cecal microbial community by 16S rRNA gene sequencing with three treatments consisting of basal diet (CTR group), basal diet with COS (COS group), and basal diet with antibiotic and ZnO (AZ group).

Methods: A total of 60 male weaned piglets (Duroc × Landrace × Large White) with an average weight of 6 kg and 21-day old were used in this study. The control (CTR) group was the piglets fed with basal diet; the antibiotics/ZnO (AZ) treatment group was those given the basal diet supplemented with aureomycin (30 mg/kg), polymyxin E (12 mg/kg) and ZnO (3000 mg/kg); the COS group was those given the basal diet supplemented with low-molecular weight (20,000–30,000 Da) of chitosan (50 mg/kg). Out of the 60 animals, each treatment consisted of five replicates and 4 piglets per replicate (totally 20 animals). The feeding trials lasted for 28 days and the contents of cecum were collected.

Results: Beta diversity analysis showed that community structure were distinctly different among above three treatments. Compared with CTR, samples of COS and AZ enriched bacteria affiliated with Prevotella but decreased Firmicutes. Moreover, COS increased the relative abundance of Succinivibrio and Anaerovibrio when compared with AZ. Through microbial functions prediction, COS had higher abundance of functions involved in metabolism of cofactors and vitamins when compared with CTR and AZ. Cecal pH in COS was lower than that in CTR or AZ.

Conclusions: COS could be a natural safe candidate feed additive as an alternative of the feed-grade antibiotics in early-weaned piglets.
Early-Life Exposure to Permethrin Food Contaminant Inducing Neurodegeneration: Is It Matter of Epigenetics?

Laura Bordoni¹, Cinzia Nasuti², Donatella Fedeli², Rosita Gabbianelli²

¹School of Advanced Studied and ²School of Pharmacy, University of Camerino, Italy

Permethrin (PERM) exposure in rats at a dose close to No Observed Adverse Effect Level (NOAEL) for 15 days during neonatal brain development leads to the accumulation of the pesticide long after exposure and to the impairment of dopaminergic system in striatum nucleus, contributing later in life to the burden of oxidative stress and behavioral changes linked to a Parkinson’s like syndrome through several pathways. Among these, epigenetics is a strong candidate mechanism to be involved because of its well-known capability to modulate the effect that environmental factors have on the genome. Previous studies showed alteration of DNMT’s expression regulation and global DNA methylation both in the exposed and in F1 generation. Anyway, the dynamic of the epigenetic contribution to the pathology onset is still not clear. In order to clarify the role of epigenetics in this Parkinson-like syndrome induced by a food contaminant, rats were gavaged through intragastric tube with PERM solubilized in corn oil at a dose of 1/50 of LD50 corresponding to 34.05 mg/kg. The compound was administered daily in the morning from PND6 to PND21 and control group was administered with vehicle. Animals where then sacrificed at 60PND and the expression of genes involved in the dopaminergic pathway, also related in Parkinson onset (Nurr1, TH, α-syn), and genes that regulate epigenetic mechanisms specifically in the brain (Mecp2, TET1, TET2, TET3) was analyzed. Global DNA methylation level assessed by ELISA and histone modifications at specific gene loci investigated through ChIP were also determined in order to clarify the epigenetic landscape. Significant alteration in terms of these markers has been identified in our model, suggesting that epigenetics can exert a pivotal role in the onset of this disease. This study focus on an emerging role of food intake, specifically in early-life, in the regulation of the onset later in life of chronic diseases through intriguing mechanisms such as epigenetics.
Author Index

Numbers refer to abstract number

Abbott, K. P8
Adamski, M. P5
Aguilar-Salinas, C.A. P41
Albu, H. P61
Albuquerque, R.C. P12
Alderete, T.L. P30
Alkayal, F. P6
Amos, A. P7
Araújo, J.N.G. P15
Arcidiacono, T. P57
Arefanian, H. P6
Arguello, L.E. P37
Arias, L. P22
Bailey, S.M. P29
Barreto, A.L. P20
Barrington, W. O1
Batinga, L.B. P17
Beckett, E.L. P8
Bennett, B. O1
Ben-Nissan, D. P9
Bjellaas, T. P39
Björkegren, J.L.M. P52
Bonaldi, A. P20
Bordoni, L. P10, P66
Bourges-Rodriguez, H. P59
Bramucci, M. P10
Brascacchio, C. P57
Brassard, D. P46
Bui, H.H. P39
Cai, W. P65
Campos-Pérez, F. P41
Campos-Pérez, W. P11
Canizales-Quinteros, S. P41
Cantor, R.M. P52
Cárcamo, J.G. P22
Cavadino, A. O3
Chan, B. P54
Cerdà, J. P39
Charugundla, S. P21, P33
Che, N. P21, P33, P52
Chen, S. P64
Chen, Z. P64
Choi, W.-S. P6
Chowanadisai, W. P25
Civelek, M. P52
Collins, J. P5
Cominetti, C. P26
Costa, G.R. P12
Couture, P. P46
Cruz, M.S.M. P15
Cuervo, M. P48
Cunha, A.T.O. P15
Daafar, M. P43
Dang, O. P13, P63
Drevon, C.A. P39, P52
Duarte, M.K.R.N. P15
Duarte, V.H.R. P15
Duerksen-Hughes, P. P16, P18
Eck, B.L.P. P35
Eroshkin, A. P16, P18
Eskin, E. P21
Federi, D. P10, P66
Ferguson, J. P8
Fernandes, C.T.O. P41
Filippov, V. P16, P18
Filippova, M. P16, P18
Fiorini, D. P10
Fisberg, R.M. P19
Fonseca, A.C. P20
Freitas, R.C.C. P15
Fujii, T.M.M. P19, P20
Fuqua, B. P21
Gabbianelli, R. P10, P66
Gallo, M.A. P22
Garcia-Lacarte, M. P23
Garcia-Sanchez, J.A. P60
Garrido, M. P8
Geary, B.A. P30
Geoffroy, C. P22
Ghaskar, R. P52
Gigleux, I. P46
Goncalves, P. P48
González-Aldaco, K. P53
González-Muniesa, P. P36
Gong, L. P30
Goyal, D. P37
Grimaldi, K. P61
Guernzel, N. P63
Guimarães, E. P43
Gunersen, T.E. P39
Guo, M. P48
Gupta, P. P24
Gupta, S. P39
Hart, M.D. P25
Hassan-Brunhstein, Y. P39, P52
Hassimotto, N.M.A. P12
Hazen, S. P41
Hazen, S.L. P34, P39
He, C. P28
Heber, D. O4, P35
Henning, S.M. O4, P35
Hernandez, S. P22
Hernandez-Pando, R. P41
Hernandez-Vazquez, A.J. P52
Hewer, A.L. P39
Horst, M.A. P20, P26
Horvath, S. P44
Hou, S. O1
Hsu, M. O4, P35
Huang, L. P27
Huang, Y. P28
Huertas-Vazquez, A. P41
Hui, S. P21, P33, P41
Hui, S.T. P29, P39
Hwang, D.H. P30
Hypothen, E. O3
Ibarra-González, I. P41
Jaceldo-Siegler, K. P16, P18
Jere鸣e, K.N. P25
Jin, Z. P64
Jones, P.J.H. P46
Jones, R.B. P30
Karani, V. O3
Kim, E.D. P52
Kim, H.-N. P31
King, K. P8
Kohlmeier, M. P13, P65
Koistinen, H.A. P52
Koplev, S. P52
Krauss, N.B.R. P34
Krishnan, K.C. P33, P39, P52
Kupzyk, T.W.K. P63
Kurtz, I. P34
Laakso, M. P52
Lajolo, F.M. P12
Lang, J.M. P34
Lee, J. P39
Lee, R. O4
León-Mimila, P. P41
Levine, M.E. P44
Li, D.Y. P39
Li, Q. P35, O4
Li, Y. O2
Li, Z. P35, O4
Lima, S.C.V.C. P15
Linares, H. P62
Long, J. P35, O4
Lopes, C.R.M. P17
López-Contreras, B. P41
López-Pascual, A. P36
Lorente-Cebrián, S.
Loskutov, V. P16, P18
Lovegrove, J.A. O3, P26
Lu, D. P21
Luca, M.D. P29
Lucchessi, A.D. P15
Lucock, M. P8
Lusis, A.J. P21, P29, P33, P34, P39, P41, P52
Ma, X. P64
Macias-Kauffer, L. P41
Macrina, L. P57
Makin, S. P52
Mansego, M.L. P23
Maraflie, S. P6
Marchioni, D.M.L. P19
Maria-Engler, S.S. P12
Martin, A.A. P52
Martin, C. P8
Martínez, J.A. P52
Martínez, J.A. P36, P48, P50
Martínez-López, E. P11
Mauldin, K. P37
Mehribain, M. P39, P52
Méndez-Sanchez, N. P41
Mendoza-Fandino, G.A. P38, P45
Meng, Y. P52
Mezzavilla, M. P57
Michels, K.B. P55
Milagro, F.I. O5, P23, P48, P50
Mongges, E. P43
Moni, A. P43
Morais, C.C. P26
Moreno-Arriola, E. P60
Moser, M. P39
Murgia, C. P5
Nagalla, B. P58
Nasuti, C. P10, P66
Niblett, S. P8
Nord, M.M. P19
Norheim, F. P21, P33, P39, P52
Novak, R. P39
Ocampo-Medina, E. P41
Ojeda-Granados, C. P42, P49, P53
Author Index

Oliveira, J.M. P15
Oliveira, V.R. P17
Ong, T.P. P12
Ortega-Cuellar, D. P60
Otoluwa, A.S. P43

Palmer, S.L. P30
Pan, C. P21, P29, P33, P39, P52
Panduro, A. P42, P49, P53
Parks, B. P21
Parks, B.W. P39, P52
Pedrosa, T.N. P12
Pérez-Robles, M. P11
Perricone, M. P42, P49, P53
Parks, B. P21
Parks, B.W. P39, P52
Pedrosa, T.N. P12
Pérez-Robles, M. P11
Perricone, M. P42, P49, P53

Qi, H. P33
Qin, H. P27
Quach, A. P44
Quassinti, L. P10
Quintero, M.D. P38, P45
Rajbhandari, P. P52
Rajendiran, E. P46
Ramírez, C. P47
Ramírez-Gomez, N.A. P60
Ramos-López, O. O5, P48
Ramprasad, V. P46
Randolph, J.M. P54
Reue, K. P52
Reyes-Fermín, L. P41
Riezu-Boj, J.I. O5, P48
Rivera-Higuiez, I. P42, P49, P53
Rogero, M.M. P19
Rojas, C.M. O1
Roman, S. P42, P49, P53
Rosenberg, G.M. P52
Sabir, S. P33
Sailoo, A.K.K. P58
Salvador-Adriano, A. P60
Samblas, M. P50
Santos, D.M.C. P15
Savrely-Kubicek, L.R. P54
Schellenberger, M. P51
Seldin, M.M. P52
Sepulveda-Villegas, M. P49, P53
Sepulveda-Villegas, M. P42
Shawaf, E.A. P6
She, Y. P46
Shen, T. P27
Shidoji, Y. P62
Shih, D.M. P52
Silbiger, V.N. P15
Silcock, L. P57
Silva, A.M.G. P15
Smith, B.J. P25
Soldati, L. P57
Song, S.W. P31
Souza, A.L.S. P15
Souza, T.O. P15
Steinberg, F.M. P54
Stevens, A.M. P54
Stiensma, L.T. P55
Strong, M.D. P25
Sun, C. O2
Surendran, S. P26
Suzuki, O. P13
Swain, T.M. P29
Tabaie, W.E. P56
Tang, T.Z. P25
Tao, T. P28
Terranegra, A. P57
Tessier-Grenier, M. P46
Thaha, A.R. P43
Threadgill, D. O1
Tiss, A. P6
Tomni, S. P57
Tontonoz, P. P52
Torres-Castillo, L. P11
Torrinhas, R.S.M.M. P15
Tremlay, A. P46
Twohig, C. P5
Varakantham, V. P58
Vasquez-Trespalacios, E.M. P38
Vega-Badillo, J. P41
Vela-Amieva, M. P41
Velásquez, L. P45
Velazquez-Arellano, A. P59, P60
Vergnes, L. P52
Veysey, M. P8
Vezzoli, G. P57
Vilhena, C. P20
Vilhena, M. P20
Villamí-Ramírez, H. P41
Vimal, K.S. P26
Vranceanu, M. P61
Vulpe, C. P21
Wang, B. P65
Wang, C. P16, P18
Wang, K.G. P25
Wang, Y. P64
Wang, Z. P64
Wilson Tang, W.H. P39
Wiltshire, T. P13
Wirdjatmanto, B. P43
Woo, S.L. P35
Woo, G. P64
Wu, X. O2
Woo, S.L. P35
Woo, G. P64
Woo, X. O2

Yamada, M. P20
Yan, X. P27
Yang, H. P28
Yang, J. P35, O4
Yang, X. P28
Yap, R.W.K. P62
Yap, W.S. P62
Yatim, H. P43
Yin, Y. P28
Young, L. P63
Yu, T. P64

Zheng, C. P64
Zhou, A. P3
Zhou, S. P28
Zhou, Z. P33
Zhu, C. P54, P64
Zhu, J. P65
Zivkovic, A.M. P54
Zule, M.A. P23, P50

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