Plenary Sessions and Workshops

Program and Time Schedule
PROGRAM OF PLENARY SESSIONS

Genomes, function and emerging application
Date: Sunday 20 July 2008
Chair: Prof. Dr. Bhanu Chowdary
Co-chair: Prof. Dr. Tosso Leeb

8:45 - 9:25 Keynote speaker. Dr. Vivek Mittal, “RNAi in mammalian systems”, CSHL, Cold Spring Harbor, USA.

9:25 - 10:00. Dr. Fiona McCarthy, “GOing from "OMICs" data to biological knowledge in livestock”, Department of Basic Sciences, Mississippi State University, USA

10.00 - 10.30 Break

10.30 - 11:05 Dr. George Liu, “Genomic Variations and Functional Elements in the Bovine Genome”, USDA Bovine Functional Genomics Laboratory, Beltsville MD, USA

11:05 - 11:40 Dr. Carole Charlier, “Highly effective SNP based association mapping and management of recessive defects in livestock”, University of Liège, Liège, Belgium

11:40 - 12:15 Dr. Leif Andersson, “Harvesting time for chicken genomics”, Uppsala University, Uppsala, Sweden.

Genetics of Immune Response
Date: Monday 21 July 2008
Chair: Prof. Dr. Susan Lamont
Co-chair: Dr. Inger Edfors

9:00 - 10:00 Keynote speaker. Dr. W. Wiersinga, "Genetics of autoimmune thyroid disease in humans", Dept of Endocrinology and Metabolism, Academic Medical Center, University of Amsterdam, the Netherlands

10:00 - 10:30 Break

10:30 - 11:00 Dr. Chris Tuggle, "Transcriptomic analysis of immune response to Salmonella infection in the pig", Department of Animal Science, Iowa State University, Ames, Iowa USA

11:00 - 11:30 Dr. James Wynn, "Molecular investigation of resistance to amoebic gill disease in Atlantic salmon", CSIRO Marine and Atmospheric Research, Hobart, Tasmania, Australia

11:30 - 12:00 Dr Rebecca Baxter, "Polymorphisms in DRB3 binding pockets affect immune response to a FMDV peptide in cattle", The Roslin Institute and R(D)SVS, University of Edinburgh
Forensics, Legal Aspects and Technical Challenges in Low Copy Number DNA amplification

Date: Wednesday 23 July 2008
Chair: Dr. Bruce Budowle, Co-chair: Dr. Paolo Garofano

09.00 - 10.00 Keynote speech: Dr. B. Budowle, “Low Copy Number DNA Typing and Considerations for Routine Laboratory Work”. Federal Bureau of Investigation, Laboratory Division, Quantico, VA, USA.

10.00 - 10.25 Break

10.25 - 10.55 Dr. Paolo Garofano, “Interlaboratory experiences: From human cancer research to post-pharmacogenetics in animals”. Coanan, Rome, Italy.


11.55 – 12.05 Dr. Sree Kanthaswamy, “Canine Population Data Generated from a Multi-Plex STR Kit for Use in Forensic Investigation”. University of California, Davis, USA.

Animal Models for Human Diseases

Date: Thursday 24 July 2008
Chair: Prof.dr. Herman Raadsma, Co-chair: Prof.dr. Larry Schook

9:00 - 9:50 Keynote speaker, Dr Chand Khanna, “A comparative approach to identify and understand metastasis-associated genes in osteosarcoma”. CCR National Cancer Institute, NIH, Rocksville, MD, USA.

9.50 - 10.10 Dr Emanuelle Bourneuf, “The pig as a biomedical model for human melanoma”, UMR INRA-CEA LREG, Domaine de Vilvert, bat 320, Jouy-en-Josas, France.

10:10 - 10:40 Break

10.40 - 11.20 Prof Larry Schook, “Large animal models for human diseases”, University of Illinois, 1203 W. Gregory Drive, Urbana, USA

11.20 - 12.00 Dr Imke Tammen, “Animal models for Neuronal Cernoid Lipofuscinosis (NCL)”, Reprogen, Faculty of Veterinary Science, University of Sydney, Camden, Australia
WORKSHOP PROGRAM

The following workshop program is scheduled:

Sunday July 20

13.30-17.30 ISAG-FAO Advisory Group on Animal Genetic Diversity
Chair: Hans Lenstra     J.A.Lenstra@uu.nl
Venue: RAI Auditorium

13.30-14.00 Steffen Weigend, Friedrich-Loeffler-Institute, Neustadt, Germany
Assessment of population structure and genetic diversity in chickens based on molecular markers
14.00-14.30 Paolo Ajmone Marsan, Università Cattolica del S. Cuore, Piacenza, Italy
A global view of livestock biodiversity and conservation
14.30-15.00 Tad S. Sonstegard, Agricultural Research Service, Beltsville, MD, USA
SNP analysis and population structure in cattle breeds
15.00-15.30 Ottmar Distl, University of Veterinary Medicine Hannover, Germany
Genetic diversity in horse breeds
15.30-15.45 Break
15.45-16.00 Paul Boettcher, FAO-AGAP, Rome, Italy
The role of the ISAG-FAO Advisory Group in the implementation of the global plan of action for animal genetic resources
16.00-16.15 Johannes A. Lenstra, Utrecht University, Utrecht, The Netherlands
Metaanalysis of microsatellite data allows a molecular classification of European cattle (poster 5041)
16.15-16.30 Christoff Knorr, University of Göttingen, Göttingen, Germany
Admixtture analysis of European and Chinese pig breeds (poster 5006)
1630-16.45 Ercan Kurar, Selcuk University, Konya, Turkey
In vitro conservation and preliminary molecular characterization of some Turkish native domestic animal genetic resources-I (TURKHAYGEN-I) (poster 5025)
16.45-17.00 Pam Wiener, Roslin Institute, Roslin, UK
Patterns of genetic diversity near genes under selection in cattle (poster 5011)
17.00-17.15 Miika Tapio, ILRI, Nairobi, Kenya
Molecular estimation of effective population size – a new perspective for the assessment of risk status of livestock breeds (poster 5027)

13.30-17.30 Poultry Gene Mapping
Chair: Richard Crooijmans     Richard.Crooijmans@wur.nl
Venue: RAI Room A

No agenda provided.
Sunday July 20 (continued)

13.30 -17.30 Satellite workshop: Domestic Animal Epigenetics
Chair: Ikhide Imumorin ikhumorin@spelman.edu
Co-chair: Hasan Khatib hkhatab@wisc.edu
Venue: RAI Room B

- Chromatin Remodeling in Porcine Embryos
  R.A. Cabot, Purdue University, West Lafayette, USA
- Epigenomic Characterization of Cloned Cattle
  A. Eggen, INRA, France
- The Mouse H19CR: DNA Methylation and Beyond
  C. Gebert, NIH-NICHD, Bethesda, MD, USA
- Imprinted Genes in Bovine Embryo-Fetal Development
  S. Hienleder, U of Adelaide, Adelaide, Australia.
- Comparative In Silico Analysis of Some Imprinted Regions in Cattle
  I.G. Imumorin, Spelman College, Atlanta, USA.
- Sequence Characteristics of Imprinted Genes in Cattle
  H. Khatib, University of Wisconsin, Madison, USA.
- Epigenetic Regulation of Short Chain Fatty Acids in Ruminants
  C-J. Li, USDA-ARS, Beltsville, MD, USA.
- Epigenetic Case Studies in Agricultural Animals
  G. Liu, USDA-ARS, Beltsville, MD, USA.
- Epigenetic Variation in Cloned Swine
  P. Piedrahita, North Carolina State University, Raleigh, USA.
- Molecular Characterization of M16iX Myostatin in Mouse
  J.M. Reecy, Iowa State University, Ames, USA.
- Interaction of H19 and IGF2 on Carcass Traits in Beef Cattle
  S.M. Schmutz, U of Saskatchewan, Saskatoon, Canada.
- MicroRNA-26a Targets Histone Methyltransferase Enhancer of Zeste homolog 2 during Myogenesis
  R. Tellam, CSIRO Livestock Industries, Australia.

13.30 -15.30 Applied Genetics Committee of Companion Animals
Chairs: Leslie Lyons lalyons@ucdavis.edu
Cindy Harper cindy.harper@up.ac.za
Venue: RAI Room C
No agenda provided.

13.30 -16.30 Comparative MHC Steering Committee
Chair: Shirley Ellis Shirley.ellis@bbsrc.ac.uk
Venue: RAI Room D

- Shirley Ellis: Institute for Animal Health, Compton, UK
  Comparative MHC committee: what have we achieved since 2002?
- Steven Marsh: Anthony Nolan Research Institute, London, UK
  IPD-MHC: a database for MHC sequences
- Rebecca Baxter: The Roslin Institute and R(D)SVS University of Edinburgh, UK
  Annotation of the bovine MHC
- Don Miller: Cornell University, Ithaca, USA
  Polymorphism and genomic organisation of the equine MHC class II region
Sunday July 20 (continued)
Comparative MHC Steering Committee (continued)

- Keith Ballingall: Moredun Research Institute, Edinburgh, UK
  Signatures of species hybridisation contributing to MHC diversity in domestic sheep
- Chris Davies: Utah State University, Logan, USA
  The non-classical MHC class I genes of cattle
- Mike Stear: University of Glasgow, UK
  Immunogenetic analysis of nematode infections in sheep
- Joan Lunney: USDA, Beltsville, USA
  Update on SLA genes and their impact on immune and disease interactions

15.30 – 17.30 Genetics of Immune Response
Chair: Herman Raadsma  h.raadsma@usyd.edu.au
Venue: RAI Room C

<table>
<thead>
<tr>
<th>Time</th>
<th>Speaker/Abstract</th>
</tr>
</thead>
<tbody>
<tr>
<td>1530</td>
<td>Herman Raadsma, introduction to workshop</td>
</tr>
<tr>
<td>1535</td>
<td>Marcos Vinicuis Silva, A whole-genome association study of major determinants for parasitic infection in Angus breed.</td>
</tr>
<tr>
<td>1545</td>
<td>Guiguigbaza Kossigan, Detection of signature of selection among 14 cattle European and African populations in genomic regions associated to trypanotolerance</td>
</tr>
<tr>
<td>1555</td>
<td>Nicholas N Jonsson, Selecting cattle for host resistance to tick (<em>Rhipicephalus microplus</em>) infestation – natural and human-directed selection.</td>
</tr>
<tr>
<td>1605</td>
<td>Eduardo Casas, A putative quantitative trait locus on chromosome 20 associated with bovine pathogenic disease Incidence</td>
</tr>
<tr>
<td>1615</td>
<td>Natasha Ellis, Fine mapping of QTL for parasite resistance in sheep</td>
</tr>
<tr>
<td>1625</td>
<td>Tracy Hadfield, High-Resolution RH Mapping of a Potential QTL Region on Sheep Chromosome 9 (OAR9)</td>
</tr>
<tr>
<td>1635</td>
<td>Jens Tetens, Expression of antimicrobial peptides in different localisations of the bovine udder.</td>
</tr>
<tr>
<td>1645</td>
<td>R.J. Leach, The quantification of genetic components controlling observed variability to immunisation with a peptide derived from Foot and Mouth disease Virus</td>
</tr>
<tr>
<td>1655</td>
<td>H. Murua Escobar, Establishment of an in vivo model for canine prostate cancer and development of an AAV mediated gene therapeutic approach</td>
</tr>
<tr>
<td>1705</td>
<td>Yu Gao, The SLA-immune long oligonucleotide set: a new tool for functional studies of immunity and disease resistance in pig</td>
</tr>
<tr>
<td>1715</td>
<td>S Mashima, Immunological Parameters of the Chicken Lines selected for the response in <em>Graft-versus-host reaction in ovo</em></td>
</tr>
<tr>
<td>1725</td>
<td>Chang-Chun Li, Understanding <em>Haemophilus Parasuis</em> Infection in Porcine Spleen Through Transcripomics Approach</td>
</tr>
<tr>
<td>1735</td>
<td>Oliver Jann, Comparative genomics of Toll-like receptor signalling reveals candidate genes for disease resistance traits</td>
</tr>
<tr>
<td>1745</td>
<td>Raadsma Herman, Election workshop committee</td>
</tr>
</tbody>
</table>
Monday July 21 (continued)

13.30-15.00  Satellite workshop: Informal Meeting regarding DNA technology in Llama and Alpaca
Chair: Cecilia Penedo  mctorrespenedo@ucdavis.edu
Venue: RAI Room C
No agenda provided.

15.30-17.30  Comparative Genomics
Chair: Chris Tuggle  cktuggle@iastate.edu
Venue: RAI Auditorium
15.30 – 15.40  Welcome and introduction
15.40 – 16.00  Oliver Jann – Comparative genomics of Toll-like receptor signalling
16.00 – 16.20  Tomasz Zabek – Phylogeny relationship of GDF5 in mammals
16.20 – 16.30  Comparative Genomics Workshop discussion
16.30 – 16.50  Shuhong Zhao – Discovery of miRNAs through comparative genomics
16.50 – 17.10  Strozzi and Williams – Analysis of non-orthologous genes between genomes
17.10 – 17.30  Denis Larkin – Chromosome rearrangements in amniote evolution

15.30-17.30  Cattle Molecular Markers and Parentage Testing
Chair: Leanne van de Goor  lgo@vhladmin.nl
Venue: RAI Room A
-  Welcome
-  Comparison Test 2007-2008
-  Next Comparison Test 2009-2010
-  Use of SNPs for Parentage Testing or Identification – Brent Woodward, Merial Limited, USA
-  Other business
-  Election of Committee
-  Close

15.30-17.30  Satellite workshop: Aquaculture Genomics and Genetics
Chair: Sylvie Quinioi  Sylvie.Quiniou@ARS.USDA.GOV
Venue: RAI Room B
15:30 - John Benzie, Shrimps genomics today.
16:10 - Pierre Boudry, Recent Advances in the development of genomic resources in Crassostrea gigas: towards the sequencing of the Pacific oyster genome.
16:30 - Filip Volckaert, Genomic tools for breeding, fisheries and evolutionary biology of European Sea Bass.
16:50 - Mathias Moen, QTL for resistance against Infectious Pancreatic Necrosis in Atlantic salmon.
17:10 - Mathew Baranski, Use of SNP-chips and selective DNA pooling for identifying disease resistance markers in Atlantic salmon.
17:30 - Sylvie Lapegue, Mapping QTL for disease or stress resistance in oysters.
Monday July 21 (continued)

15.30-17.30  Dog Genome Mapping
Chair Kathryn Graves  ktgraves@email.uky.edu
Venue: RAI Room C

- Leif Andersson "Genome-wide association analysis - a powerful method for finding genes
underlying phenotypic diversity in the dog." 45-60 minute presentation
- Poster Presentations: 15 minutes each plus question time
- - Alan Wilton  Poster 2178 Disease Gene Identification in Dogs: Use of SNP arrays and
candidate genes
- Barbara Zangerl  Poster 2130 CFA9 association study to identify potential modifier loci for
progressive rod-cone degeneration (PRCD) in the dog.
- Cord Drögemüller Poster 2193 Positional cloning of the canine hairless mutation reveals an
essential role for the FOXI3 gene in ectodermal development

Tuesday July 22

08.30-12.00  Cattle, Sheep and Goat Gene Mapping
Chair: Eduardo Casas  Eduardo.Casas@ARS.USDA.GOV
Venue: RAI Auditorium

- Welcome and Business meeting (E. Casas, USDA, ARS, Clay Center, Nebraska, USA).
- The Genome Sequence of Taurine Cattle: A Window to Ruminant Biology and Evolution (H.
Lewin, Institute for Genomic Biology, Urbana, Illinois, USA).
- Dutch Milk Genomics Initiative (A. Schennink, Animal Breeding and Genomics Centre,
Wageningen University, The Netherlands).
- Genome wide association study of feed efficiency in beef cattle (L. Sherman, University of
Alberta, Alberta, Canada).
- Genomic signatures of artificial selection in US Holstein cows (T. Sonstegard, USDA, ARS,
Beltsville, Maryland, USA).
- FecXr, a deletion in the BMP15 locus associated to increased prolificacy in the Rasa Aragonesa
(L. Montenegro, University of Zaragoza, Zaragoza, Spain).
- Gene expression profiles in CNS of naturally scrapie-affected sheep (I. Martin-Burriel,
Universidad de Zaragoza, Zaragoza, Spain).
- Using a 1.5K ovine SNP array to expand the sheep linkage map (J. Maddox, University of
Melbourne, Victoria, Australia).
- Gene discovery for reproduction rate in tropically adapted Australian beef cattle (R. Hawken,
CSIRO St Lucia, Australia).
- Profile of circulating DNA in healthy cattle (J. Beck, Chronix Biomedical GmbH, Goettingen,
Germany).
- Functional analysis of the c.1892+19 T>C SNP of the PPARGCIA gene which is associated with
variation in milk fat yield (R. Fuerbass, Research Institute for the Biology of Farm Animals,
Dummerstorf, Germany).
- Analysis of gene expression in horned and polled tissue from Brahman cattle using Agilent bovine
microarray (M. Mariasegaram, CSIRO St. Lucia, Australia).
- Goat mammary expressed genes representative of immunity function during pregnancy before
acquisition of its secretory phenotype (F. Faucon, INRA Jouy en Josas, France).
Tuesday July 22 (continued)

08.30-12.00 Avian Immunogenetics
Chair: Janet Fulton
Venue: RAI Room A
JFulton@hyline.com
No agenda provided.

08.30-12.00 Animal Forensics
Chair: Sree Kanthaswamy
Venue: RAI Room B
skanthaswamy@ucdavis.edu

- Present Guidelines from Budowle et al. 2005
- Forensic and non-forensic testing
- Inter-lab collaboration
- Proficiency tests v. Comparison tests
- Accreditation/Certification
- Databases
- Nominate new standing committee members
- Other business

08.30-12.00 Meeting of the Editorial Board of Animal Genetics
Moderator: The Editor
Venue: RAI Room C
Chris.Moran@vetsci.usyd.edu.au
No agenda provided.

15.30-17.30 Applied Genetics in Sheep and Goats
Chair: Jill Maddox
jllm@rubens.its.unimelb.edu.au
Venue: RAI Room B

15:30 Sandy Sgorlon, poster 2208: Nutrigenomics studies in sheep using a custom microarray platform
15:45 Amparo Martínez Martínez, poster 2231: Genetic relationships between native goat populations from Spain, Portugal and America
16:00 José Antonio Bouzada Rey, poster 2152: High exclusion probability in ovine genealogical control through 18plex and 20plex PCR
16:15 Jill Maddox on behalf of Mike Heaton and the ISGC "Update on developing sheep SNP markers for parentage testing from the ovine genome project" and "Update on the Sheep HapMap project"
16:30 Susan Dileanis ISAG 2008 Goat Comparison Test Results
16:50 Jill Maddox ISAG 2008 Sheep Comparison Test Results
17:20 General discussion and election of new committee
17:30 End of session
Tuesday July 22 (continued)

15.30 – 17.30  Equine Genetics and Thoroughbred Parentage Testing
   Chair: Alan Guthrie  alan.guthrie@up.ac.za
   Venue: RAI Room A
   No agenda provided.

15.30 – 17.30  Satellite workshop: Coat Colour Genetics
   Chair: Sheila Schmutz  sheila.schmutz@usask.ca
   Venue: RAI Auditorium

   13:30  White patterns in Horse and Cattle
   Tosso Leeb, Switzerland

   13:50  White and Belt Phenotypes in Chinese Pigs (KIT)
   Lusheng S Huang, China

   14:10  The Beta-Defensin Gene in Dogs – Black and Brindle
   Greg Barsh, USA

   14:30  The Mutation for Progressive Gray in Horses
   Leif Andersson, Sweden

   14:50  Lavender plumage colour phenotype in Quail and Chicken : same MLPH gene but very
   different mutations.
   Bertrand Bed'Hom, France

   15:10  Dilute Phenotypes in Cattle (SILV)
   Sheila Schmutz, Canada

Wednesday July 23

15.30-17.30  Domestic Animal Sequencing
   Chair: Lawrence Schook  Schook@uiuc.edu
   Venue: RAI Auditorium

   15:30 to 15:45  Call to order, Review of Previous Meeting and Workshop Objectives, Larry
   Schook

   15:45 to 16:00  Harris Lewin, Bovine Genome Project

   16:00 to 16:15  Richard Crooijmans, Chicken Genome Project

   16:15 to 16:30  Jill Maddox, Ovine Genome Project

   16:30 to 16:45  Ed Smith, Turkey Genome Project

   16:45 to 17:00  Alan Archibald, Pig Genome Sequencing Project

   17:00 to 17:15  Jim Reecy, QTLdb

   17:15 to 17:30  Discussion and Business Meeting
                   - Transition from sequencing to annotation and bioinformatics
                   - Animal phenotypes
Wednesday July 23 (continued)

15.30-17.30  **Horse Genome Mapping**  
Chair: Imke Tammen  
Venue: RAI Room A

15.30  **Leif Andersson* and Claire Wade** on behalf of the Equine Genome Sequencing Consortium  
Preliminary analysis of the equine genome  
20min + 5 min discussion

15.55  **Terje Raudsepp*, Nandina Paria and Bhanu P. Chowdhary**  
Sequencing-ready physical map of the horse Y chromosome  
15min + 5 min discussion

16.15  Suzanne Eivers, Beatrice McGivney, David MacHugh, Cormac Taylor, Katie Mason, James MacLeod, Lisa Katz and **Emmeline Hill*  
Exercise-induced gene expression in equine skeletal muscle.  
10min + 5min discussion

16.30  **Virgine Lampe*, Claudia Dirks and Ottmar Distl**  
Insights into equine osteochondrosis based on genetic markers in Hanoverian warmblood horses  
10min + 5min discussion

16.45  **Sarah Blott*, L.Y. Fox-Clipsham, E. Temperton, J.E. Swinburne and M. Vaudin**  
Single nucleotide polymorphism (SNP) density and predicted mapping resolution of complex trait genes in the Thoroughbred horse  
10min + 5min discussion

17.00  Brief updates  
**Ottmar Distl** - Equine BAC – physical map  
**Bhanu Chowdhary** - Expression arrays

17.10  General discussion and group updates

17.20  Election of chair and steering committee  
* = presenting author

---

**15.30-17.30  Pig Gene Mapping and Applied Genetics**  
Chair: Gary Rohrer  
Venue: RAI Room C

No agenda provided.

---

**15.30-17.30  Early Career Scientist Workshop. Steps to publish a paper in Animal Genetics.**  
Moderators  
Venue: RAI Room D

No agenda provided.
The following poster abstracts have been presented incorrectly in the abstract book:

**Poster 2198, Maternal dietary methionine supplementation modulates foetal hepatic expression profiles**

Presenting Author: Klaus Wimmers, Research Institute for the Biology of Farm Animals (FBN), Research Unit ‘Molecular Biology’, 18196 Dummerstorf, Germany

Other authors (name only):
1. Eduard Murani
2. Wirawan Nuchchanart
3. Siriluck Ponsuksili

**Abstract:**
Impact of maternal nutrition during pregnancy on gene expression profiles and phenotypes of offspring was shown experimentally and in epidemiological studies in model animals and humans. The aim of the study is demonstrating of foetal programming in pigs, listing of genes sensitive to foetal programming and evidencing and quantifying of the role of DNA-methylation in this phenomenon. Therefore, 36 sows were fed methionine supplemented (‘MET’) vs. control (‘CON’) gestation diets. Liver samples of fetuses of three developmental stages (d35, d63, d91 p.c.) were subsequently monitored for diet-dependent differential gene expression by using Affymetrix microrarrays. Statistical and bioinformatics evaluation of the expression profiles revealed largest effect of gestation diets at late foetal stages (91 dpc) when more than 1600 transcripts were differentially regulated. Functional annotation clustering highlighted the GO terms biological quality, cellular and anatomical structure morphogenesis, and negative regulation of metabolic processes. Interestingly, members of the IGF axis were found being diet-dependent regulated that play a key role in prenatal growth and are sensitive to regulation by DNA-methylation. Quantification of the DNA methylation at these loci will shed light on the role of this epigenetic mechanism as the molecular basis of genotype-environment (diet) interactions.

**Poster 2236, Expression profiling and eQTL analysis reveal biological pathways and candidate genes affecting water holding capacity of muscle**

Presenting Author: Siriluck Ponsuksili, Research Institute for the Biology of Farm Animals (FBN), Research Group ‘Functional Genome Analysis’, 18196 Dummerstorf, Germany

Other authors (name only):
1. E. Jonas
2. E. Murani
3. C. Phatsara
4. T. Srikanchai
5. C. Walz
6. M. Schwerin
7. K. Schellander
8. K. Wimmers

**Abstract (maximum 200 words):**
There is considerable variation in the water holding capacity (WHC) of meat, which is an important economic factor in pork production. WHC depends of numerous genetic and environmental factors that affect muscle fibre properties and metabolic processes ante and post slaughter. We aimed to identify candidate genes for WHC by QTL mapping, expression profiling and expressionQTL (eQTL) analysis. Expression microarray analysis of M. longissimus dorsi RNAs of 74 F2 animals of a resource population showed 1,279 transcripts with trait correlated expression. Negatively correlated transcripts were enriched in functional categories and pathways of extracellular matrix, receptor interaction, and Ca-signalling; positively correlated transcripts dominantly represented oxidative phosphorylation, mitochondrial pathways, and transporter activity. A eQTL analysis of trait correlated transcripts revealed 897 eQTL, with 104 eQTL matching previously identified QTL for WHC; 96 transcripts were trans regulated and 8 had cis acting regulation. Based on these findings, a priority list of genes was established out of the orchestra of genes of biological networks relevant to the liability to elevated drip loss. The complex relationships between biological processes in live skeletal muscle and meat quality are mainly influenced by the energy reserves and their utilisation in the muscle and by the muscle structure itself.
Poster 1001

Title: Retention of Intron-1 in Cathelicidin-4 mRNA of Egyptian Native and Frisian Crossbred Cattle

Ahlam A. Abou Mossallam,
Cell Biology Department, National Research Center, El Behoose Street, P.O. Box 12622, Dokki, Cairo. Tel: 002-027605691 Fax: 002-023370931, E-mail: ahlammasry@yahoo.com

Other authors:
1 Eman R. Mahfouz
2. Mona A. Bibars
3. Soheir M. El Nahas

Abstract (maximum 200 words):
The antimicrobial peptide gene cathelicidin-4 (CATHL-4) was investigated in native and Frisian crossbred cattle reared in Egypt. The gene was tested for reaction with cDNA of various cattle tissues such as blood; lung; trachea; intestine; muscle; lymph and liver using PCR. A positive PCR product was seen in blood, lung, trachea, liver and lymph tissues, but not in intestine of both native and crossbred cattle. NCBI-Blast analysis of native and crossbred cattle CATHL-4 cDNA sequences (303 and 316bp respectively) with Bos taurus CATHL-4 mRNA (gi: 31341226) and BTA22 (gi: 76649266) were almost the same, they both showed high sequence identity. Analysis of PCR products of CATHL-4 cDNA revealed the retention of intron-1 in both native and crossbred cattle, which is absent in Bos taurus CATHL-4 mRNA (gi: 31341226). This may have resulted from alternative splicing which increases the genetic diversity of the genome without increasing the overall number of genes. It may play a role in increasing their innate immunity by increasing gene expression, since intron-containing genes are expressed more efficiently than intronless ones. The results indicate that both native and crossbred cattle reared in Egypt may express cathelicidin gene more efficiently than Bos taurus or Bubalus bubalis reared abroad.

Poster 1002

Title: Evaluation as therapeutics horse of KISO pony by behavioral related gene (DRD4 gene) polymorphism

Presenting Author (name & address): HARUTAKA MUKOYAMA, Nippon Veterinary and Life Science University, Tokyo, Japan

Other authors (name only):
1  MARIE ENDO
2  KYOKO KURODA
3  HIROKI FURUTA
4  TATSUYUKI YOSHIDA

Abstract (maximum 200 words):
The ancestor of KISO pony which is a kind of Japanese native horses came from Mongolia grassland horse at the second century. KISO pony has been used as an employment animal for a long term. And this pony is utilized for animal therapy recently. The behavioral related gene polymorphism has been reported to be associated with the personality trait of novelty-seeking in humans. We evaluated the temperament as therapeutics horse of KISO pony by the polymorphism of dopamine D4 receptor (DRD4) gene which is one of behavioral related gene. The sequences of DRD4 exon III was determined 504bp in 21 KISO pony. All KISO pony samples had 9 repeats: no other repeat-type polymorphism of variable number of tandem repeats (VNTR) in DRD4 exon III was found. However, two SNPs was found in the amplified regions (147bp and 292bp).

Poster 1003

Title: Epigenetic Case Studies in Agricultural Animals

George Liu
Abstract:
In many biological processes, the regulation of gene expression involves epigenetic mechanisms. An altered pattern of epigenetic modification is central to many animal diseases. Using animal disease models, we have studied one of the major epigenetic components: DNA methylation. We characterized the impacts of tissue types, ages and generations on the DNA methylation status of multiple genes in disease-resistant and susceptible animals. These results indicate that differential levels of DNA methylation of important candidate genes are related to the distinct traits of the animal models and could contribute to their resistance or susceptibility to diseases. We are also probing recent technological advances, which allow epigenetics to be studied at a genome-wide scale. Now it is time to apply epigenomics approaches to animal epigenetics research.

Poster 1004
Title: Genome wide SNP discovery in pig using 1G Genome Analyzer: How to play around with sequence length, quality level and mapping qualities
Presenting Author (name & address): Andreia J. Amaral - Animal Breeding and Genomics Centre, Wageningen University, P.O. Box 338, Wageningen, 6700 AH, The Netherlands.

Other authors (name only):
1. Hindrik H.D. Kerstens
2. Hendrik-Jan Megens
3. Bert Dibbits
4. Richard P.M.A. Crooijmans
5. Johan T. den Dunnen
6. Martien A.M. Groenen

Abstract (maximum 200 words):
Large scale sequencing of reduced representation libraries using the 1G Genome Analyzer has proven to be a reliable, fast method for the identification of large numbers of SNPs. Challenges arise, due to varying sequence quality, which is crucial during sequence selection. For instance, quality scores decrease with increasing sequence length, thereby increasing the number of unique sequences due to sequencing errors and leading to the identification of false SNPs in the 3’end of the sequence. On the other hand, shorter sequences lead to a higher proportion alignment of paralogous sequences.
This study shows the impact of varying sequence length and quality level on mapping quality and identification of true SNPs. A DNA pool of 5 animals from a boar line was digested with DraI and fragments of around 200bp were end-sequenced using 1G Genome Analyzer, yielding 70,348,064 sequences of 36bp. Output from BUSTARD, was used to select sequences which were uniquely mapped to a reference genome. Sequences with unknown bases and polymers were removed and sequence uniqueness ranged from 35% at 29bp to 55% at 35bp indicating a high error rate. Results, show the reliable identification of large number SNPs in spite of a relatively high error rate.

Poster 1005
Title: CRB GADIE, a Biological Resources Centre dedicated to livestock genomics
Order of the authors: ESQUERRE Diane, MARTHEY Sylvain, NEAU André, GAO Yu, ROGEL-GAILLARD Claire, HUGOT Karine
Presenting Author (name & address): Diane ESQUERRE 1, 2
1 INRA DGA, UMR314, Laboratoire de Radiobiologie et Etude du Génome, Jouy-en-Josas, 78350, France.
2 CEA/DSV/iRCM/SREITT/LREG, Jouy-en-Josas, 78350, France

Other authors (name only):
1. MARTHEY Sylvain
2. NEAU André
3. GAO Yu
4. ROGEL-GAILLARD Claire
5. HUGOT Karine

Abstract (maximum 200 words):
CRB GADIE has a double-objective related to livestock genomics: (1) to maintain and valorize DNA collections and (2) to prospect, develop and distribute high throughput genomics tools. CRB GADIE conserves 2.5 millions BAC and cDNA clones for eight species including pig, chicken, cattle, trout, horse, rabbit, goat and sheep. Until 2001, our team has set up protocols, automates and LIMS to manage and screen libraries in optimal and secure conditions. More than 150 worldwide laboratories have been using INRA libraries thanks to CRB GADIE activities. Until 2004, CRB GADIE produced and distributed generic cDNA and oligonucleotide microarrays dedicated to pig, cattle, chicken and trout species. The CRB GADIE has been identified by REX EADGENE as a reference platform to produce chicken and pig microarrays and is currently working with two other European platforms (ARK-Genomics/Roslin, UK and University of Aahrus, DK) to develop common quality controls. In addition to a generic activity on clone management and chip production, CRB GADIE is involved in the development of promising new tools and methods such as genomic tiling arrays and sequence enrichment of targeted genomic regions by capture on oligonucleotide microarrays. CRB GADIE activities are presented at http://www-crb.jouy.inra.fr/BRC/index.html.

Poster 1006

Title: Automated SNP calling overestimates the number of SNP in the bovine genome

LAERCIO R. PORTO-NETO¹,² and WILLIAM BARENDSE²
Cooperative Research Centre for Beef Genetic Technologies
¹The University of Queensland, School of Animal Studies, St. Lucia 4072, Australia;
²CSIRO Livestock Industries, Queensland Bioscience Precinct, St. Lucia 4067, Australia.

Single nucleotide polymorphisms (SNP) have been identified from the Bovine Genome Sequencing Project and over 2 million have been identified at the Baylor College of Medicine (ftp://ftp.hgsc.bcm.tmc.edu/pub/data/Btaurus/). The SNP were identified using text based methods that included a series of filters as described in the data set. We generated assays for 42 SNP from the genome sequence SNP set with the ABI SNPlex™ technology. We sampled genotypes from 804 animals of taurine dairy and 379 animals of taurine, zebu and composite beef breeds. Two of the SNP assays did not work and 22 of the SNP were monomorphic in all breeds, both taurus and zebu. That gives an estimate of the proportion of 0.45 ± 0.08 (SE) of the genome sequence SNP being valid. There were significant differences in the proportion of successful SNP assays from the three genetic regions that were tested. The sequence traces for the successful SNP, when compared to the traces of the unsuccessful SNP, showed that automated text based SNP calling had overestimated the number of SNP in the genome sequence database. Using a simple algorithm based on the sequence traces we were able to reduce the false calls to zero in our data set.

Financial support: L.R.P.N. is a Ph.D. student at the University of Queensland and receives an Endeavour International Postgraduate Research Scholarship, a UQ International Student Living Allowances and a scholarship from the Beef CRC. W.B. receives financial support from CSIRO, the Beef CRC, and Meat and Livestock Australia.

Poster 1007

Title: From public sequences to custom microarrays: an easy and affordable tool.

Presenting Author (name & address): Lorraine Paret, Department of Animal Production, Università della Tuscia, Viterbo, Italy

Other authors (name only):
1. Giovanni Chillemi
2. Susana Bueno
3. Gianluca Prosperini
4. Silvia Bongiorni
5. Alessio Valentini
Abstract (maximum 200 words):
Expression data of more than 100 species are present in Gene Expression Omnibus at NCBI, with most platforms represented by spotted DNA/cDNA or oligonucleotides. This evidences a weak point in microarray generation, since starting from cDNA libraries or oligonucleotides synthesis implies a considerable lag between the availability of sequences and microarray preparation. In situ generated oligonucleotides are quicker to develop, but being mostly produced by big companies, only species for which there is high interest worldwide are considered.

We developed a pipeline of software instruments that allow to start from unannotated, redundant sequences, as those found in public databases or generated by parallel sequencing, to yield oligonucleotides suitable for in situ generation on chip. As an example, we generated a chip from sheep (Ovis aries) ESTs deposited at NCBI. The chip carries about 23,000 non-redundant features in quadruplicate and represents the first fully annotated sheep microarray with a large genome covering. In slide replicates show a coefficient of variation<0.25 for genes that are differentially expressed with P<0.01 of fold change.

The procedures are being installed in a portal with public access and will allow researchers to develop their own microarray from any species provided there are available sequences.

Poster 1008

Title: The validity of SNP markers for Bovine Parentage Test in Japan


Other authors (name only):
1. KAZUHITO KUROGI
2. YUUSUKE KOZONO
3. SIN-ICHI SHIMANUKI
4. KAZUHIRO SHIMIZU
5. SYOUTA NISHIMURA
6. AKIKO TAKASUGA
7. MITSUO MORITA

Abstract (maximum 200 words):
We have been conducting parentage testing using 17 microsatellite (MS) markers that include ISAG preferred 9 markers, to carry out tests with PE1* scores at > 0.99998. Recently, we have been able to confirm progress in the technology for the SNP typing that has also resulted in lower costs than before. In this study, we selected 91 SNP markers for parentage test in order to calculate excluding probability. The SNP were selected based on the date of 87 Japanese Black and 84 Holsteins genotyped using Bovine Mapping 10K SNP Kit (Affymetrix GeneChip®) under the following conditions, (1) heterozygosity at 0.4, (2) located in a unique position on the genome, (3) 2~4 SNPs that were more than 5.7Mb apart on each chromosomes. Samples were 175 major sires (104 Japanese Black and 71 Holsteins) used in Japan. SNP was genotyped using the Illumina GoldenGateAssay technique, and analyzed by the Illumina BeadStadio3 software. PE1* score using the 91 SNP markers was >0.99999997. Equivalent scores for the 17 MS markers were achieved using 60~65 SNP markers. Finally, we were able to achieve higher degrees or precision on parentage test in Japan using the SNP markers.

PE1*:Given two parents and one offspring, excluding a parent

Poster 1009

Title: Comparative analysis of a BAC-based physical map of the horse genome

Authors: A. Wöhlke1, G. Nordsiek2, M. Scharfe2, M. Jarek2, F. Schrader2, J. Wrede4, J.P. Pook1, B. Zhu3, P. J. De Jong2, B. P. Chowdhary1, T. Lee13, H. Blöcker2, O. Distl1

Presenting Author (name & address): Ottmar Distl, 1Institute of Animal Breeding and Genetics, University of Veterinary Medicine Hannover, Bünteweg 17p, 30559 Hannover, Germany

Abstract (maximum 200 words):
Abstract (maximum 200 words):
The second horse genome assembly is based on whole genome shotgun sequences (Broad Institute) and BAC end sequences (Hannover). We developed a high-resolution BAC-based physical map to further improve the horse genome assembly and to provide further SNPs. This physical map of the horse genome has been created using a combination of fluorescent fingerprinting and end sequencing of 150,000 BAC clones from the CHORI-241 equine BAC library (10X genome coverage). The results of 314,972 BAC end sequences are accessible at http://www.tiho-hannover.de/einricht/zucht/hgp/index.htm and public databases (NCBI, Ensembl). Fluorescent fingerprints were obtained by using the 4-restriction enzyme 4-color technique and separating the resulting fragments on capillary sequencers. Simultaneously with the collection of raw data we optimized the parameters for the assembly of the fingerprint contigs using the software FPMiner 2.0 and FPC V8.5.3. In addition, we used BAC end sequences and horse genome draft sequences for anchoring the fingerprint contigs, for closing gaps and for selecting the minimal tiling path of the physical horse map. So we can compare an assembly solely based on fingerprints with an assembly using fingerprints and equine sequences. We will present an updated status of this fingerprint map and comparative analyses with other genomes.

Poster 1010
Title: The SLA-immune long oligonucleotide set: a new tool for functional studies of immunity and disease resistance in pig

Presenting Author (name & address): Yu GAO 1, 2
1INRA, DGA, UMR314, Laboratoire de Radiobiologie et Etude du Génome, Jouy-en-Josas, France.
2CEA, DSV, iRCM, SREIT, LREG, Jouy-en-Josas, France

Other authors (name only):
1. Laurence FLORI
2. Mark STAM
3. Karine HUGOT
4. Francois LEFEVRE
5. Isabelle OSWALD
6. Claire ROGEL-GAILLARD

Abstract (maximum 200 words):
Several genome wide generic arrays have been developed in pig but they only partially cover the genome and lack many immune response genes. Our aim was to design and validate a long oligonucleotide set dedicated to the immune response. The set comprises 3773 unique probes and includes all genes and putative transcripts localized in the swine leukocyte antigen (SLA) complex region (826 probes) as well as immune response genes outside SLA (2947 probes). The SLA subset contains targeted sense and anti-sense sequences of 407 transcripts and 6 non-coding RNA genes. The SLA-immune set represents 3104 porcine genes or transcripts among which 95% have one assigned GO term at least. The immune response pathways referred in KEGG are all covered and 38 other pathways are represented. The SLA-immune set was spotted onto glass slides together with the Qiagen-NRSP8 set and a series of control elements. We have analyzed transcriptome variations between unstimulated PBMCs and PBMCs stimulated by either LPS or PMA/ionomycin. We will present our results using this generic chip enriched with probes targeting immune response and will discuss the porcine SLA-immune oligonucleotide set as a promising tool for studying immunity and disease resistance in pig.

Poster 1011
Title: Development of a 25-plex SNP assay for traceability in cattle

Presenting Author (name & address): Dr. Baruch Karniol. Bactochem LTD 2 Hacharash St. Ness-ziona 70400 ISRAEL Tel.- 972 8 9308308 Fax.- 972 8 9401439 bkarniol@gmail.com

Other authors (name only):
1. Andrei Shirak
2. Eyal Baruch
3. Avi Tal
4. Aviv Cahana
5. Michal Kam
6. Yitzchak Skalsky  
7. Charlote Singrün  
8. Gotfried Brem  
9. Joel I. Weller  
10. Micha Ron  
11. Eyal Seroussi  

Abstract (maximum 200 words):  
Single Nucleotide Polymorphisms (SNPs) are amenable to automation and therefore become the marker of choice for DNA profiling. SNaPshot, a primer extension-based method was used to multiplex 25 SNPs that have been previously validated as useful for identity control. Detection of extended products was based on four different fluorochromes and extension primers with oligonucleotide-tails of differing lengths, thus controlling the concise length of the entire chromatogram to 81 bases. Allele frequencies for Holstein, Simmental, Limousine, Angus, Charolais and Tux Cattle were estimated and significant positive Pearson-correlation coefficients were obtained among the analyzed breeds. The probability that two randomly unrelated individuals would share identical genotypes for all 25 loci varied from $10^{-8}$ to $10^{-10}$ for these breeds. For parentage control, the exclusion power was found to be 99.9% when the genotypes of both putative parents are known. A traceability test of duplicated samples indicated a high genotyping precision of greater than 0.998. This was further corroborated by analysis of 60 cases of parent-sib pairs and trio families. The 25-plex SNaPshot assay is adapted for low and high-throughput capacity and thus presents an alternative for DNA-based traceability in the major commercial cattle breeds.

Poster 1012
**Poster 2001**

**Title:** “Nos2A Evaluation of Nos2A & SLC9A3R1 as positional and biological candidate genes for litter size in pigs”

Authors: Amanda Fernández Rodríguez, Animal Genomics Department, INIA, Ctra Coruña km 7.5, 28040 Madrid, Spain

Other authors (name only):
1. Carmen Rodríguez
2. María Martínez Giner
3. Ingrid Balcells
4. Cristina Óvilo
5. Ana Isabel Fernández

**Abstract:**

The aim of the present study was to identify the responsible genes of two epistatic QTL detected on SSC12 for litter size trait in a Meishan x Iberian F2 cross. Two functionally related genes, Slc9a3r and Nos2a, have been selected as functional and positional candidates to underlay these QTL affecting number of piglets born alive and total number of piglets born. Slc9a3r is a factor required for Nos2 isoform activity and both, Slc9a3r and Nos2a variants have been related with reduction of the litter size in rodents. The complete cDNA sequences of both porcine genes have been characterized in four F2 animals. Nine SNPs were detected in Slc9a3r, four of them changing aminoacid composition; and 13 SNPs and three INDELs were detected in Nos2a, five out of them changing aminoacid composition. Physical (IMpRH) and linkage mapping were carried out and both genes were mapped on porcine chromosome 12, within epistatic QTL intervals. Significant associations for Nos2a and Slc9a3r polymorphisms on piglets born alive and total number of piglets born have been found with marker assisted association tests.

**Poster 2002**

**Title:** Genetic study of common variants at the ABCA1 and CETP genes in vervet monkeys

Authors: Gift Chauke and Jürgen Seier

Presenting Author: Gift Chauke, PO box 19070, MRC Primate Unit, Tygerberg, Cape Town, South Africa

Other authors (name only):
1. Peter C. Thomson
2. Herman W. Raadsma

**Abstract:**

In the last few years there has been growing evidence of the influence of genetic variation in the determination of plasma lipid concentrations, especially for genes involved in lipid transport and metabolism. More recently, a pharmacogenetic approach has been applied to polymorphisms in these genes and demonstrated that they are also associated with response to lipid-lowering drugs. The objective of this study was to identify common variations in 2 important genes related to lipid transport and with potential to be genetic determinants of niacin (nicotinic acid) responsiveness in vervet monkeys (Cercopithecus aethiops).

20 monkeys were genotyped for polymorphisms in the genes encoding ATP-binding cassette transport A1 (ABCA1) and cholesteryl transfer ester protein (CETP). An average of 5 single nucleotide polymorphisms (R219K, V771M, V825I, I883M and I405V) was genotyped within these genes. Based on the genotype results, 4 monkeys were selected from the 20 and treated with niacin at an escalating dosage. Their mean lipid-lowering response following drug therapy was examined, compared to those with the same genotype in a placebo group. Data obtained from this study will contribute to potential benefits of using the vervet monkey as a model to evaluate the mechanism of niacin as a therapeutic approach that can exploit the benefits of HDL in coronary heart disease.

**Poster 2003**

**Title:** Identification of positional functional candidate genes for protein percentage and lactation persistency

Presenting Author: Mini Singh, University of Sydney, NSW 2570

Other authors (name only):
1. Peter C. Thomson
2. Herman W. Raadsma

**Abstract:**

Positional and functional evidence for importance of genes for lactation persistency and protein percentage will be identified by integrating positional candidate genes from QTL fine mapping and genes identified against location of functional candidates obtained from transcript profiling. Previously identified QTL regions associated with these traits in sheep on OAR3 and OAR20 have been fine mapped using 28 additional markers (12 on a 10 cM region in OAR3 and 16 on a 20 cM region in OAR20) with Linkage Analysis/Linkage Disequilibrium analysis (LA/LD) over 4 sire backcross and double backcross families of approximately 700 daughters. From standard QTL analyses animals with contrasting genotypes 'Q' and 'q' for desired QTLs have been identified. RNA extracted from tissue samples from mammary glands
of these animals at three different times corresponding to pre-, peak-, and post lactation, have been used in functional (transcriptome) analysis using the Affymetrix Bovine Genechip array to obtain a list of differentially expressed genes. The use of combining QTL mapping and functional analyses using animals of defined QTL genotype will allow a list of positional functional candidate genes for further investigation.

Poster 2004

Title: New splice variants in the porcine PPARG1A gene

Presenting Author: Tim Erkens, Heidestraat 19, 9820 Merelbeke, Belgium

Other authors:
1. Karel Bilek
2. Alex Van Zeveren
3. Luc J. Peelman

Abstract:
Peroxisome proliferator-activated receptor γ coactivator 1α (PPARG1A) is a versatile coactivator with many functions, but is primarily involved in fat and energy metabolism. Therefore, it is a very interesting candidate gene for meat quality, which is one of the most important criteria in pig selection today. PPARG1A however, does not have the same function in every tissue and there is still much unknown about its regulation. Therefore a detailed transcription profile of PPARG1A in 34 tissues and 3 embryonic developmental stages in the pig was constructed in order to detect any alternative splicing. This was done by extracting mRNA from each tissue, cDNA synthesis and screening for splice variants with exon-spanning primers. Two new splice variants in the pig were discovered in which exon 8 was partly or completely spliced out. Both splice variants potentially give rise to a much shorter protein of respectively 359 and 337 AA, of which the first 291 AA would be the same compared to the complete protein (796 AA). Considering the functional domains of PPARG1A, it is very likely these variants have a large impact on the function of the protein and could provide an explanation for the regulation of the diverse functions of PPARG1A.

Poster 2005

Title: Acox1: A candidate gene for fatty acid composition in pigs

Presenting Author: María Muñoz - Departamento de Mejora Genética Animal -INIA. 28040 Madrid, Spain

Other authors (name only):
1. Estefanía Alves
2. Luis Varona
3. Jordi Estellé
4. Carmen Rodríguez
5. Luis Silió

Abstract:
Two QTL related with backfat fatty acid composition were previously detected on SSC12 in an Iberian x Landrace F2 cross. Acox1 was selected as candidate gene to explain one of these QTL, because human-pig comparative mapping shows the human localization could match in the QTL interval and Acox1 is the first enzyme of the fatty-acid β-oxidation in peroxisomes. Alternatively spliced transcript variants encode two different enzymes with the exons 3-II and 3-I respectively. The complete porcine cDNA sequences of both isoforms were sequenced in two Iberian, two Landrace and three pigs of a composite Chinese-European line, and seven SNP were detected. Two missense SNP were genotyped in the Iberian x Landrace intercross, and a new linkage map of the chromosome based on the segregation of 11 markers was obtained. The Acox1 gene mapped at 17.5 cM between the S0143 and GH markers. New linkage analysis identified three significant QTL: QTL1 affects the percentage of myristic, QTL2 the average chain length and the percentages of palmitic, linolenic and gadoleic and QTL3 the percentages of palmitoleic, stearic and vaccenic. No evidence was obtained of association of analyzed missense Acox1 SNP with fatty acid composition.

Poster 2006

Title: Fine Mapping of QTL for Parasite Resistance in Sheep

Presenting Author: NATASHA A ELLIS, Centre for Advanced Technologies in Animal Genetics and Reproduction (ReproGen), The University of Sydney, PMB3, Camden, NSW, 2570, Australia

Other authors (name only):
1. KAREN J FULLARD
2. SEYIT A KAYIS
3. DINESH KHATKAR
4. GINA ATTARD
5. KEN J BEH
6. DAVID PIEDRAFITA
7. HERMAN W RAADSMA

Abstract:
Gastro-intestinal nematodes (GIN) are estimated to cost Australian sheep producers over $AUD200 million annually through loss of production and death. Effective control of GIN is compromised by parasite
Resistance to drenches, so selecting sheep that are naturally resistant to parasitic infection is a feasible alternative to chemical control. We have fine mapped two QTL regions for parasite resistance in a backcross Indonesian Thin Tail x Merino population that were twice challenged with H. contortus. Twelve and fourteen new microsatellites were added to the 15 and 14 markers previously mapped on OAR1 and 3 respectively. QTL for FEC (P<0.01) and body weight change (P<0.05) at first challenge were detected on OAR1 around positions 169cM and 312cM using linkage analysis. QTL for FEC (P<0.05) and PCV changes (P<0.05) during the first challenge were also detected on OAR3, around positions 180cM and 160cM. Caution should be used when interpreting these positions as confidence intervals were large; LA/LD analysis will be used to address this. A meta-analysis is underway with the results to be used in a collaborative effort to further fine map with a targeted SNP panel. Additional positional candidate gene studies are being conducted by integrating mapping studies with gene expression analyses.

Poster 2007

Title: Mapping of Quantitative Trait Loci for lactation persistency traits in ewes

Presenting Author: Elisabeth Jonas; 425, Werombi Road; Camden 2570, NSW; Australia

Other authors (name only):
1. Peter C. Thomson
2. D. McGill
3. Mary K. Lam
4. Herman W. Raadsma

Abstract:
The persistency is a trait of great consideration and well investigated in dairy cattle; but only a few studies were performed targeting the productivity in dairy sheep. It was found in different studies that cows with greater persistency tend to incur less feed, health, and reproductive costs and that cows affected with mastitis tend to have less persistent lactations. In this study samples of animals of an Awassi-Merino backcross population were used for a linkage analysis using regression and maximum likelihood methods. The curves of the milk, protein, fat, lactose and energy yield were calculated using the Wood model. The three parameters derived from the model were further biological interpreted to calculate the yields and persistence of the traits. QTL for milk, protein, fat, lactose and energy yield, the persistency of the different traits and additional for the somatic cell score could be detected on different ovine chromosomes with the most interesting region on OAR3. The results of the genome-wide detection leading to the first study of persistency for milk composition traits in dairy sheep. Future fine mapping and candidate gene approaches may lead to the detection of marker for the improvement of the productivity and health in dairy sheep.

Poster 2008

Title: Deletion in a conserved CCR5 motif is associated with approximately doubling proviral load of maedi-visna/ovine progressive pneumonia virus

Presenting Author: Stephen N. White, USDA-ARS-ADRU, PO Box 646630, 3003 ADBF, WSU, Pullman, WA 99164

Other authors (name only):
1. Michelle R. Mousel
2. Gregory S. Lewis
3. Lynn M. Herrmann-Hoesing

Abstract:
CCR5 is a chemokine receptor that serves as a coreceptor for human immunodeficiency virus (HIV) and may have a role in regulating maedi-visna/ovine progressive pneumonia virus (OPPV) infection. Both of these lentiviruses are macrophage-tropic, have similar genomic structures, and cause lifelong persistent host infection. A human CCR5 coding deletion (termed delta-32) results in strong resistance to HIV infection by preventing expression of active CCR5 protein on the cell surface, and polymorphisms in CCR5 regulatory regions have been implicated in delayed progression to acquired immune deficiency syndrome (AIDS). To evaluate the impact of CCR5 on OPPV, proviral loads of OPPV were measured in 383 naturally exposed Rambouillet, Polypay, and Columbia sheep with approximately equal numbers from each breed. The ovine CCR5 genomic sequence was determined, and polymorphisms were obtained from the open reading frame and surrounding regulatory sites. One haplotype was associated with approximately doubling proviral load among positive animals (P<0.01), and the haplotype contains a 4 base deletion within a conserved motif in a regulatory region. While none of the polymorphisms was associated with differing odds of infection, the association with reduced proviral load suggests CCR5 may play a role in restricting the degree of OPPV infection.

Poster 2009

Title: Whole Genome Scan in Commercial Angus Cattle for QTL Influencing Carcass Value Traits: Carcass Weight, Fat Thickness, Marbling, and Ribeye Area.
Presenting Author: Matthew McClure.
Division of Animal Sciences, University of Missouri, Columbia, Missouri 65211, USA

Other authors (name only):
1. Jae W. Kim
2. Natalya S. Morsci
3. Stephanie D. McKay
4. Megan M. Rolf
5. Jared E. Decker
6. Robert D. Schnabel
7. Jeremy F. Taylor

Abstract:
We have genotyped two large Angus mapping populations for 415 microsatellites and 9 SNP markers for all *Bos taurus* autosomes and the X chromosome and have tested for quantitative trait loci (QTL) influencing marbling, ribeye area, carcass weight, and fat thickness. The first population is comprised of 1,678 AI sires born after 1955; the second is comprised of 36 halfsib families containing 2,546 commercial steers produced by the Circle A Ranch and MFA Inc. Data were analyzed using QTLExpress under a halfsib design and by LOKI using the full pedigree. At a chromosomewise P<0.01 significant level, every autosome was found to harbour multiple carcass related QTL. We estimate there to be as many as 22 QTL for carcass weight, 24 QTL for fat thickness, 21 QTL for marbling and 31 QTL for ribeye area within the Angus genome. Comparing our data to published results suggests that the majority of these QTL segregate within all *Bos taurus* breeds of cattle, but that novel QTL could exist within specific breeds. These results support our population-based approach to QTL mapping within commercially relevant populations.

Poster 2012
Title: Association of SNPs with carcass measurements and taste panel assessed meat quality traits
Presenting Author: …
Jennifer Gill, Roslin Institute, Midlothian, UK, EH25 9PS
Other authors (name only):
1. Stephen C Bishop
2. Caroline McCorquodale
3. Pamela Wiener

Abstract:
Various single nucleotide polymorphisms (SNPs) have been shown to be associated with meat and carcass quality traits in cattle. Examples include leptin, associated with fatness, and calpain and its inhibitor calpastatin, associated with meat tenderness. SNPs from these genes have been incorporated into commercial genetic tests for predicted meat quality. This study aimed to validate the effects of these six SNPs and to test two further SNPs, in the growth hormone receptor (GHR) and DGAT1 genes, also suggested to have effects on meat quality. Traits included a mechanical measure of tenderness (tenderometer), 22 carcass quality traits and a unique set of seven taste-panel-assessed sensory traits. Data were collected from 483 commercial Aberdeen Angus-cross animals. Statistical analyses used REML. Fixed effects included farm, sex, genotype and the genotype-sex interaction; random effects included the interaction of farm, sire and slaughter date. Significant associations between SNP alleles and traits were found for four SNPs (p<0.05 for all). An association was confirmed between one of the calpain SNPs and tenderness, assessed by both taste panel and tenderometer, in agreement with previous findings. Other significant associations included a leptin SNP with overall liking and fat level, DGAT1 with fat level and GHR with steak odour.

Poster 2013
Title: Genetics markers involved in productive traits in Romanian swine breeds
Presenting Author: Costache Marieta, Splaiul Independentei 91-95, Bucharest 5, 050095, Romania
Other authors (name only):
1. Maria Adina Manea
2. Stelian Kevorkian
3. Georgescu Sergiu Emil
4. Anca Dinischiotu

Abstract:
Research of the gene polymorphisms with major economical effects is currently the best-known application of pig genome mapping in breeding practice. In swine breeds, two major genes are well-known: the ryanodine receptor 1 gene (RYR1), associated with meat quality, and estrogen receptor gene (ESR) whose polymorphism is correlated with reproduction traits. Our objective was to develop straightforward methods to examine the two loci in swine breeds from Romania. We used the PCR-RFLP technique to identify two point mutations: one for RYR1 gene (C→T) correlated with malignant hyperthermia and another for ESR gene (T→G) associated with reproduction traits. DNA amplification was carried out through PCR using normal primers and amplicons were digested with restriction endonuclease *HhaI* and *Aval*. Restricted products were analyzed via electrophoresis in agarose gel stained with ethidium bromide. Our results were confirmed through sequencing.
The major focus of this study was to identify the normal or affected homozygous and carrier individuals in Romanian swine breeds and to implement a useful diagnosis methodology in order to assist veterinarians and breeders in disease control. The method presented above is reliable, fast, and can be successfully applied in the wide-scale screening of different pig populations.

Poster 2014

Title: Genetic characterisation of Charolais and Limousin cattle breeds using microsatellites markers in Slovakia.

Presenting Author: Jakabová Daniela, The Breeding Services of the Slovak Republic, state enterprise, Hlohovská 5, 95141 Nitra- Lužianky, Slovakia

Other authors (name only):
1. Trandžík Jozef
2. Kozlík Peter
3. Fečková Monika
4. Mindeková Silvia
5. Massányi Peter
6. Ryba Štefan
7. Židek Radoslav
8. Jakab František

Abstract:
The polymorphism of 11 microsatellite markers (BM1824, BM2113, ETH3, ETH10, ETH 225, INRA23, SPS115, TGLA53, TGLA122, TGLA126, TGLA222) was investigated in the Slovak population of 199 Charolais and 66 of Limousin cattle. DNA was amplified in one multiplex PCR for amplifying these microsatellites (StockMark for Cattle, AB). The automatic analysis was performed using an ABI 310 sequencer and size of analyzed DNA fragments was determined in base pairs using computer package GeneScan v.3.7 (AB). Statistical parameters were calculated using Powermarker v.3.25 software. Total 99 alleles were identified in the Charolais breed and 82 in the Limousin breed. The number of allele per each locus ranged from 4 (ETH10) to 13 (TGLA53, TGLA122) in Charolais breed and in Limousin from 4 (BM1824) to 12 (TGLA227). The average value of $H_e$ was 0.714 and $H_o$ was 0.752 in Limousin. In Charolais, heterozygosities were lower: $H_e$ was 0.667 and $H_o$ was 0.644. Mean PIC values were similar: 0.6371 in Charolais and 0.6737 in Limousin cattle breed. The combined exclusion probability was (PE>0,999) in both breeds. The values of PE confirmed the usefulness of this set of microsatellite markers in parentage testing of Charolais and Limousin cattle in Slovakia.

Poster 2015

Title: Increasing accuracy of estimated breeding value using DNA parentage test information in beef natural service populations

Presenting Author: Z. Wang, Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, AB, Canada, T6G 2P5.

Other authors (name only):
1. D. Kolbehdari
2. G. Plastow
3. S.S. Moore
4. J. S. Smith

Abstract:
Genetic markers have been used for assigning progeny to their sires in beef cattle populations where natural service is the norm. The genetic evaluation for such population can only use maternal information since the paternal information is unknown. Sufficient single nucleotide polymorphisms (SNP) are now available for efficient and accurate assignment of parentage. The value of such parentage assignment is discussed in reviews, but no quantified research can be found. The IGENITY cattle marker panel contains 200 SNP was used to assign sires in nine herds of seed stock beef cattle population. On average the success rate of the parentage test in this population was between 82-87%. A total of 2082 progenies out of 2558 were assigned to 110 bulls across the herds. The objective of this study was to quantify the increasing in accuracy of estimated breeding value by adding the parentage test information. Two performance traits, birth weight (BWT) and weaning weight (WWT) were analyzed in this study. By adding the paternal information, the accuracy of EBV for these two traits was increased by approx. 10% and 17% on average for young calves and sires respectively.

Poster 2016

Title: Mapping quantitative trait loci for embryo survival and litter size in pigs

Presenting Author: Silvia Hernandez, The Roslin Institute and Royal (Dick) School of Veterinary Studies, University of Edinburgh, Roslin, Midlothian EH25 9PS, Scotland, UK.

Other authors (name only):
1. Heather A. Finlayson
2. Cheryl Ashworth
3. Chris S. Haley
4. Alan L. Archibald
Abstract:
Reproductive performance is a critical component of sustainable animal production systems. Increasing the number of viable and productive offspring per reproductive female reduces financial and environmental costs and improves the sustainability of the system. The Chinese Meishan breed is one of the most prolific pig breeds known, displaying greater litter size than commercial Western breeds. Meishan pigs deliver this increased litter size through higher levels of prenatal survival for a given ovulation rate. To date only a fraction of the molecular genetic variation that underpins the superior reproductive performance of the Meishan has been identified. In an earlier study we mapped a QTL for the related traits of litter size and embryo survival to the distal end of pig chromosome 8. Additional genetic markers were developed in the QTL region and genotyped across the three-generation Large White – Meishan F2 population in order to improve the resolution with which the litter size and embryo survival QTL were mapped. The confidence interval for the QTL has been aligned with the emerging pig genome sequence and homologous regions of the human and murine genomes in order to reveal positional candidate genes (including secreted phosphoprotein 1, SPP1) for litter size and embryo survival.

Poster 2017

Title: A putative quantitative trait locus on chromosome 20 associated with bovine pathogenic disease incidence

Presenting Author: Eduardo Casas, USDA, ARS, U.S. Meat Animal Research Center, P.O. Box 166, Clay Center, NE 68933. USA

Other authors (name only):
1. Gary D. Snowder

Abstract:
The objective was to detect QTL associated with the incidence of multiple pathogenic diseases. Four F1 sires were used to produce offspring: Brahman × Hereford (BH; n = 547), Piedmontese × Angus (PA; n = 209), Brahman × Angus (BA; n = 176), and Belgian Blue × MARC III (BM; n = 246). Records for bovine respiratory disease, infectious keratoconjunctivitis (pinkeye), and infectious pododermatitis (footrot) were available for all the offspring. The incidences of these three microbial pathogenic diseases were combined into a single binary trait to represent an overall pathogenic disease incidence. Offspring diagnosed and treated for one or more of the pathogenic diseases were coded as a 1 for affected and 0 for untreated. A putative QTL was detected in the family derived from the BH sire. The maximum F-statistic (F = 13.52; P = 0.0003) was located at centimorgan 18. The support interval of the QTL spanned from centimorgan 9 to centimorgan 28. Offspring inheriting the Hereford allele, in the BH-sired family, and the Angus allele, in the PA-sired family, were less susceptible to incidence of pathogenic diseases.

Poster 2018

Title: STRs vs. SNPs in horse parentage testing

Presenting Author: Teruaki Tozaki*, Department of Molecular Genetics, Laboratory of Racing Chemistry, Utsunomiya, Tochigi 320-0851, Japan, ttozaki@lrc.or.jp

Other authors (name only):
1. Hironaga Kakoi*
2. Hitoshi Gawahawa
3. Kei-ichi Hirota
4. Nobushige Ishida
5. Telhisa Hasegawa
* Authors contributed equally to this work

Abstract:
At present, about 1.4 million SNPs were identified on the horse genome due to recent advances of the horse genome project. Interest in utilizing SNPs for parentage testing is rapidly increasing, but there is no enough information to evaluate the utility, such as their power of discrimination. In this study, two genotyping systems using each STRs and SNPs were used for the evaluation. At first, two multiplex assay groups were constructed to genotype 53 SNPs and genotypes of 95 thoroughbreds were determined using Sequenom’s MassARRAY system. Total PE of the 53 SNPs was 0.99996, and MAF was 0.36 on average. The SNP system was applied for the questions of the ISAG 2006 horse comparison test, and 5 of the 53 SNPs excluded for a pseudo Sire-Foal combination. Next, 30 STRs used in our routine and additional parentage testing were genotyped using the 95 thoroughbreds. Total PEs were 0.99997 and 0.99991, respectively. Allele frequency was 0.721 on average. The STR systems were also applied for the same questions, and 5 and 6 STRs excluded for the pseudo combination, respectively. Those results indicate that about 50 SNPs are required to reach equivalent power of discrimination as well as the routine STR systems.
**Poster 2019**

**Title: Nutrient-induced modulation of genome profile and genetic network**

Presenting Author: Congjun Li, Bovine Functional Genomics Laboratory, USDA-ARS, 10300 Baltimore Ave, BARC EAST, Beltsville, MD 20705 USA

Other authors (name only):
1. Robert W. Li
2. Ted H. Elsasser

**Abstract:**
Short-chain fatty acids participate in metabolism both as nutrients and as regulators of histone deacetylation. The major biochemical change that occurs in cells treated with butyrate is the global hyperacetylation of histones. Utilizing gene expression profiling, our studies indicated that butyrate induces many significant changes in the expression of genes associated with regulatory pathways that are critical to cell growth, immune response and signal transduction. We identified 450 genes significantly regulated by sodium butyrate at a very stringent false discovery rate (FDR) = 0%. When relaxing the stringency to FDR = 10%, there were 3662 genes significantly regulated (3662/45383 = 8%). The functional category and pathway analyses of the microarray data revealed that four canonical pathways (Cell cycles checkpoint; pyrimidine metabolism; G1/S checkpoint regulation and purine metabolism) were significantly perturbed. The biologically relevant networks and pathways of these genes were also identified. IGF2, TGFB1, TP53, E2F4, and CDC2 were established as being centred in these genomic networks. Butyrate induced biological effects in bovine cells provide an example of epigenetic regulation of genome and a basis for understanding the full range of the biological roles and the molecular mechanisms that butyrate may play in animal cell growth, proliferation, and energy metabolisms.

**Poster 2020**

**Title: Relationship of the Bovine Growth Hormone Gene to Carcass Traits in Japanese Black Cattle**

Presenting Author: Ken Tatsuda (K. Tatsuda), 1533 Befucyo Kasai-City Hyogo 679-0198, Japan, Tel. : +81-790-47-2430, Fax : +81- 790-47-0549, E-mail : Ken_Tatsuda@pref.hyogo.lg.jp

Other authors (name only):
1. A. Oka
2. E. Iwamoto
3. Y. Kuroda
4. H. Takeshita
5. H. Kataoka

**Abstract:**
The bovine growth hormone gene (bGH) possesses three haplotypes, A, B and C, that differ by amino acid mutations at positions 127 and 172 in the fifth exon: (leucine127, threonine172), (valine127, threonine172) and (valine127, methionine172), respectively. The correlation between meat quality or carcass weight and these haplotypes was investigated in Japanese black cattle. Altogether, 940 bGH haplotypes were compared with respect to six carcass traits; carcass weight, longissimus muscle area, rib thickness, subcutaneous fat thickness, beef marbling score and beef color. The frequency of the B haplotype was higher (0.421) than that of A (0.269) and C (0.311). High carcass weight and low beef marbling were associated with haplotype A (p<0.05 and p<0.01, respectively), whereas beef marbling was increased by haplotype C (p<0.05). Estimated regression coefficient of the A haplotype substitution effect for carcass weight and beef marbling score were 5.55 (13.1% of the phenotypic S.D.) and -0.31 (17.0%), respectively. That of the C haplotype for beef marbling score was 0.20 (11.0%). The other traits showed no relationship to the haplotypes examined. The results of this investigation suggest that information pertaining to bGH polymorphisms in Japanese black cattle could be used to improve the selection of meat traits.

**Poster 2021**

**Title: Genome wide association study of feed efficiency in beef cattle.**

Presenting Author: Laura Sherman, Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, AB, T6G 2P5, Canada

Other authors (name only):
1. Donald Nkrumah
2. Stephen Moore

**Abstract:**
Feed is the highest variable cost in beef production making feed efficiency an important trait to study. Feed efficiency is measured as residual feed intake (RFI), which is the difference between actual feed intake of the animal and expected feed intake based on weight and growth rate. 2,633 SNPs covering the bovine genome were analyzed in 464 steers for associations with RFI. 150 SNPs were identified with $P < 0.05$. To create a panel of SNPs that were maximally informative for RFI from these 150 SNPs, two methods were tested. First, a sequential molecular breeding value (MBV) was created for each animal by sequential addition of the SNP effects and only SNPs which improved the model were included. This panel
contained 79 SNPs. Second, the SNPs were combined in a single multivariate model and a backward elimination was used until all SNPs left in the model were significant with $P < 0.05$, which left 32 SNPs. Regression of an MBV of the 32 SNPs on RFI produced $r^2 = 0.497$ and regression of the 79 SNP MBV produced $r^2 = 0.497$. These SNP panels provide progress towards identification of markers for use in marker-assisted selection of RFI in beef cattle.

**Poster 2022**

**Title**: Genetic effects mapped in a Charolais X Holstein cross cattle population.

BEATRIZ GUTIÉRREZ-GIL¹, JOHN L WILLIAMS², PAMELA WIENER³

¹Dpto. Produccion Animal. Facultad de Veterinaria. Universidad de León. 24071 León, Spain.
²Parco Tecnologico Padano, Via Enstein, Lodi 296900, Italy.
³Division of Genetics and Genomics, The Roslin Institute and R(D)SVS, University of Edinburgh, Roslin, Midlothian, EH259PS United Kingdom.

**Abstract**: An experimental cattle population to identify QTL influencing cattle production traits was established at the Roslin Institute, starting in 1995 by crossing Charolais sires and Holstein dams. Phenotypic measurements for a wide range of traits were made on the 500 second-generation individuals of the herd (F2, Charolais Backcross and Holstein Backcross individuals). We present here a summary of the results obtained for the traits related to meat production, temperament and coat colour dilution. A QTL affecting proportion of bone in the carcass on chromosome 6 was the most significant QTL identified for production traits. Many other genome-wide significant QTL were detected for yield and carcass composition traits. Several QTL were also found with effects on technological properties of the meat (e.g. moisture, pH) and consumer satisfaction (taste panel traits and fatty acid composition). For the temperament traits studied, which assessed the fearfulness of the animals to approaching humans and to social separation, a single genome-wide significant QTL was found on chromosome 29. The coat colour records were used to map the Charolais dilution locus, and to assess the relationship of a mutation in the SILVER gene with the coat colour dilution characteristic of the Charolais breed.

**Poster 2023**

**Title**: Searching QTL affecting milk traits on chromosome 22 in Spanish Churra sheep.

Presenting Author: ELSA GARCÍA-GÁMEZ,
Departamento Producción Animal – Facultad de Veterinaria, Campus de Vegazana s/n, 24071 – León, Spain

Other authors (name only):
1. BEATRIZ GUTIÉRREZ-GIL
2. MARTA GARCÍA
3. LUIS FERNANDO DE LA FUENTE
4. JUAN JOSÉ ARRANZ

**Abstract**: After a preliminary genome scan carried out in Churra sheep to detect QTL influencing milk production traits, additional families are now being analyzed to confirm some of the identified effects. We present here the analysis performed on chromosome 22 for 15 additional families, which included 798 Churra ewes. Phenotypic measurements for milk yield, protein percentage, fat percentage and somatic cell scores (SCS) were used to calculate yield deviations used in the analysis. Eight microsatellite markers evenly distributed across the chromosome were genotyped for all the population and used to construct a linkage map. A multimarker regression method was implemented through the QTL Express software. Chromosome-wise critical values were calculated through 10,000 phenotype permutations. The average information content across the linkage map was 0.78. The across-family association analysis revealed a 5% chromosome-wise significant QTL for the SCS (p-value = 0.026), between markers INRA81 and TGLA429. None of the production related traits showed significant linkage association on this chromosome. The SCS QTL was found to be segregating in three of the 15 analyzed families. The joint analysis of these data with those generated through the genome scan experiment will be needed to confirm the QTL effect here reported.
In farms, the red deer mating programs consists of one male which services 30-40 hinds; working usually with 12 groups of enclosed hinds. In our country, red deer maternity assessment is still based on visual observation during the post-parturient phase. The behaviour of hinds and calves, mainly mother care and nurse, is used to assess maternity relationships. Adoption and allosuckling can affect, however, the accuracy of this procedure, which also is very hard and expensive. Like in other species, DNA typing by microsatellites could be the alternative to reduce the mismothering rate. Due to the absence of a set of markers for typing service in red deer, our lab developed a panel of 11 markers useful for identification and parentage testing. All the markers are of public domain, derived from cattle and deer genomes. A total of 130 alleles were found on 90 samples tested. Assuming one known parent, the individual exclusion power ranged between 0.17 and 0.73, while the cumulative exclusion probability reached 0.999947 for the whole panel. The panel was validated on a trial assay, involving 34 hinds, 32 calves and 7 males. The results of DNA parentage testing were confirmed by previous visual observation assignment.

Poster 2025

Title Polymorphisms in Positional Candidate Genes on BTA14 and BTA26 affect Carcass Quality in Beef Cattle


Other authors (name only):
1. Donald Nkrumah
2. Laura Sherman
3. Stephen Moore

Abstract:
Previous analysis on Bovine Chromosome 14 (BTA14) and 26 (BTA26) have indicated a number of quantitative trait loci (QTL) affecting meat quality traits. Association studies of 2, 4 dienoyl CoA reductase 1 (DECR1) in pigs and core binding factor alpha domain 2 (CBFA2T1) in humans have shown associations between polymorphisms in those genes with lipid metabolism. Sequencing analysis in cattle identified nine polymorphisms in DECR1; including four exonic, two of which produced changes, while four intronic SNPs were identified in CBFA2T1. Multiple sequence alignment of DECR1 among cattle, human and mice showed that four mutations lie in conserved regions across these species. Single locus analysis produced associations (P < 0.05) with ultrasound marbling score (CBFA2T1) and ultrasound backfat (DECR1). Recent studies have linked fibroblast growth factor 8 (FGF8) to several QTL affecting obesity in mice which indicated its potential for regulating adiposity in other species. In FGF8, four polymorphisms were identified, two intronic and two exonic. Single locus analysis resulted in significant associations (P < 0.01) with carcass backfat and lean meat yield. This study hopes to contribute to a growing list of genes with specific functions affecting the final meat quality in beef cattle.

Poster 2026

Title: Effect of SCD, SREBP and FASN genotypes on fatty acid composition in adipose tissue of Japanese Black cattle.

Presenting Author: Shinji SASAZAKI. Laboratory of Animal Breeding and Genetics, Graduate School of Agricultural Science, Kobe University, Kobe, Japan 657-8501.

Other authors (name only):
1. Atsuko TANAKA
2. Hideki OHSAKI
3. Kenji OYAMA
4. Fumio MUKAI
5. Hideyuki MANNEN

Abstract:
Fatty acid composition of beef adipose tissue is one of important traits because high proportion of monounsaturated fatty acid is related with favorable beef flavor and tenderness. In this study, we investigated the effects of genetic factors such as stearoyl-CoA desaturase (SCD), sterol regulatory element binding protein (SREBP) and fatty acid synthase (FASN) on beef carcass traits including fatty acid composition using two cattle populations (N = 417 and 233, respectively). Sire effect was significantly related to almost all traits except BMS, suggesting that the trait examined in this study is highly controlled by genetic factors. The effect of SCD genotypes on fatty acid composition was detected remarkably in both cattle groups, especially on stearic (C18:0), oleic (C18:1) acids and monounsaturated fatty acid (MUFA) content. FASN genotypes were significantly associated with almost all fatty acids (C14-18) except stearic (C18:0) and linoleic acid (C18:2). However, effect of SREBP genotypes was not identified in this study. Our results also suggested there were other genetic factors relevant to fatty acid metabolism. In conclusion, these results would contribute to bring an insight into molecular mechanism of fatty acid metabolism in cattle.
Poster 2027

Title: The ubiquitin ligase gene (WWP1) is responsible for the chicken muscular dystrophy.

Presenting Author: Hirokazu MATSUMOTO, Laboratory of Animal Breeding and Genetics, Graduate School of Agricultural Science, Kobe University, Kobe, Japan 657-8501.

Other authors (name only):
1. Hideaki MARUSE
2. Yumi INABA
3. Kanako YOSHIZAWA
4. Shinji SASAZAKI
5. Akira FUJIWARA
6. Masahide NISHIBORI
7. Akinori NAKAMURA
8. Shin’ichi TAKEDA
9. Nobutsune ICHIHARA
10. Tateki KIKUCHI
11. Fumio MUKAI
12. Hideyuki MANNEN

Abstract:
Muscular dystrophy is defined as a group of inherited diseases producing progressive weakness and degeneration of skeletal muscles. It is well known that abnormalities of muscle proteins to compose the linkage between sarcolemma and basal lamina lead to muscular dystrophies, but genes responsible for several muscular dystrophies and related diseases have not yet been identified. The gene responsible for chicken muscular dystrophy with abnormal muscle (AM) remains unclear, either. Our previous study revealed seven functional genes as AM candidate genes, none of which were determined to be responsible for other muscular dystrophies. In current study, sequence and expression comparison between normal and dystrophic chickens was conducted to detect a mutation responsible for the disease. We detected three synonymous mutations and one missense mutation in AM candidate genes, while no remarkable alteration of expression patterns was observed. The missense mutation was observed in WWP1 and detected only in dystrophic chickens in several tetrapod species. These results strongly suggested WWP1 would be responsible for chicken muscular dystrophy. The information must be useful for determining the corresponding human dystrophy and for providing new insights for understanding muscular dystrophies.

Poster 2028

Title: Development of SNPs markers for individual identification and parentage test in Japanese Black cattle population.

Presenting Author: Kazuhiro HARA, Laboratory of Animal Breeding and Genetics, Graduate School of Agricultural Science, Kobe University, Kobe, Japan 657-8501.

Other authors (name only):
1. Hideki WATABE
2. Shinji SASAZAKI
3. Fumio MUKAI
4. Hideyuki MANNEN

Abstract:
Individual identification and parentage control are essential for consumer protection and efficient management of animal population. Since single nucleotide polymorphisms (SNPs) are abundant in genome, genetically stable and amenable to high-throughput automated analysis, these are expected to be efficient DNA markers for the diagnostic. This study describes the development of SNP markers in order to apply the identification and parentage test in Japanese Black cattle population. Amplified fragment length polymorphism method was employed to detect informative candidate markers and yielded 44 SNPs markers from the 220 primer combinations. Unlinked 30 SNPs out of them were finally selected as diagnostic markers. The allele frequencies for each marker were estimated by using PCR-RFLP in Japanese Black population. Based on the frequency data, estimated identity power of these markers was 1.18E-12. Parentage exclusion probability was 0.97243 with only one sampled parent. Sire exclusion was 0.99747 when the dam’s genotype was known. This panel of SNP markers is theoretically sufficient for individual identification of any individual in Japanese Black cattle. These markers could be useful for identification and parentage test, and would contribute to management of beef industry in Japan.

Poster 2029

Title: Polymorphisms of fatty acid related genes and association between genotypes and fatty acid composition in Japanese Black cattle.

Presenting Author: Shogo HOASHI, Laboratory of Animal Breeding and Genetics, Graduate School of Agricultural Science, Kobe University, Kobe, Japan 657-8501.

Other authors (name only):
1. Atsuko TANAKA
2. Hideki OHSAKI
3. Shinji SASAZAKI
4. Kenji OYAMA
5. Fumio MUKAI
6. Hideyuki MANNEN
Abstract:
The fatty acid composition of adipose tissue in cattle has become more important in the beef industry. Fat tissue containing abundant monounsaturated fatty acid (MUFA) reflects lower fat melting points, leading to favorable beef flavor and decreasing the blood concentration of LDL-cholesterol. In this study, we searched polymorphisms in full length CDS of the six genes, FABP4, LXRα, CYB5, ACSL1, ACSL4 and DGAT2, associated with fatty acid and lipid metabolism. Sequence comparison among eight animals, including five Japanese Black and three Holstein cattle, revealed 12 single nucleotide polymorphisms. Four of them, I74V, G51E, V133I in FABP4 and G51E, V133I in LXRα, were predicted to cause amino acid substitutions. We investigated associations between these genotypes and fatty acid compositions of carcass fat in Japanese Black cattle. The genotype I74V in FABP4 was significantly associated with palmitoleic acids (C16:1, P<0.01) content of intramuscular fat, while remarkable effect was not detected in the other genotypes. In conclusion, I74V in FABP4 gene would contribute to improve fatty acid composition in beef cattle.

Poster 2030

Title: Assessment of genetic diversity in Percheron horses registered in Canada using genealogical records and multiloci microsatellite data.

Presenting Author: Tara McParland, Maxxam Analytics Inc., 2 - 335 Laird Road, Guelph, ON, Canada, N1H 6J3

Other authors (name only):
1. Yves Plante

Abstract:
The Percheron was also a very popular stage-coach horse. In Canada, between 1908 and 1950, the Percheron registry counted twice as many horses registered. Over the last ten years, the Percheron has regained some popularity with average yearly registrations of 318 stallions and 655 mares. Individual microsatellite allelic richness and the polymorphism information content stayed constant through time. In conclusion, the Canadian registered Percheron horse population exhibits a relatively high level of genetic diversity even though it has gone through a reduction in population size between 1950 and 1975.

Poster 2031

Title: Understanding Haemophilus Parasuis Infection in Porcine Spleen Through Transcriptomics Approach

Chang-Chun Li, * Hong-Bo Chen, * Ming-Di Fang, * Meng-Jin Zhu, Xin-Yun Li, Rui Zhou, † Kui Li, ‡ Shu-Hong Zhao *

*Key Lab of Agricultural Animal Genetics, Breeding, and Reproduction of Ministry of Education, †Division of Animal Infectious Disease, State Key Laboratory of Agricultural Microbiology, Huazhong Agricultural University, Wuhan 430070, P R China; ‡Department of Gene and Cell Engineering, Institute of Animal Science, Chinese Academy of Agricultural Sciences, Beijing 100094, P R China

Abstract:
Haemophilus parasuis (HPS) is a prominent swine pathogen that causes Glässer’s disease characterized by fibrinous polyserositis, meningitis and arthritis. The molecular mechanisms underlying disease pathogenesis related to HPS infection is poorly understood, particularly the host counteraction to HPS invasion by the immune system. In this study, we firstly investigated the global expression changes in spleen following HPS infection using the Affymetrix Porcine Genechip™. Differential expression of selected genes was confirmed by QRT-PCR analysis. Differentially expressed genes involved in subsystems/modules including inflammasomes, acute-phase proteins and complement, cell differentiation, adhesion molecules, transcription factors and many others which may play central roles in several cascade. Also, down-regulations of both MHC class I and MHC class II genes are likely the main mechanisms for HPS evasion. We further detected the expression change of differentially expressed genes in LPS or Poly (I:C) treated porcine PK-15 cells. Interestingly, mRNA levels of S100A8, S100A9, and S100A12 increased in a sustained manner within 48h after administered LPS and Poly (I:C) respectively, suggesting they may play important roles in porcine systemic inflammations. In silico mapping of differentially expressed genes to porcine immune trait QTL regions showed that acute-phase protein HP and transcription factor CEBPD may be important candidate genes which merit further investigation.
Altogether, our findings indicate previously unrecognized gene transcription changes in case of HPS infection in vivo and should provide new clues for identification of candidate genes related to HPS resistance.

Poster 2032

Title: FecX\(^R\), a deletion in the BMP15 locus associated to increased prolificacy in the Rasa Aragonesa sheep breed.

Presenting Author: Luis V. Montecagudo Ibañez, Department of Anatomy, Embriology and Genetics, Faculty of Veterinary Sciences of Zaragoza (Spain), C/Miguel Servet 177, 50013-ZARAGOZA (Spain)

Other authors (name only):
1. Isidro Sierra
2. M. Teresa Tejedor
3. Adolfo Laviña
4. Ricardo Ponz

Abstract:
Since 1990, the National Association of Rasa Aragonesa Sheep Breeders (ANGRA) maintains a genetic improvement program for prolificacy. Data on 1.5 million were obtained during the field production control. We selected a small group of ewes showing high prolificacy performances in the resulting database. The exons for GDF9 (Growth Differentiation Factor 9) and BMP15 (Bone Morphogenetic Protein 15) were PCR amplified and sequenced. We identified a 17 bp deletion in the second exon of the BMP15 pro-protein. This deletion modifies the open reading frame in the sequence, so that only the first 45 amino acids of the wild variant are conserved. Then, premature stop codons appear at the positions 100 and 101. As a consequence the region of the pro-protein conserved in the mature protein is not synthesized (it normally starts in the position 120). Ewes heterozygous for the mutation (named FecX\(^R\)) show increased prolificacy, due to an increase in the number of duplets and triplets. In fact, the highest prolificacy in the ANGRA records was obtained by one of these ewes (3.67 lambs as a mean in three births). At present we are extending our search to flocks all over Aragón to detect further carriers of the deletion.

Poster 2033

Title: Biosafety assessment of markers genes in transgenic cloned cattle

Presenting Author: Yan Liu, State Key Laboratory for Agrobiotechnology, China Agricultural University, Yuanmingyuan West Road 2, Beijing, China

Other authors (name only):
1. Ning Li

Abstract:
Both enhanced green fluorescence protein (EGFP) and neomycin phosphotransferase type II (NPTII) are widely used in transgenic studies, but their side effects have not been extensively investigated. In this study, we evaluated the expression profiles of the two marker genes and the relationship between their expression and organ abnormality (dead cattle)/health profile (living cattle). In dead cattle, EGFP and NPTII protein expression level ranged from 0.3 to 5ug/g in heart, liver and lungs, and the expression profiles exhibited differential or mosaic pattern between the organs, the pathologic symptoms of which were identified, but were similar to those of age-matched cloned cattle (p>0.05). All data indicated that the expression of EGFP and NPTII is not associated with organ abnormality in transgenic cloned cattle. With respect to living cattle, all parameters derive from serum samples in which EGFP and NPTII expression will be measured by RIA and ELISA. Meanwhile, hormones and clinical parameters, such as complete blood counts and biochemical analyses will be checked every two months during half a year. All the parameters from transgenic cloned cattle will be compared with the ones from age-matched cloned cattle and age-matched normal cattle. The study is in progress.

Poster 2034

Title: Identification of differentially expressed genes in rumen tissues of dairy cattle fed a low or high grain diet

M. Taniguchi\(^1\), G.B. Penner\(^1\), K.A. Beauchemin\(^3\), M. Oba\(^1\), L.L. Guan\(^1\)

\(^1\)Department of Agricultural Food and Nutritional Sciences, University of Alberta, Edmonton, Alberta, T6G 2P5 Canada.
\(^2\)Animal Genome Research Unit, Division of Animal Sciences, National Institute of Agrobiological Sciences, 2 Ikenodai, Tsukuba, Ibaraki 305-8602, Japan.
\(^3\)Agriculture and Agri-Food Canada, Lethbridge, Alberta, Canada

Abstract:
A fundamental understanding of molecular mechanisms of ruminal epithelial cell adaptation to highly fermentable diets is essential to prevent ruminal acidosis (RA). In this study, we aimed to gain a broad
overview of the involved molecular mechanisms by
detecting differentially expressed genes (DEG) in
rumen tissues from lactating dairy cows fed a low
(8%) or a high grain diet (64%) using microarray
analysis (a bovine 24K panel). A total of 7035
elements were detected as DEG with >1.5-fold
expression change ($P < 0.05$): 2993 and 4042 were up-
and down-regulated in the cows fed a high grain diet,
respectively. Molecular function of the DEG analyzed
with gene ontology and KEGG databases indicated
that 49 (22 up/27 down), 26 (8 up/18 down) and 19
DEGs (9 up/10 down) are involved in MAPK,
calcium, and insulin pathways, respectively. The
down-regulated gene encoding cAMP-dependent
protein kinase subunit beta ($PRKACB$) was common
among the three pathways. This suggests that
intracellular signals induced by cAMP may be
negatively affected in cows fed highly fermentable
diets. Functional analyses of DEG to elucidate the
interactions between diet fermentability and metabolic
adaptation by animals, leads to a better understanding
of the role of absorptive metabolisms in mitigating
RA.

**Poster 2035**

**Title:** Molecular analysis and SNP development of three candidate genes ($IL6$, $IL10$, $TF$) for disease resistance in pigs

Presenting Author: Emilia Danilowicz, Institute of Genetics, University of Berne, Bremgartenstrasse 109a, 3001 Berne, Switzerland

Other authors (name only):
1. Mansoureh Akouchekian
2. Rosa Martinez-Arias
3. Inga Probst
4. Gerald-F. Gerlach
5. Tosso Leeb
6. FUGATO-consortium IRAS

**Abstract:**
Different cytokines such as interleukin 6 ($IL6$) and $IL10$ are secreted in response to specific microbial molecules referred to as pathogen associated molecular patterns (PAMPs) and play a central role in the immunological response. Transferrin-mediated provision of iron is essential for bacterial growth and iron-depletion of transferrin is a first line defense against bacterial infections.

We obtained the genomic structure and complete DNA sequence of the porcine $IL6$, $IL10$ and $TF$ genes and identified polymorphisms on a panel of ten different pig breeds. Comparative intra- and interbreed sequence analysis revealed 8 polymorphisms in the porcine $IL6$, 21 in the $IL10$, and 57 in the $TF$ gene, which include single nucleotide polymorphisms (SNPs) and insertion deletion polymorphisms (indels).

In the $IL6$ gene one polymorphism was located in an exon (c.92G>T) and predicted to cause an amino acid exchange (p.R31L). In the $IL10$ gene, two polymorphisms were located in the coding sequence, and one (c.371A>T) was predicted to cause an amino acid exchange (p.Q124L). In the $TF$ gene 7 of the 57 polymorphisms were located in the coding sequence. Two SNPs (c.1417A>G, c.1810A>C) were predicted to cause amino acid changes (p.K473E, p.N604H). All polymorphisms were submitted to the NCBI dbSNP.

**Poster 2036**

**Title:** Assignment of the swine MHC alleles and biological traits in selective breeding Duroc pigs

Presenting Author: ASAKO ANDO, Department of Molecular Life Science, Division of Basic Medical Science & Molecular Medicine, Tokai University School of Medicine, 143 Shimokasuya, Isehara, Kanagawa 259-1193, Japan

Other authors (name only)
1. OK KAR SOE
2. YASUNORI OHBA
3. NORIAKI IMAEDA
4. NAOHITO NISHII
5. MASAKI TAKASU
6. GOU YOSHIOKA
7. HISAKO KAWATA
8. ATSUKO SHIGENARI
9. HIROHIDE UENISHI
10. HIDETOSHI INOKO
11. HITOSHI KITAGAWA

**Abstract:**
Pigs with defined swine major histocompatibility complex (MHC: SLA) haplotypes and characterized immunological traits are useful for studies of transplantation and immune responses. To develop SLA homozygous pigs with novel SLA haplotypes, a pair of pigs was chosen from a commercial Duroc pig line, and substantial breeding within progenies was carried out for eight generations. The theoretical inbreeding coefficient at the eighth generation was 78.5%. In the selective breeding Duroc (SBD) pigs, only two SLA haplotypes including novel two class I haplotypes, Hp-27.30 and Hp-60.13 were assigned by SLA-DNA typing methods. The haplotypes were also identified by genetic polymorphisms of 36 microsatellite markers within the SLA region in the SBD pigs. Despite inbreeding for eight generations, litter sizes of the SBD pigs in the all generations were comparable to those of non-SBD pigs. Weaning weights from the fifth to eighth generation produced progenies significantly lighter ($p<0.01$) than those in the non-SBD pigs. Concerning the lymphocyte subsets, the percentage of CD4+ T-lymphocytes in
peripheral blood of the SBD pigs was significantly higher than that in the non-SBD pigs. Our newly established SBD pig lines with novel SLA haplotypes will be useful in further studies on immune responses against various foreign antigens.

Poster 2037

Title: Analyses of serum antibody titers against swine erysipelas vaccines and hepatitis E virus infection in SLA-defined selective breeding Duroc pigs

Presenting Author: HITOSHI KITAGAWA, Laboratory of Veterinary Internal Medicine, Faculty of Applied Biological Sciences, Gifu University, 1-1 Yanagido, Gifu 501-1193, Japan

Other authors (name only):
1. ASAKO ANDO
2. NORIAKI IMAEDA
3. OK KAR SOE
4. YASUNORI OHBA
5. GOU YOSHIOKA
6. TOSHIHIRO USHIJIMA
7. HIROAKI OKAMOTO
8. HOZUMI TANAKA
9. EIJI KOBAYASHI

Abstract:
To clarify the relationship between swine leukocyte antigen (SLA) haplotypes assigned by DNA typing techniques and antibody productivity, we determined antibody titers against swine erysipelas live and inactivated vaccines and natural infection of hepatitis E virus (HEV) in SLA-defined selective breeding Duroc pigs with 2 SLA haplotypes; Hp-27.30 and Hp-60.13. Serum antibody titers against a swine erysipelas live vaccine determined by a viable cell agglutination test were lower in order in Hp-27.30 homozygotes, Hp-27.30/60.13 heterozygotes and Hp-60.13 homozygotes. Mean titer in the homozygous pigs with Hp-27.30 was significantly (P<0.05) lower than that in non-selective breeding pigs. Titers against inactivated vaccine for swine erysipelas were determined by ELISA. Mean titer in Hp-27.30 homozygous pigs was lowest, and significantly (P<0.05) lower than in Hp-27.30/60.13 heterozygotes and Hp-60.13 homozygotes. Mean titer in the homozygous pigs with Hp-27.30 was significantly (P<0.05) lower than that in non-selective breeding pigs. Titers against inactivated vaccine for swine erysipelas were determined by ELISA. Mean titer in Hp-27.30 homozygous pigs was lowest, and significantly (P<0.05) lower than in Hp-27.30/60.13 heterozygotes and Hp-60.13 homozygotes. Mean titer in the homozygous pigs with Hp-27.30 was significantly (P<0.05) lower than that in non-selective breeding pigs. Titers against inactivated vaccine for swine erysipelas were determined by ELISA.

Poster 2038

Title: Syndactyly in a German Holstein cow – a case report

Presenting Author: Claudia Floren, Institute of Veterinary Medicine, University of Goettingen, Burckhardtweg 2, 37077 Goettingen, Germany

Other authors (name only):
1. Ekkehard Schütz
2. Bertram Brenig

Abstract:
Syndactyly in Holstein cattle, also called Mulefoot disease (MFD), is an autosomal recessive abnormality characterized by the fusion of the functional digits. An identity-by-descent linkage study mapped the disorder to BTA 15. The region is homologous to a segment of murine chromosome MMU 2 containing the low density lipoprotein receptor-related protein 4 gene (LRP4). Studies in mouse showed that LRP4 plays an essential role in the process of digit differentiation in mammalian species. Recently, six mutations associated with syndactyly have been detected in cattle. All newly described mutations affect different conserved protein domains but do not explain all analyzed cases of syndactyly. Here, an affected Holstein cow, with two syndactylous hind foot claws has been investigated for these six mutations. In addition, exons 2 to 38 of LRP4 and the respective splice/donor sites were comparatively sequenced using the affected animal, six unaffected randomly chosen animals and one heterozygous carrier for MFD. Surprisingly, the affected cow did not reveal any of the described SNPs associated with the phenotype. This confirms the supposition of additional mutations causative for syndactyly. Further experiments will focus on the 5'UTR and exon 1 to clarify whether these regions might be involved in this case of syndactyly.

Poster 2039

Title: SNP within the porcine NAMPT/PBEF1/visfatin gene is associated with fat and carcass traits

Stanislav Cepica1, Heinz Bartenschlager2, Cristina Ovilo1, Almudena Fernandez2, Jana Zrustova4, Martin Masopust1, Ales Knoll4, Hermann Geldermann2

1 Institute of Animal Physiology and Genetics ASCR, Libecheov, Czech Republic
2 Department of Animal Breeding and Biotechnology, University of Hohenheim, Stuttgart, Germany

Abstract:
Syndactyly in Holstein cattle, also called Mulefoot disease (MFD), is an autosomal recessive abnormality characterized by the fusion of the functional digits. An identity-by-descent linkage study mapped the disorder to BTA 15. The region is homologous to a segment of murine chromosome MMU 2 containing the low density lipoprotein receptor-related protein 4 gene (LRP4). Studies in mouse showed that LRP4 plays an essential role in the process of digit differentiation in mammalian species. Recently, six mutations associated with syndactyly have been detected in cattle. All newly described mutations affect different conserved protein domains but do not explain all analyzed cases of syndactyly. Here, an affected Holstein cow, with two syndactylous hind foot claws has been investigated for these six mutations. In addition, exons 2 to 38 of LRP4 and the respective splice/donor sites were comparatively sequenced using the affected animal, six unaffected randomly chosen animals and one heterozygous carrier for MFD. Surprisingly, the affected cow did not reveal any of the described SNPs associated with the phenotype. This confirms the supposition of additional mutations causative for syndactyly. Further experiments will focus on the 5'UTR and exon 1 to clarify whether these regions might be involved in this case of syndactyly.
muscles measured immediately after, 1 hour after and 3 hours after exercise. The transcriptional profiles are global transcriptional profiles in porcine skeletal muscle exposed to a single bout of endurance exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a singlebout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investigate the impact of a single bout of endurance exercise on the phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The phenotype of skeletal muscle cells is highly adaptive to different types of stimulus such as stress or exercise. The aim of the present study is to investi...
other variations were synonymous. These and future results will be the base of subsequent statistical analyses that will test the possible influence of the identified polymorphisms on milk production traits.

**Poster 2042**

**Title:** SNaPshot™ assay successfully determines SNP for chestnut coat color in historic Thoroughbred horses

Presenting Author: *M. Whitten (Max-Plank Institute for Evolutionary Anthropology, Deutscher Platz 6, D-04103, Leipzig, Germany)

M.G. Campana (University of Cambridge, Cambridge, United Kingdom)

Other authors (name only):
1. M.A. Bower
2. A.M. Murphy
3. C.J. Edwards
4. M. Binns

*Mark Whitten and Michael G. Campana are co-presenting this poster at ISAG 2008. Mark Whitten shall be listed as first author, with M.G. Campana being second.

**Abstract:**
A single missense mutation at position C901T in the melanocyte-stimulating hormone receptor gene (MC1R) causes chestnut coat color in horses. Since detailed historic records have been kept by Thoroughbred horse breeders from the 17th century, we used this mutation to address the reliability of the SNaPshot™ technology, a single base extension SNP-genotyping assay, on ancient DNA extracts from the bones and teeth of historic Thoroughbred horses. In a blind test, we compared the coat colors recorded in Thoroughbred pedigree records with the genotypes obtained from bones and teeth of historic Thoroughbred horses. In all cases, the genotypes of the historic Thoroughbred horses were consistent with the phenotypes derived from historical records. We propose that using a SNaPshot™-genotyping assay is a simple, robust and reliable method for recording SNP data in historic samples. This methodological development opens the door to in-depth studies of past phenotypic and genotypic diversity, and allows us to address questions such as the prevalence of inheritable disease and selective breeding in the past.

**Poster 2043**

**Title:** Polymorphism of 17 microsatellite loci in Akhal-Teke, Arabian and Thoroughbred horses in Russia

Presenting Author: LYUDMILA KHRABROVA, The All-Russian Research Institute for Horse Breeding, Laboratory of Genetics, P.O. Divovo, Ryazan region, Russia

Other authors (name only):
1. LILIYA KALINKOVA
2. MARINA ZAITCEVA
3. ALEXANDER ZAITCEV

**Abstract:**
For studying genetic diversity within three famous horse breeds, Akhal-Teke (n=109), Arabian (n=125) and Thoroughbred (n=476), a set of 17 microsatellites (StockMarks®) was used. Comparison between these populations demonstrated that each breed has some alleles, which were not found in other breeds yet. The oldest Akhal-Teke breed had the highest genetic variability – 139 alleles (among them 28 unique alleles) in 17 loci. There were more individual genetic variations in Akhal-Teke horses (Ae=3,72, H0=62,7%), than in other two breeds. In Arabian and Thoroughbred horses 102 alleles were found, among them 4 and 7 unique ones, respectively. The population of Arabian horses in Russia, traced to Tersk stud, was characterized by high consolidation (Ae=2,92). Horses of three studied breeds had specific allele spectrum of satellite DNA for 16 loci and similarity was found only in locus HMS2. Estimated coefficients of genetic similarity demonstrate more close relationship between Arabian and Thoroughbred horses (0,780). Akhal-Teke horses show less significant resemblance to Thoroughbred (0,727) and Arabian breeds (0,729). High polymorphism of microsatellite, blood group and biochemical markers in Akhal-Teke horses proves the ancient origin of the breed.

**Poster 2044**

**Title:** Genomic DNA markers analysis in hybrid lines of pigs

Lenka Putnova1, Irena Vrtkova1, Josef Dvorak1, Nadezda Kernerova2, Vaclav Matousek2

1 Mendel University, Laboratory of Agrogenomics, Zemedelska 1, 613 00 Brno, Czech Republic
2 University of South Bohemia, Faculty of Agriculture, Studentska 13, 370 05 Ceske Budejovice, Czech Republic
Abstract:
Genetic variability of four porcine genes (FUT1, MC4R, MYF4 and RYR1) with considerable economic importance and ten microsatellite markers (S0068, SW24, S0107, S0355, S0380, SW353, SW936, SW72, S0070 and TNFB) was studied. SNP and microsatellite genotyping was done in 59 animals of crossbred pigs (CL x CLW, CLW x P and CLW x D). The following allele frequencies were observed: 0.25 for allele A and 0.75 for allele G of FUT1 gene, 0.50 for allele A and 0.50 for allele B of MC4R gene, 0.79 for allele A and 0.21 for allele B of MYF4 gene and 0.92 for allele N and 0.08 for allele n of RYR1 gene. A total number of 72 distinct alleles were obtained for microsatellite loci. The number of alleles at individual loci ranged from 4 (S0355) to 10 (TNFB). The highest heterozygosity (above 80%) was observed for locus S0107, SW24, S0070 and TNFB. The highest polymorphism information content (PIC>0.80) was determined for locus S0107 and S0070. The probabilities of paternity exclusion/one parental genotype unavailable/and parentage exclusion were for this microsatellite panel 99.96%/99.00%/99.99% in all hybrid pigs. The study was supported by grants of Ministry of Agriculture of the Czech Republic 1G58073 and QG60045.

Poster 2045
Title: Mapping genes associated with Recurrent Airway Obstruction (RAO) in horses
Presenting Author: June Swinburne, Centre for Preventive Medicine, Animal Health Trust, Lanwades Park, Kentford, Newmarket, Suffolk CB8 7UU, UK

Other authors (name only):
1. Sarah Blott
2. Helen Bogle
3. Jolanta Klukowska
4. Elizabeth Temperton
5. Vince Gerber

Abstract:
RAO is a severe inflammatory airway disease typically affecting middle-aged horses and resulting in coughing and exercise intolerance due to increased mucus production and bronchoconstriction. It is initiated by the inhalation of dust and may be a model for human asthma. Once sensitised, a horse becomes hypersensitive to airborne particles and requires careful management. RAO is a common disease caused in part by inherited factors and influenced by environmental conditions. We have undertaken extensive sampling from Swiss Warmblood horses consisting of two half-sibling families based around RAO-affected stallions. These offspring (n=132; n=98) have been graded using a general scale 1-4 which incorporates the typical clinical symptoms, and also for individual symptoms such as coughing and nasal discharge. We have performed a whole genome scan, using a panel of microsatellite markers (average spacing 8.5cM), in order to locate genes influencing this condition. The genotyping data is being analysed using a regression interval QTL mapping method for out-bred half-sib families. We will present our data showing the positions of QTLs associated with this condition. In addition unrelated horses, and other European Warmblood breeds, have been collected in order to perform an association study using the recently available 60,000 SNP chip.

Poster 2046
Title: Analysis of gene expression in horned and polled tissue from Brahman cattle using the Agilent Bovine microarray.
Presenting Author: Maxy Mariasegaram, CSIRO, St Lucia, Australia

Other authors (name only):
1. Sigrid Lehnert
2. Toni Reverter
3. Wes Barris
4. Kishore Prayaga

Abstract:
We present results from a study examining the gene expression differences involved in the development of horned and polled phenotypes in Brahman cattle. Tissue biopsies were obtained from the skull region of new born calves to obtain a total of 11 samples representing phenotypes corresponding to 3 horned, 4 scurred and 4 polled animals (3 treatments). Hybridization against the 4x44K Agilent bovine microarray was conducted by the SRC microarray facility (IMB, University of Queensland). Each chip contains 44,000 unique 60mer oligos representing 21,475 individual genes which are deposited using a non-contact inkjet system. The analysis was conducted using a mixed model approach with array, chip, dye and sex as fixed effects; probe, probe*treatment, probe*sex and residual as random effects. A total of 733 differentially expressed probes were identified across various contrasts of interest i.e. Poll vs. Horn, Poll vs. Scur, and Horn vs. Scur. Preliminary inspection points to the existence of several differentially expressed pathways specific to the polled phenotype. One pathway of significant interest relates to genes involved in cell adhesion such as desmocollin (DSC), desmoglein (DSG) and related cadherin genes that appear to be up regulated in polled animals versus horned.
**Poster 2047**

**Title:** miR-26a targets the histone methyltransferase Enhancer of Zeste homolog 2 during myogenesis

Presenting Author: Ross L. Tellam, CSIRO Livestock Industries, Queensland Bioscience Precinct, 306 Carmody Rd, St Lucia 4067, QLD, Australia

Other authors (name only):
1. Chung Fai Wong

**Abstract:**
MicroRNA (miRNA) are important regulators of many biological processes. However, the biological targets for most miRNA are still poorly defined. In this study we profiled the expression of miRNA during myogenesis from proliferating myoblasts through to terminally differentiated myotubes. Microarray results identified six significantly differentially expressed miRNA that were more than two-fold different in expression level in myotubes compared to proliferating myoblasts. From this list miR-26a was further examined. Over-expression of miR-26a in murine myogenic C2C12 cells induced creatine kinase activity, an enzyme that markedly increases during differentiation of the cells into myotubes and the myogenic transcription factors myoD and myogenin. These results indicated that increased expression of miR-26a promoted myogenesis. Through a bioinformatics approach we identified the histone methyltransferase, Enhancer of Zeste homolog 2 (Ezh2), as a potential target of miR-26a. Over-expression of miR-26a in murine myogenic C2C12 cells induced creatine kinase activity, an enzyme that markedly increases during differentiation of the cells into myotubes and the myogenic transcription factors myoD and myogenin. This research links a miRNA with epigenetic mechanisms controlling myogenesis.

**Poster 2048**

**Title:** Genetic diversity and individual assignment in major cattle breeds registered in Canada.

Presenting Author: Kevin Lang, Genserve, Saskatchewan Research Council, 15 Innovation Blvd., Saskatoon, SK, Canada, S7N 2X8.

Other authors (name only):
1. Yves Plante

**Abstract:**
Base-line knowledge of genetic diversity in food producing animals provides valuable information to genetic conservation programs. Surprisingly, large population surveys of major beef and dairy cattle breeds are lacking in Canada. Here we explore genetic diversity in the Angus, Blonde d’Aquitaine, Gelbvieh, Hereford, Holstein, Limousin, Simmental, and Belgian Blue breeds. A total of 3,548 individuals were genotyped at twelve ISAG recommended bovine microsatellites. The effective number of alleles ranged from 3.00 (Hereford) to 3.92 (Belgian Blue). The average allelic richness ranged from 5.36 (Hereford) to 6.76 (Gelbvieh), whereas Nei’s gene diversity indices ranged from 0.64 (Hereford) to 0.72 (Belgian Blue). Eighteen private alleles were detected with at least one such allele found in any given breed. A frequency-based approach could assign 98.6% of Angus cattle tested to the predefined breed but only 84.4% of the Limousin cattle were properly assigned. A Bayesian approach identified eight natural clusters. Over 98% of Holsteins could be assigned to a unique cluster, but only 76% of the Blondes and Limousin cattle could be assigned to unique groups. Genealogical records and multiloci genotype databases are now used to monitor temporal changes in genetic diversity in these and other minor breeds registered in Canada.

**Poster 2049**

**Title:** Associations of IGF1 and FASN positional candidate genes with fat deposition and carcass merit traits in three unrelated beef cattle populations

Presenting Author: Changxi Li, Agriculture and Agri-Food Canada, Lacombe Research Centre, 6000 C&E Trail, Lacombe, Alberta, Canada T4L 1W1, and Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, Alberta, Canada, T6G 2P5

Other authors (name only):
1. Khandker Islam
2. Michael Vinsky
3. Ronda Crews
4. Erasmus Okine
5. Stephen Moore
6. Denny Crews

**Abstract:**
Two positional candidate genes for fat deposition and carcass merit traits, insulin-like growth factor-1 (IGF1) on bovine chromosome 5 and fatty acid synthase (FASN) on bovine chromosome 19, were examined for associations with hot carcass weight, carcass average fat, lean meat yield, carcass ribeye area and carcass marbling score in three unrelated beef populations.
cattle populations. Two gene-specific single nucleotide polymorphisms (SNPs), one for each gene, were genotyped on 463 beef steers from a hybrid population, 206 purebred Angus, and 187 purebred Charolais which had the phenotypic traits recorded. A preliminary association analysis showed that the IGF1 SNP was significantly associated with carcass average fat and lean meat yield in the Angus population, and animals with the “TT” genotype had significantly lower carcass average fat and higher lean meat yield (P<0.05). However, no association was found between the IGF1 SNP in the other populations. The FASN SNP showed no associations with carcass traits in any of the populations examined. Different IGF1, FASN SNP, as well as SNP in other genes will need to be developed to further characterize genetic control of fat-related carcass traits in beef cattle.

Poster 2050

Title: Polymorphisms in porcine TLR1, TLR2, and TLR6

Presenting Author: Ingrid-Maria Bergman, School of Pure and Applied Natural Sciences, University of Kalmar, S-391 82 Kalmar, Sweden

Other authors (name only):
1. Amelie Johansson
2. Inger Edfors

Abstract:
The Toll-like receptors (TLRs) recognize pathogen-associated molecular patterns (PAMPs) and initiate the innate as well as the adaptive immune defences. Several TLR polymorphisms associated with susceptibility to infectious diseases have been identified in man. The complete coding sequences of porcine TLR1-10 have been reported, as well as several non-synonymous SNPs in TLR1, 2, 4, 5, and 6. So far, there are few studies on associations with disease in pigs, but the TLR2/6 heterodimer is known to be activated by Mycoplasma hyopneumoniae. The present study aims to investigate polymorphisms in porcine TLR genes mapping to chromosome 8, close to earlier identified QTLs for immune-related traits. Currently, polymorphisms in TLR1, TLR2, and TLR6 are being investigated amongst animals of different breeds (Wild boar, Hampshire, Landrace, and Yorkshire). Our preliminary results indicate that fewer polymorphisms are present in TLR2 than in TLR1 and TLR6. So far, no polymorphisms have been detected in TLR2 in the Wild boars. In TLR6, most of the SNPs detected are clustered in the part of the sequence corresponding to the transmembrane domain and its vicinity, while polymorphisms are evenly distributed along TLR1 and TLR2.

Poster 2051

Title: Microsatellite markers for resistance to facial eczema disease in New Zealand cattle.

Presenting Author: C.A. Morris, AgResearch, Ruakura Research Centre, PB 3123, Hamilton New Zealand.

Other authors (name only):
1. N.G. Cullen.

Abstract:
Facial eczema (FE) is caused by the toxin, sporidesmin, from the Pithomyces chartarum fungus, found on pastures in summer/autumn in the North Island of New Zealand. In susceptible dairy cows, sporidesmin causes liver injury, and reduced milk production and survival. A secondary effect is photosensitisation, leading to visible eczema symptoms; hence the common name of the disease. Susceptibility (liver injury), measured using the enzyme GGT from peripheral blood, is a heritable trait in cattle (latest estimate 0.40 ± 0.04). In 2004-08, 11664 commercial animals in 55 Holstein-Friesian (F), Jersey (J) and FxJ herds experiencing FE following natural challenge were blood-sampled (to assay GGT activity and to extract DNA). Pedigree information was obtained from the herd owners. Breeding Values for susceptibility were returned to owners of the cows and artificial-insemination sires.

Earlier genomics work in a research beef herd affected with FE had pinpointed two chromosomal regions; 384 elite dairy cows and cow-sires were then genotyped using 26 and 12 surrounding microsatellite markers. Significances of the best markers/chromosome associated with phenotype were respectively P<0.002 and P<0.008 overall (P<0.0008 and P<0.02, within breeds). The best two markers here could be part of an FE test, but additional markers are sought.

Poster 2052

Title: Fishy-egg tainting by brown-shelled layers is recessively inherited under typical commercial conditions

Presenting Author: Fiona Buchanan, Department of Animal and Poultry Science, University of Saskatchewan, 51 Campus Drive, Saskatoon, Sask., Canada S7N5A8

Other authors (name only):
1. Alison Ward
2. Henry Classen

Abstract:
Fishy-egg taint has long been a problem associated with feeding canola meal (CM) to brown-shelled laying hens. It is caused by a SNP in the flavin-containing monooxygenase 3 gene (FMO3 984c.A>T). This mutation prevents the fishy-smelling trimethylamine (TMA) from being oxidized to the non-odourous trimethylamine N-oxide (TMAO), leading to an accumulation of TMA in developing egg yolks. TMA is produced from the bacterial fermentation of choline in the gut. Conflicting results from previous studies have found egg tainting to be recessive or additive. Both studies fed high concentrations of choline chloride to induce tainting, which does not reflect commercial production practices. Our objective was to characterize the inheritance pattern of fishy-egg tainting when hens are fed CM, reflecting typical industry practices. Diets consisting of 0, 6, 12, 18, or 24% CM were fed to 6 hens per genotype per diet (n=90). Three eggs were collected per hen and the yolks were analyzed for TMA concentration. The effects of diet, genotype, and their interaction were all significant (P<0.0001). Only hens of the TT genotype displayed increasing yolk TMA concentration with increasing CM. We therefore conclude that fishy-egg tainting is recessively inherited under standard CM feeding practices.

Poster 2053

Title: Analysis of DNA sequence variants in candidate genes for bovine spongiform encephalopathy (BSE) susceptibility located in a QTL region on bovine chromosome 17

Presenting Author: Bertram Brenig, Institute of Veterinary Medicine, University of Göttingen, D-37077 Göttingen, Germany


Abstract:
With the occurrence of the BSE crisis, several studies have evaluated sequence variations in the prion protein gene (PRNP) to assess their associations with BSE. In contrast to other mammals no functional polymorphisms in PRNP leading to a susceptibility/resistance towards BSE have been reported. Recently, associations between polymorphisms in the regulatory region of PRNP and BSE have been shown. In addition, several genome-wide DNA marker scans have been performed and quantitative trait loci (QTL) regions significantly linked to BSE susceptibility (BTA 5 and 17) were detected. We have analyzed functional and positional candidate genes located in one of these QTL regions on chromosome 17q23-24. The region flanks several cM upstream and downstream marker INRA025, which showed a genome-wide significant linkage disequilibrium in BSE positive cattle. DNA from 139 BSE cattle and 484 unaffected control cattle (Holstein, Fleckvieh, and Braunvieh) were genotyped for mutations in RNP24, PSMD9, PITPNM2, and B3GNT4. SNPs were detected in all four genes, and allele frequencies were calculated for each SNP stratified according to disease. Different statistical analysis showed that several of the SNPs are significantly associated with BSE susceptibility/resistance. Acknowledgments: The study was supported by the German Research Foundation DFG (BR992/14-1,2) and German TSE platform.

Poster 2054

Title: Polymorphisms associated with residual feed intake in growing cattle

Presenting Author: Denis Fidalis N. Mujibi, Department of Agriculture, Food and Nutritional Science, 4-10 AgroForestry Centre, University of Alberta, Edmonton, T6G2P5, AB, Canada.


Abstract:
Residual feed intake (RFI) has a moderate heritability. However, collection of feed intake data is expensive. Effective selection would therefore be enhanced if marker assisted evaluation were to be used. In this study 44 SNP, previously identified to be of interest were evaluated for association with RFI. The data consisted of 464 steers tested for feed efficiency over a period of 3 years. A mixed inheritance model was used for analysis. Marker assisted breeding values (MEBV) were calculated as the sum of polygenic EBV and SNP effect whereas accuracy of prediction was estimated as the correlation between the (M)EBV and phenotype. Three SNPs were found to be highly associated with RFI (P < 0.05). The allelic substitution effects for these SNP were -0.19 ± 0.07, -0.21 ± 0.06 and -0.23 ± 0.08 kg/d. Genotypic effects were
substantial with a difference of 0.59, 0.48 and 0.55 kg/d between the two homozygotes of the SNP. Together, the SNP accounted for 21% of the total phenotypic variation for RFI. Single trait heritability for RFI was increased by 13% when all three SNP were included in the model. Validation of these SNP in a different population will determine their utility for marker-assisted selection.

Poster 2055

Title: Mitochondrial DNA - an imperfect tool to study the origin of horse populations

Presenting Author: Iwona Głażewska, Dep.of Genetics, University of Gdańsk, Kładki 24 80-822 Gdańsk, Poland,

Other authors (name only):
1. Barbara Gralak

Abstract:
A popular tool used in the study of the origin of breeds and species is mtDNA analysis. However, since mtDNA is passed exclusively through the female line, it is an incomplete source of information on present-day population ancestors. The genetic analysis of Polish Arabian horses highlights some of the limitations of this method:

- The percentage of the mares from which mtDNA originates is only 7.0% of the total number of population founders, and the contribution of their genes to the present gene pool is 8.3%.
- The presence and frequency of haplotypes in the population is strictly dependent on the decisions of breeders as to preferred dam lines. This is reflected both in the different number of haplotypes in particular studs and in the large disproportion in haplotype frequency (0.2% - 31.6%).
- Genetic relations between particular dam lines revealed by mtDNA analysis are not reflected in protein or microsatellite marker analyses. If differences between the results of the latter two can be attributed to distinct selection pressure, the discrepancy between the dendrograms constructed using mtDNA data and those based on the remaining markers seems to arise from the imperfection of the mtDNA method.

Poster 2056

Title: Y-Chromosome diversity in cattle.

Presenting Author: Luis J. Royo; SERIDA-Somió, Camino de los Cuevales 604, E-33203 Gijón (Spain)

Other authors (name only):
1. Lucía Pérez-Pardal
2. Alain Beja-Pereira
3. Isabel Álvarez
4. Ivan Fernández
5. Amadou Traoré
6. Carmen Rincón
7. Rodolfo J. C. Cantet
8. F. Abel Ponce de León
9. Félix Goyache

Abstract:
A total of 350 male samples belonging to 41 cattle populations of Europe (244), Africa (46) and Asia (60) were genotyped for 5 Y-specific microsatellites totaling 6 different loci. The microsatellites used were tested for non-amplification on female DNA. The number of alleles identified was 6, 3, 11, 7, 8 and 6, respectively, for locus A, B, C, D1, D2 and E. Analyses allowed identifying a total of 55 different haplogroups with 182 samples (52%) having the same haplotype (H55). Most of the haplotypes found (21) were unique. Correspondence analysis carried out on the Y-chromosome haplotypes allowed the identification of three different factors explaining, respectively, 11.7%, 9.9% and 7.4% of the variance. These factors let us separate haplotypes of zebuine origin (11) from those of taurine origin (44). Within the taurine cluster the correspondence analysis would allow separating two subgroups: the main subgroup included most individuals belonging to Continental European, Iberian and African cattle breeds, whereas the second subgroup was more frequent in Northern European cattle breeds. This grouping pattern was confirmed via Network analysis. Median-joining trees informed that both the zebuine and the main taurine clusters resembled star-like shapes. Research partially funded by grants MEC CGL2005-03761/BOS and USDA-NRI 2002-35205-11627.

Poster 2057

Title: Further characterization of equine KIT mutations in horses with dominant white or variable white coat color

Presenting Author: Bianca Haase, Institute of Genetics, Vetsuisse-Faculty, University of Bern, Switzerland

Other authors (name only):
1. Gabriela Obexer-Ruff
2. Samantha Brooks
3. Teruaki Tozaki
4. Judith Howard
5. Vinzenz Gerber
6. Dominique Burger
7. Pierre-Andre Ponceet
8. Stefan Rieder
9. Telhisa Hasegawa
10. Cecilia Penedo
11. Tosso Leeb

Abstract:
We recently ascertained the genomic structure of the equine KIT gene and demonstrated that multiple independent mutations within this gene are responsible for the dominant white coat color in several modern horse breeds. Sequencing of the coding regions of KIT in additional unrelated Thoroughbred, Quarter Horse, and draft horse families revealed new mutations associated with depigmentation phenotypes. In the mouse, KIT mutations that lead to white or white spotted coat color phenotypes are often associated with pleiotropic effects such as anemia and male sterility. We analyzed various blood parameters in dominant white Franches-Montagne horses heterozygous for the KIT Y717X mutation. The results showed that Franches-Montagnes horses carrying the KIT Y717X did not have any significantly altered blood parameters with respect to age-matched solid colored Franches-Montagnes horses. Our data indicate that KIT mutations may have different effects in mice and horses, and that at least the KIT Y717X mutation does not have a major negative effect on the hematopoietic system of dominant white horses.

Poster 2059

Title: Fine-mapping of ATPP in Japanese Black cattle and search for candidate genes with microarray

Presenting Author: Takashi Hirano, Shirakawa Institute of Animal Genetics, Odakura, Nishigo, Nishi-shirakawa, Fukushima 961-8061, Japan

Other authors (name only):
1. Yoichi Sato
2. Naoya Ihara
3. Toshio Watanabe
4. Yoshikazu Sugimoto

Abstract:
Abnormal teat patterning phenotype (ATPP) is characterized by one (moderate form) or two (severe form) absent teat(s) in cattle. We have mapped ATPP to the centromeric region on BTA17 (ATPP-1), and the telomeric and centromeric regions on BTA1 (ATPP-2 and -3) with 152 affected progenies and 454 normal progenies of Japanese Black Bull A (Ihara et al., 2006; Anim. Genet. 38, 15–19). In this study, we performed a fine-mapping of the ATPP-1 region and search for candidate genes. Because severe ATPP animals had the ATPP-1 risk-haplotype from Bull A, we searched shared risk-haplotype in paternal haplotypes of all 136 severe ATPP animals, which could narrow down the ATPP-1 region to a 3.15 Mb region. To search for genes relating with teats development, we performed a microarray analysis using teats and skin of three-day-old female mice. In the corresponding region to the ATPP-1, 2 genes exhibited expression differences between teats and skin with the microarray analysis. These genes may be candidate genes for ATPP.

Poster 2060

Title: Gene discovery for reproduction rate in tropically adapted Australian beef cattle

Presenting Author: Rachel Hawken, CRC for Beef Genetic Technologies, University of New England, Armidale NSW, 2351 Australia and CSIRO Livestock Industries, Queensland Bioscience Precinct, 306 Carmody Road, St Lucia, QLD, 4064, Australia

Other authors (name only):
1. Prayaga K C
2. Collis E
3. Johnston D
4. Holroyd R G
5. Sim W
6. Williams P
7. Corbet N
8. Fordyce G
9. Tier B
10. Burns B M
11. Reverter A
12. Burrow HM

Abstract:
Female reproductive performance is a key driver of profitability of beef cattle systems, but is generally lowly heritable and only expressed relatively late in life. The Cooperative Research Centre for Beef Genetic Technologies (Beef CRC) has established a research population consisting of both Brahman and Tropical Composite cows and various productive, reproductive, adaptive and feed efficiency measures have been recorded. Two key components of reproduction rate of interest to industry are age at puberty and post partum anoestrus interval. Age at puberty is an issue for Brahmans in Northern Australia where they can be 3 or even 4 years of age before their first parturition. Furthermore, post partum anoestrus can delay ovulation beyond the mating season. Both of these traits result in increased production costs associated with the maintenance of barren cows and affecting their lifetime reproductive performance. The Beef CRC cattle resource, and research to identify genetic markers for components of reproduction rate will be described.
**Poster 2062**

**Title: The Single Nucleotide Polymorphisms of Chicken 70-kilodalton Heat Shock Protein (HSP70) Gene**

Qingbin Luo, Xiquan Zhang, Guanfu Yang
( College of Animal Science, South China Agriculture University, Guangzhou 510642, P.R. China)

**Abstract:**
The heat shock protein 70, which coded by the 70-kilodalton heat shock protein (HSP70) gene, is very important for the genesis of the heat tolerance of chicken. In the research of resistance breeding, HSP70 gene can be considered as the candidate loci of improving the sensibility of heat shock. But in present, researches about single nucleotide polymorphisms (SNPs) exist in chicken HSP70 gene are very limited. In this research, 591 individuals from 20 different Chinese native chicken breeds were studied. Denaturing High Performance Liquid Chromatography (DHPLC) technology was used to screen unknown SNPs existed in chicken HSP70 gene sequence. From every typical DHPLC peak, 2 individuals were selected to sequence. Homologous analysis was carried out based on the sequence information of samples that amplified from same primer to identity new unknown SNPs. From the compare of the typical DHPLC peak and sequence information, the genotype of every individual in each SNPs site was deduced. The genotype and gene frequency of every SNPs site in different chicken breeds were calculated and the difference exist among different breeds were tested. Average heterozygosity of different breeds, genetic diversity and its associate parameters, standard genetic distance (DST) and genetic identity parameters were calculated to measure the genetic variation exist among and between different chicken breeds in HSP70 gene. Based on the genotype of every individual in each SNPs sites, the PHASE program, which compiled by the theory of most likelihood, was used to affirm the haplotype and the distribution frequency of every haplotype in different chicken breed was calculated. In the PCR products amplified by primer A5, A6, A7 and A4, 10 SNPs sites (A258G, C276G, C507T, C1040A, G1044A, C1431A, T1476C, G1500A, A1529G and C1722T) were detected. All of these SNPs exist in the code region of HSP70 gene. Especially, the mutation of C→A happened in 1040 site lead to a substitution of the amino acid histidine for proline at residue 347 and the mutation of A→G happened in 1529 site lead to a substitution of the amino acid arginine for lysine at residue 510 in HSP70. From the compare of the typical DHPLC peak and sequence results, the genotype of every individual in each SNPs site were confirmed. To those not sequenced individuals, the genotypes in each SNPs site were confirmed according to their DHPLC peak, same peak mean same genotype. Thus the relationship between genotypes and DHPLC peak was formed. When consider the distribution frequency of genotype and gene in each SNPs site, significant difference (P<0.05) or highly significant difference (P<0.01) of mutation gene among different chicken breeds were found in eight sites (A258G, C276G, C507T, G1044A, T1476C, G1500A, A1529G, C1722T). But these frequency differences were not accordant with the difference of each chicken breed’s geographic latitude. Generally speaking, in 3 to 4 sites, high frequency of mutation gene were found and significant difference (P<0.05) exist among breeds in 10 chicken breeds, they were Xinghua, Silkie, Taihe, Chongren Partridge, Yugan sooty, Qingyuan Partridge, Huiyang Bearded, Silkie, Guangdong, Taoyuan, Taibai and Henan game. In A1529G and C1722T, the mutation allele frequency in chicken breeds distributed in two southern provinces, Guangdong and Jiangxi, were significantly higher (P<0.05) or even highly significantly higher (P<0.01) than chicken breeds distributed in northern provinces. The average heterozygosity of every chicken breed in each SNPs site of the HSP70 gene were not high, the highest one was 0.2885 which exist in Huiyang Bearded and the smallest one was 0.1813 which exist in Lueyang Black. The gene divergence (H_F) of each SNPs site were different greatly, the highest H_F was 0.4971 that exist in C276G site. The H_F values in G1500A and A258G sites were also high; they were 0.4900 and 0.4387 respectively. The G_ST values of these sites were with the same trend. They indicated that high variation exist in these sites, not only within but also among breed groups. These three SNPs sites, C276G, G1500A and A258G, were suit to be used as marker sites to judge the genetic diversity of HSP70 gene in different chicken breeds. 56 haplotypes were infered in the 10 SNPs sites of HSP70 gene exist in all of these 20 chicken breeds. 28 of them were truly exited in at least one individual. In these 28 truly exited haplotypes, H14●H21●H5●H6●H7●H9●H10●H12●H
15 □ H20 □ H25 □ H26 □ H31 □ H34 were the main types and distributed in most breed groups with relative high frequency. These 28 truly exited haplotypes hold 81% of all haplotypes frequency in this study; only 19% of haplotypes cannot be confirmed. In the truly exited haplotypes, 14 main types hold 85% of all haplotypes frequency, and the remaining 14 haplotypes hold 15% of all haplotypes and be considered as rare haplotypes. Great differences of the distribution frequency of different haplotypes among breed groups were found. In all of these haplotypes, only H14 and H21 were distributed in every breed. The distribution frequency diversity of haplotypes base on 10 SNPs in HSP70 gene were very abundant in the 20 chicken breeds.

Poster 2063

Title: Expression of genes associated to omega 3 fatty acid content in bovine skeletal muscle

Presenting Author: Isabel Tupac-Yupanqui. Dpt. Animal Production. Veterinary Faculty. UCM. Madrid Spain

Other authors (name only):
1. Raquel Pérez Gómez
2. Isabel Tupac-Yupanqui
3. Javier Cañón
4. Susana Dunner

Abstract:
Fatty acids are determinant of the sensorial quality of meat. Moreover, long chain unsaturated fatty acid (PUFA) are important for health purposes. We analyzed 120 bovine individuals for fatty acids profile and different candidate genes with the aim of associating the expression of particular genes to the amount of PUFA in skeletal muscle. Gene expression was measured through real time RTqPCR in samples from *Longissimus dorsi* of animals presenting different levels of long chain fatty acids omega 3 (LCFAω3), with the result of a significant higher expression of AANAT and UCP2 and down-regulation of AHA1 in those animals showing a high LCFAω3 phenotype. AANAT is an enzyme which catalyses the transformation of serotonin to N-acetylserotonin, the limiting step in melatonin synthesis. Both N-acetylserotonin and melatonin have been described as having protective effects against lipid peroxidation. UCP2 protein, placed on the inner mitochondrial membrane, has the ability to dissipate an excess of membrane potential produced by a strong proton gradient that induces the genesis of reactive oxygen species (ROS), rising the oxidative stress levels in the entire cell. During lipid peroxidation, fatty acids (mostly the PUFAs) lose insaturations and suffer oxidative degradation. Up-regulation of genes AANAT and UCP2 seems to prevent against ROS aggressions avoiding LCFAω3 to be degraded. Moreover, the low expression of AHA1 (a molecular co-chaperone) in samples of high LCFAω3 phenotype, reflects a reduced cellular oxidative stress.

Poster 2064

Title: Y-chromosome haplotype diversity in cattle

Presenting Author: Catarina Ginja 1,2,3
1 Departamento de Genética e Melhoramento Animal, Estação Zootécnica Nacional, Portugal
2 Instituto Superior de Agronomia, Tapada da Ajuda, Portugal
3 Veterinary Genetics Laboratory, University of California, Davis, U.S.A.

Other authors (name only):
1. Lilia Melucci
2. Jorge Quiroz
3. Roberto Martinez
4. Maria Antónia Revidatti
5. Amparo Martinez-Martinez
6. Luís Telo da Gama
7. Maria Cecília T. Penedo

Abstract:
Cattle Y-chromosome diversity was investigated in Portuguese and Spanish native breeds, and related Creoles from the Americas. European and Zebu breeds which may have influenced Creole cattle were also analyzed. Haplotypes defined by five SNPs, one indel and 7 microsatellites in the non-recombining region of the chromosome were obtained for 748 males representing breeds from Argentina (2), Brazil (1), France (2), India (2), Mexico (2), Paraguay (1), Portugal (13), Spain (3), The Netherlands (1), United Kingdom (5) and United States of America (1). The 24 haplotypes identified were classified according to defined patrilines Y1 and Y2 of *B. taurus* and Y3 of *B. indicus*. The highest diversity over all breeds was found within Creoles (0.484±0.090). Y3 haplotypes were present in high frequency in Creoles consistent with widespread male-mediated *B. indicus* introgression. Among European cattle, the Iberian breeds were the most diverse (0.234±0.259) showing 7 private haplotypes (H2Y1, H5Y1, H7Y2, H8Y2, H10Y2, H12Y2, H23Y1). The Y1 patrilines found in Iberian cattle, and also reported in aurochs, could represent ancient local haplotypes. The presence of alleles INRA189-90 in Caracu and INRA189-104 in Portuguese, Canary Island and British White breeds suggested influence of African cattle in the genetic makeup of these breeds.
Poster 2065

Title: Application of 19 microsatellites DNA markers for paternity control in dogs.

Presenting Author: Anna Radko, National Research Institute of Animal Production, Immuno and Cytogenetics Department, Krakowska 1, 32-083 Balice, Poland

Other authors (name only):
1. Ewa Słota

Abstract:
Microsatellite DNA become the international standard for individual identification and parentage verification in canine breeds. Only one commercial kit prepared by Applied Biosystems company was available on market so far for canine genotypes. This set included 10 microsatellite markers, which was not always enough for confirmation of parentage. At present the assay of 19 microsatellite markers is available for canine parentage testing, offered by Finnzymes Diagnostics company. This kit contains the following 19 loci: AHTk211, CXX279, REN169O18, INU055, REN54P11, INRA21, AHT137, REN169D01, AHTk260, AHTk253, INU005, INU030, FH2848, AHT121, FH2054, REN162C04, AHTh171, REN247M23 and Amelogenin. These markers are included in the panel of loci recommended by the ISAG. The aim of this study was to analyse the polymorphism of 19 microsatellite markers and their usefulness for parentage verification in 28 individuals of Borzoi breed in Poland. Amplified PCR products were analysed by automated sequencer ABI PRISM 3130xl Genetic Analyser. The number of alleles per locus varied from 3 (INU005) to 8 (REN162 and AHT171) alleles. Except the locus INU005, which showed the lowest variation (PIC=0.396, H=0.438) the calculated PIC values exceeded 0.6 and H values ranged from 0.677 (INUQ55) to as much as 0.84 (REN162). Based on PE it was found that incorrect pedigree in this breed can be excluded with 99.9998% accuracy using DNA analysis.

Poster 2066

Title: POLYMORPHISM OF THE OPN GENE AND mRNA EXPRESSION PATTERNS OF THE OPN IN SOWS OVIDUCT

Presenting Author: ...Agnieszka Korwin-Kossakowska, Institute of Genetics and Animal Breeding, Polish Academy of Sciences, Jastrzębiec, Poland

Other authors (name only):
1. Dorota Goluch
2. Mariusz Pierzchala
3. Joanna Wyszyńska-Koko
4. Grażyna Sender

Abstract:
The aim of our study was to find the relationship between the polymorphism of the OPN (osteopontin) gene, it’s expression in female ovaries and oviduct and reproductive performance traits of sows. Sixty sows after first mating was slaughtered and tissues samples of the ovaries and oviduct were taken. The total RNA was extracted from the frozen tissues according to the protocol of Chomczynski and then subjected to Real Time PCR analysis. The polymorphisms within OPN gene for all sows were examined, simultaneously. The polymorphism in intron 6 of this gene was examined base on literature data (but with PCR fragments size 498 and 193 bp). The new polymorphism in promoter region was found. The fragment of 274 bp was chosen because of its contents of tree specific sites: type II collagen “silencer” sequence at -682 glucocortycoid response site at -658 and CAAT box -592. The results of first parity (number of piglets born alive and piglets weaned, litter weight at born and at weaning, age at farrowing) were also collected. The statistical analysis between the polymorphism of the OPN gene, the expression of this gene in ovaries and oviduct tissues and reproductive performance traits of the sows is under evaluation.

Poster 2067

Title: DNA Assignment for profitable production decision

Rogberg-Muñoz, A1; JP Lirón1; MV Ripoli1; EI Francisco1; EE Villegas-Castagnasso1; DM Posik1,2; P Peral-García1; G Giovambattista1

1 IGEVET, CCA La Plata, FCV-UNLP-CONICET. Calle 60 y 118 S/N CP: 1900 - La Plata - Buenos Aires - Argentina
2 Comisión de Investigaciones Científicas (CIC). Calle 526 entre 10 y 11 CP: 1900 - La Plata - Buenos Aires – Argentina

Abstract:
Extensive animal production can compromise relation between productivity and information; when both are necessary, technology can be useful. Wagyu breed is known for its marbling capacity and for improving carcass quality. Feedlot feeding system is more expensive than pasture, making economically important to decide which animal will be feedlot fattened. This study was done in a pasture extensive, commercial herd. 164 Angus and 26 half Wagyu/Angus dams were artificial inseminated with Wagyu, and mated after with Angus bulls. At 15 months, animals were moved for 300 days to a feedlot and Breed assignment was performed (there was no
more semen available for paternity). DNA was extracted from hair and eleven microsatellites were genotyped. Wagyu percentage was determined using Structure 2.2 software, Angus and Wagyu animals were included as control populations. From the 190 dams, 29 animals were Angus sired and 161 Wagyu sired. Ultrasound IMF is being scanned every two months, and all the animals Wagyu sired assigned (50% Wagyu or more) have satisfied expected grade and this was significantly higher; this will be confirmed at slaughter stage. This method appears to be adequate for select animals when breed assignment is economically important, i.e. for animal buyers for long term feedlots.

**Poster 2068**

**Title: Whole genome association study of adaptive traits in tropical beef cattle**

Prayaga KC1, Chan EKF2, Reverter A2, Johnston DJ3, Hawken RJ2, Barendse W2, Fordyce G4, Burrow HM5

1CSIRO Livestock Industries, JM Rendel Laboratory, Rockhampton, QLD, 4702, Australia
2CSIRO Livestock Industries, Queensland Bioscience Precinct, 306 Carmody Road, St Lucia, 4064, Australia.
3Animal Genetics and Breeding Unit, University of New England, Armidale NSW, 2351 Australia
4Queensland Department of Primary Industries and Fisheries, Charters Towers, QLD, 4820, Australia
5CRC for Beef Genetic Technologies, University of New England, Armidale NSW, 2351 Australia

**Abstract:**
Tropical adaptation in northern Australia is characterised by an animal’s ability to survive stressors such as ecto-parasites (ticks and buffalo flies), endo-parasites (worms), heat, humidity and seasonal nutrition. The Cooperative Research Centre for Beef Genetic Technologies has developed a resource population and quantified several of these tropical adaptive measures in terms of tick scores, faecal egg counts, fly lesion scores, rectal temperatures under hot conditions, coat score, coat colour, and flight time as a measure of temperament. A subset of this resource population constituting 582 Brahman females was genotyped using the Affymetrix 10K SNP panel. Genome wide association analyses were conducted using a variety of parametric and non-parametric methods. Significant chromosomal regions across the genome were identified for various adaptive traits under study. In this report, we present these broader chromosomal regions and highlight the possibility of developing genetic markers for implementation in the selection programmes.

**Poster 2070**

**Title: Effect of a single nucleotide polymorphism within promoter region of the bovine myogenic factor 5 (Myf5) gene on the gene expression in bovine m. longissimus dorsi.**

Authors; Presenting Author: Dagmara ROBAKOWSKA-HYZOREK, Institute of Genetics and Animal Breeding, Polish Academy of Science (IGAB PAS), Jastrzębie, 05-552 Wólka Kosowska, Poland

Other authors (name only):
1. Beata ŻELAZOWSKA
2. Lech ZWIERZCHOWSKI

**Abstract:**
Myogenic factor 5 (Myf5), belongs to the MRF family of transcription factors which control the progress of myogenesis and are candidates for molecular markers of meat production in farm animals. We detected nine single nucleotide polymorphisms (SNPs) located in the promoter region of the bovine Myf5 gene. The polymorphisms found are: G-1569A, A-823G, T-472G, G-882T, C-741G, T-511G, C-1185T, G-1466A (RFLP/AvaII), C-1128T (RFLP/NsiI). Using a Real Time PCR method we analyzed the possible influence of the promoter mutation G-882T (represented by all 3 genotypes - GG, GT, TT) on the expression levels of the Myf5 gene in the muscle (m. longissimus dorsi) of 12-months-old bulls of different breeds. As a reference the expression of the TBP gene was used. Statistically significant differences (evaluated by ANOVA method) were found in the Myf5 expression levels between GG and TT genotype bulls (p<0.001), and between GT and TT genotype (p<0.01) bulls. TESS software analysis have shown that the G→T substitution creates the Hoxa5 transcription factor binding site, while in the G allele the LEF-1 site is created. These results suggest that the G-882T polymorphism may affect Myf5 expression at the transcript level in the adult bovine muscle.

**Poster 2071**

**Title: UCP2: Structural and expression studies in Casertana pigs**

Presenting Author: MARIASILVIA D’ANDREA
University of Molise, Campobasso, Italy e-mail: dandrea@unimol.it

Other authors (name only):
1. ANGELA CAMPANELLA
2. FABIO PILLA
Abstract:
Casertana is an autochthonous pig breed characterized by massive accumulation of backfat and low growth rate. In this study Casertana pigs were compared with Large White focusing on genetic variation in Uncoupling Protein 2 (UCP2). UCP2 is an uncoupling protein that functions as an uncoupler of oxidative phosphorylation, thus dissipating energy as heat. The UCP2 gene is expressed in adipose tissue and skeletal muscle. Several studies found that uncoupling proteins are associated with the biological traits of body weight, resting metabolic rate and energy conversion.

17 Casertana and 16 Large White, of the same age and raised outdoors in the same environmental conditions, were analysed.

Sequencing the whole gene in Casertana and Large White 3 new SNPs were found. Two PCR-RFLP methods and a SNaPShot method were used to genotype the population.

The Real-Time PCR was carried out by amplifying, in singleplex, the target (UCP2) and the reference (TATABP) genes. Results were analysed by using the relative standard curve method.

The difference between the two groups in UCP2 mRNA levels was statistically (p<0.01) supported in muscle and adipose tissues as well. These findings suggest that UCP2 is a valid candidate gene for meat production traits.

Poster 2072

Title: GENETIC CHARACTERIZATION OF ORLOV TROTTER

Presenting Author: LILIYA KALINKOVA, The All-Russian Research Institute for Horse Breeding, Laboratory of Genetics, Divovo, Ryazan region, Russia

Other authors (name only):
1. LYUDMILA KHRABROVA
2. MARINA ZAITCEVA
3. ALEXANDER ZAITCEV

Abstract:
Orlov Trotter is the oldest Russian cultural breed of horses. The history of the breed goes back to 1776 when the stud at Khrenovoe was established by the count Alexei G. Orlov. Founder animals were imported from different countries of the world: Arabia, Denmark, the Netherlands, Italy, Spain, England, Germany, Poland, Persia and also from regions of the Don River, the Caucasus and Central Asia. In the XX century the unique breed suffered an extreme decrease in population size. In this study genetic variation of Orlov population (n=803) was evaluated in 8 blood polymorphic systems (EAA, EAC, EAD, EAK, AI, Es, Tp, Xk). In addition genetic diversity within a group of leading stallions (n=20) was measured with the use of 17 microsatellite loci (StockMarks®). DNA was isolated from frozen semen stored in liquid nitrogen and hair roots. Allele frequencies, the number of alleles per locus and heterozygosities were determined. The results were compared to genetic variation observed in other horse populations of Russia (28 breeds). The level of genetic diversity in Orlov Trotter was relatively high. The panel of microsatellites evaluated in the group of Orlov stallions showed very high heterozygosity (Ho=42,11 to 100%, the mean Ho=70,13%). Effective numbers of alleles were the highest at loci ASB17 (5,48), VHL20 (5,16), HMS2 (4,56), HMS3 (4,46), ASB23 (4,13). The substantial genetic variability of Orlov Trotter may be due to extraordinary various origins of historical founders of the breed.

Poster 2073

Title: Genetic characterization of the Jeju horse compared with other horse breeds reared in Korea by means of microsatellites

GIL J. CHO1, SUNG K. CHOI1, EUN W. LEE1, CHANG Y. CHO2 & SUNG H. YEON2

1 College of Veterinary Medicine, Kyungpook National University, Daegu, Korea.
2 Animal Genetic Resources Station, National Livestock Research Institute, Namwon, Korea.

The Jeju horse (JH) is a native Korean breed selected for riding, racing and meat; its name comes from Jeju Island in South Korea. According to Jeju Province, the JH comprise about 850 herds. They have been separated into distinct areas, and have been used as a racing horse at the Jeju race park of the Korea Racing Authority. Genetic characterization of the JH was performed to construct a correct pedigree of the JH family. A total of 581 horses including 278 JH, 134 Thoroughbred-TB, 118 Jeju racehorse-JR, 31 Mongolian-MG, 12 Warmblood-WB and 8 Miniature-MN were genotyped using 20 microsatellite loci (AHT4, AHT5, ASB2, ASB17, ASB23, ASB25, CA425, HMS1, HMS2, HMS3, HMS5, HMS6, HMS7, HTG4, HTG6, HTG7, HTG10, LEX3, TKY297, VHL20). The CERVUS and DISPAN software were used to calculate allele frequencies, heterozygosity, PIC, PE, genetic distances and phylogenetic trees. All microsatellites were polymorphic in this study. The number of alleles varied from 5 to 13 (mean 8.45) in the JH. The observed heterozygosity and expected heterozygosity ranged from 0.4176 to 0.8647 and from 0.3731 to 0.8809, respectively. The polymorphic information contents (PIC) ranged from 0.3470 to 0.8670. AHT4, AHT5, ASB2, ASB17, ASB23, ASB25, CA425, HMS1, HMS2, HMS3, HMS5, HMS6, HMS7, HTG4, HTG6, HTG7, HTG10, LEX3, TKY297, VHL20 loci had
relatively high PIC values (0.7). The total exclusion probability (PE) of the 20 microsatellite loci was 0.9999 in the JH. Phylogenetic trees constructed using Neighbour-joining method showed two clear separate clusters: the first includes the JH, MG and MN, the second contains the JR, WB and TB. These results provide basic information for developing an accurate pedigree and will be useful in making decisions regarding conservation of the JH.

**Poster 2074**

**Title:** Assignment of porcine full-length cDNA sequences and ESTs on the pig draft genome sequences

**Presenting Author:** Hirohide Uenishi, National Institute of Agrobiiological Sciences 2-1-2 Kannondai, Tsukuba, Ibaraki 305-8602, Japan

**Abstract:** We have determined more than 10,000 complete sequences of pig cDNA clones derived from full-length-enriched cDNA libraries. These sequences are useful for exploration of candidate genes for economically important traits in pigs, as well as functional analysis of pig genes. We estimated chromosomal locations of cDNA sequences that had been fully determined and contig sequences of ESTs used for development of SNPs, which had been localized on a linkage map. Meanwhile, pig genome sequencing is now performed by the international consortium using BAC clones, and draft sequences of nearly half of the genome have been open to the public. We used draft sequences of the BAC clones mapped on the pig chromosome for the alignment of the cDNA and EST contigs. We could determine the relationship of 5,739 cDNA clones with the BAC sequences among 10,147 clones. Furthermore, among 967 sequences used for development of SNPs that were mapped on chromosome, we could tag 542 to the BAC sequences, and the order of the assigned sequences on chromosomes well coincide with the BAC fingerprint map of the consortium. These data demonstrate usefulness of full-length cDNA sequences and ESTs for SNP development in particular regions or genes of interest.

**Poster 2075**

**Title:** Nutritional effects on epigenetic modification and their inheritance – F0 generation

**Presenting Author:** Martin H. Braunschweig, Institute of Genetics, University of Berne, Bremgartenstrasse 109A, CH-3001 Berne

**Conference registration number of Presenting Author:** 0070

**Other authors (name only):**
1. Nasikhat Stahlberger-Saitbekova
2. Martin H. Braunschweig

**Abstract:** In a three generation pig feeding experiment we are investigating whether differential nutrition of F0 boars affects gene expression and carcass traits in the next but one F2 generation. It is hypothesized that the nutrition related transgenerational effects are epigenetic in nature and the transmission is through the male line. This implies that nutrition induced epimutations are not reprogrammed in gametogenesis or in the early embryo resulting in transgenerational epigenetic inheritance. Two groups of eight F0 boars received either a control diet or a methyl donor supplemented diet. Upon puberty the F0 boars were mated to sows to produce the F1 male generation, which was then later used to produce the final F2 generation. We measured IGF2 and IGF2R gene expression in muscle, kidney and liver tissues of F0 boars. The imprinting status of the IGF2 gene in these tissues was analysed by means of the microsatellite SWC9. DNA methylation patterns were determined in differentially methylated regions (DMR) in the IGF2 gene. The expression levels of IGF2 and IGF2R genes do not differ between the two groups of F0 boars. DNA methylation patterns around the IGF2 locus in tissues of F0 boars receiving either the methyl donor supplemented diet or the control diet are presented.

**Poster 2076**

**Title:** Mitochondrial DNA variability and genetic relationships in four Italian donkey breeds (*Equus asinus*)

**Presenting Author:** M.C. Cozzi, Dipartimento di Scienze Animali - Sezione di Zootecnica Veterinaria - Università degli Studi di Milano, Via Celoria, 10, 20133 Milano – Italy

**Other authors (name only):**
1. P. Valiati
2. R. Cherchi
3. M.G. Strillacci
4. A. Corona
5. G. Pertica
6. B. Gandolfi
7. M. Polli
8. M. Longeri
9. L. Guidobono Cavalchini
Abstract:
The genetic variability of mitochondrial D-loop DNA (mtDNA) in 91 individuals from 4 donkey breeds reared in Italy (Asinara donkey ASD, Sardinian donkey SAD, Romagnolo donkey ROD and Ragusano donkey RAD) were analysed. A 478 bp fragment of the mtDNA was amplified and sequenced. Sequences obtained were aligned with the complete donkey mtDNA sequence (GenBank#X79547), using the CLUSTALX software. 20 polymorphic sites and 10 haplotypes were identified. Overall and per breed haplotype diversity (Hd), nucleotide diversity (π), average number of nucleotide differences (k) and Fu’s Fs statistic were calculated using DnaSP 4.2 software. Hd ranged from 0.638 (ASD) to 0.867 (RAD), overall 0.822; π ranged from 0.005 (RAD) to 0.020 (ROD), overall 0.017; k ranged from 2.200 (RAD) to 9.750 (ROD) overall 8.315. These results indicated moderate haplotype diversity. Italian donkey mtDNA haplotypes were compared with about 400 worldwide donkey mtDNA sequences from on-line database. Genetic distances were estimated by the Kimura-two parameter method and a phylogenetic tree was constructed using the UPGMA algorithm by MEGA 4.0 software package. Median-joining network and mismatch analysis were calculated in order to investigate haplotypes relationship using NETWORK 4.5.0.1. The results were discussed to investigate origin of the Italian donkey breeds.

Poster 2078
Title: Analysis of μ-Calpain and Calpastatin genes in Piemontese breed
Presenting Author: Liliana DI STASIO, Dip. Scienze Zootecniche - Via L. da Vinci 44 - 10095 Grugliasco (Italy)
Other authors (name only):
1. Claudio LISA
2. Alberto BRUGIAPAGLIA

Abstract:
Mutations at the μ-Calpain (CAPN1) and Calpastatin (CAST) genes have been shown to affect beef tenderness and the reported associations have been confirmed by validation studies. As no data exist on purebred Piemontese subjects, we analysed the variability at the two loci in the breed for evaluating the usefulness of integrating the markers into selection programmes. Fifty-three unrelated Piemontese subjects, representative of the breed, were tested for CAPN1 g.5709G>C (AF252504), CAPN1 g.6545C>T (AF248054), and CAST g.282C>G (AY008267) mutations, with PCR-RFLP procedures set up in our laboratory, and the allele frequencies were calculated. For a preliminary field evaluation of the marker effects, seventy meat samples from commercial Piemontese animals were genotyped for the three SNPs and tenderness was measured as shear force (WBs) at different days post-mortem. The frequencies of the C alleles, shown to be associated with improved tenderness, were 0.14 for CAPN1 5709G>C, 0.52 for CAPN1 6545C>T, and 0.63 for CAST 282C>G. The observed genotypic distributions were in agreement with HW equilibrium. Significant associations with meat tenderness (P=0.01) were observed only for the CAST 282 marker, with a favourable effect of the C allele, consistent during the ageing (~1.997 kg at day 1 to ~0.956 kg at day 11).

Poster 2079
Title: The effect of gene CAST polymorphism on productive traits in pigs
Presenting Author: ...Paweł Urbański, Institute of Genetics and Animal Breeding, Polish Academy of Sciences, Jastrzębiec, Poland
Other authors (name only):
1. Mariusz Pierzchała
2. Arkadiusz Terman
3. Jolanta Kuryl
4. Marian Kamyczek
5. Marian Różycki

Abstract:
Calpastatin is an endogenous inhibitory protein acting on calpain, the calcium dependent cysteine proteases in animal cells and the system calpain-calpastatin is important in postnatal skeletal muscle cells. The porcine CAST gene has been mapped to chromosome 2. The aim of this study was to evaluate the effect of two mutations localized in exon 14 porcine CAST gene, described by Ciobanu on growth rate and carcass traits. The study covered about 620 gilts of two breeds: Polish Large White (PLW) and Polish Landrace (PL) and synthetic line L990. The pigs were fattened and slaughtered at the Pig Testing Station of the National Research Institute of Animal Production. In our studies we observed a significant relation between mutation identified with PvuII restriction endonuclease and different productive traits. Respecting to second studied of mutation, recognized by Hpy188I enzyme we did not the polymorphism. The animals which we analysed in our studies were monomorphic. In our opinion the calpastatin gene may be have an important role in pig breeding and selection, but a further study should be continued on other pig breeds and lines.
Title: Localisation of the causative gene for reflex epilepsy in chicken

Presenting Author: Marine Douaud, UMR génétique Cellulaire, INRA Chemin de Borde-Rouge, Auzelle BP 52627, 31326 CASTANET-TOLOSAN CEDEX, FRANCE

Other authors (name only):
1. ...Katia Fève
2. ...Michèle Tixier-Boichard
3. ...David Gourichon
4. ...Agathe Vieaud
5. ...Cesira Batini
6. ...Robert Naquet
7. ...Alain Vignal
8. ...Frédérique Pitel

Abstract:
In human, genetic reflex epilepsy is a type of idiopathic epilepsy in which a particular sensory stimulus evokes paroxysmal manifestations only in genetically predisposed subjects. The Fayoumi Fepi strain of chickens carries a recessive autosomal mutation causing generalized reflex epilepsy in homozygous animals after stroboscopic stimulation. The mutation was discovered in several animals in 1970 by Crawford who decided to build a synthetic strain of Fepi. The objective of this project is to identify the causative recessive mutation "epi", determining the epileptic phenotype. Two informative families were obtained by crossing two carrier sires with 5 or 6 affected homozygous dams each, giving a total of 209 offspring. Almost half of the offspring were affected. A genome scan was performed using informative molecular markers chosen from public databases or developed specifically. The marker showing significant linkage with the epi mutation mapped to a microchromosome where the genome sequence is incomplete. By developing additional informative microsatellite and SNP markers, mainly through comparative mapping with human, we were able to locate the epi locus in a 6cM region. Thus, insufficient coverage of the microchromosomes in the genome sequence may limit strongly the possibility to map genes of major effect or QTLs.

Poster 2081
Title: Whole porcine genome linkage map based on the 7K porcine SNP Chip

Presenting Author: Rikke Kirstine Kaee Vingborg, Department of Genetics and Biotechnology, Faculty of Agricultural Sciences, University of Aarhus, Blichers Allé 20, P.O. box 50, DK-8830 Tjele, Denmark

Other authors (name only):
1. Vivi Raundahl Gregersen
2. Anette Høj
3. Frank Panitz
4. Zahn Bujie
5. Henrik Hornshøj
6. Kirsten Kræup Sørensen
7. Christian Bendixen

Abstract:
Constructions of high-density linkage maps are required for identification of QTLs across genomes. The generation of porcine sequences has provided a solid base for SNPs (Single Nucleotide Polymorphisms). The 7K porcine Illumina Custom
Infinium® Bead SNP Chip was used to genotype a previously described family material, consisting of offspring from 12 different boar families, encompassing 236 sows and 5,195 piglets. The data set was evaluated, and SNPs that deviated from Hardy-Weinberg equilibrium, contained extra alleles, demonstrated software or genotyping limitations or did not segregate were eliminated. Only SNPs genotyped in more than 90% of the animals were included, resulting in a data set comprising 4366 SNPs. The calculations were performed on sow families, since all sows had litters with more than one boar. This established the possibility for generating genetic maps of all chromosomes, even though a close network of SNPs segregating in all families was missing. The SNPs were physically assigned to all 20 porcine chromosomes. All sequences were blasted against the available porcine chromosomes assembly database, and through comparison with the human porcine comparative database. Here we present the preliminary evaluations of SNPs.

**Poster 2082**

**Title: Association of SNPs from a QTL region for mastitis with proliferation of S. aureus in mammary epithelial cells and udder health phenotypes**

*Presenting Author: Hannes Joerg, Institute of Animal Sciences, ETH Zurich, Switzerland*  
*Conference registration number of Presenting Author:*  
*Other authors (name only):*  
1. Hannes Joerg  
2. Fredi Janett  
3. Bruno Dietrich  
4. Urs Schuler  
5. Dagmar Steiger  
6. Gerald Stanzinger  

**Abstract:**  
Adherence to and invasion of epithelial cells in the bovine udder play a significant role in the pathogenesis of *Staphylococcus aureus* mastitis. In the present study, primary mammary epithelial cell cultures of 67 cows were incubated with *S. aureus* strains and the proliferation of *S. aureus* bacteria were categorized. The cows were daughters of 4 bulls having different QTL for mastitis on chromosome 2. The 67 cows were separated in two groups due to their paternal allele of 12 SNPs belonging to the QTL region. The highest differences of proliferation of bacteria were between the groups separated by the SNP +777 G/C of the CXCR2 gene. The group with paternal G-allele and C-allele had a mean proliferation of 1.80 ± 1.16 and 3.15 ± 1.18, respectively. The same groups did also show significant differences in udder health phenotypes. The least square mean of somatic cell score corrected for bull and lactation effects were 3.60 for the C-allele and 2.56 for the G-allele. The huge differences in somatic cell score is also due to the small number of cows in this project, but even if the effect will be reduced in the population it is still an interesting QTL for mastitis.

**Poster 2083**

**Title: An alternative angiogenesis pathway seems to be activated in Canine Inflammatory mammary carcinoma**

*Presenting Author: Susana Dunner  Dpt. Animal Production. Veterinary Faculty. UCM. Madrid Spain*  
*Other authors (name only):*  
1. Ana Sánchez-Archidona  
2. Laura Peña  
3. Susana Dunner  

**Abstract:**  
Canine inflammatory mammary carcinoma (IMC) can be considered as a spontaneous animal model of human inflammatory breast carcinoma (IBC). It is the most malignant type of canine mammary cancer with extremely poor prognosis. In IMC, growth, proliferation and metastasis of breast cancer are not angiogenesis-dependent processes, as an alternative pathway called Vasculogenic Mimicry (VM) has been reported. In VM tumour cells form tube-like structures similar to micro-vascular channels in order to feed themselves. To throw light on the active IMC angiogenesis pathway, we used RNA samples from 23 bitches (11 IMC, 6 “non-inflammatory” mammary carcinoma with histological malignant grade III, and 6 healthy mammary tissues) and measured the expression through real time quantitative PCR of different candidate genes commonly associated with angiogenesis. Some of these genes are down-regulated in IMC when compared to the other samples, specially VGFD which expression is significantly low (p<0.008), the receptor VGFR3 (p<0.01) and also VGFA and TIE2 (p<0.03). These results would indicate that a different pathway is activated in this special type of mammary cancer that allows metastasis to develop through VM in which proper tumoral cells are transformed into endothelial-like cells producing a rapid, efficient way to generate a vascular net that contributes to the high malignity showed by IMC and IBC.

**Poster 2084**

**Title: Insights into equine osteochondrosis based on genetic markers in Hanoverian warmblood horses**
Presenting Author: Virginie Lampe, University of Veterinary Medicine Hannover, Institute for Animal Breeding and Genetics, Buenteweg 17p, 30559 Hannover, Germany

Other authors (name only):
1. Claudia Dierks
2. Ottmar Distl

Abstract:
The objective of this work is fine mapping of QTL regions for osteochondrosis (OC) and osteochondrosis dissecans (OCD) in horses using intragenic single nucleotide polymorphisms (SNPs) and microsatellites in order to identify genes involved in the pathogenesis of equine OC. We developed SNPs in positional candidate genes and microsatellites located in quantitative trait loci (QTL) regions identified through a whole genome scan in Hanoverian warmblood horses. The whole genome scan included 14 paternal half-sib families with 211 horses and more than 260 highly polymorphic microsatellites. Traits regarded were OC and OCD in fetlock and hock joints. Genome-wide significant QTL for OC were located on ECA2, 4, 5 and 16. Comparative human-equine maps were employed to select candidate genes in the QTL regions. SNP markers were developed using equine BAC end, EST, whole genome shotgun sequences and equine contigs. We were able to identify significantly associated SNP markers within these QTL regions for OC. The SNPs were significant in their genotypic effects on the fetlock OC and/or hock OC. Further SNPs will be tested using the Illumina Equine BeadChip. This work is an important step towards an equine SNP marker set to be employed in horse breeding.

Poster 2085

Title: Bovine placenta transcript analysis using 454 sequencing

Presenting Author: Frank Panitz, Dept. of Genetics and Biotechnology, University of Aarhus, Blichers Allé, 8830 Tjele, Denmark

Other authors (name only):
1. Pernille K. Andersen
2. Christian Bendixen

Abstract:
The recent introduction of new high-throughput sequencing technologies (e.g. 454 LifeSciences; Solexa) allows generation of high quality transcriptional profiles from large-scale sequencing of cDNA fragments. Here, we compared different strategies for cDNA synthesis with respect to transcript abundance, gene content and polymorphic variation: By using either oligo-dT or random hexamer priming during cDNA synthesis two cow placenta cDNA libraries were obtained and sequenced in a single run on the GS20 sequencer with one region for each priming method. The sequences were clustered and compared to various databases in order to annotate the transcripts. Finally, single nucleotide polymorphisms (SNPs) were identified and mapped against the cattle genome to assess the genetic variation.

Poster 2086

Title: Identification of differentially expressed proteins during an Actinobacillus pleuropneumoniae infection by proteome analysis of BALF and lung tissue

Dennis Kahlisch, Department of Physiological Chemistry, Bünteweg 17, 30559 Hannover, Germany

Other authors (name only):
1. Doris Hoeltig
2. Falk Buettner
3. Thomas Rehm
4. Gerald Gerlach
5. Hassan Naim
6. FUGATO-consortium IRAS

Abstract:
Acute respiratory diseases pose severe economical problems in pig breeding. With FUGATO (Functional Genome Analysis in the Animal Organism) and its subproject IRAS the German Federal Ministry of Education and Research (BMBF) intends to support the identification of genetic markers that facilitate breeding with regard to immune defence and resistance. To identify potential biomarkers associated with a reduced susceptibility to bacterial respiratory tract pathogens an aerosol infection model for Actinobacillus pleuropneumoniae was utilized. Lung tissues from infected animals were subjected to proteomics analysis using two different strategies: The first employed lectin affinity chromatography of bronchoalveolar lavage fluid (BALF) to compare glycoprotein profiles. The second examined the recruitment of proteins into lipid rafts during infection. Mass spectrometry of identified protein spots on 2-D gels revealed differential expression of seven glycosylated proteins (Fetuin A, α-1-acid glycoprotein, α-1-antichymotrypsin, hyaluronidase, surfactant protein D, hemoglobin, haptoglobin) and two raft-associated proteins (lactoferrin, protein similar to S100 A9). Fluorescence Difference Gel Electrophoresis (2-D DIGE) was performed to detect race varieties in the glycoprotein expression at defined point of times during the infection. These proteins are currently under further investigation regarding their
identity and their putative role as genetic resistance markers in the respiratory tract.

**Poster 2088**

**Title** “Positional cloning of the causative mutation for bovine dilated cardiomyopathy (BDCMP)”

Presenting Author: Marta Owczarek-Lipska, Institute of Genetics, Vetsuisse Faculty, University of Berne, Bremgartenstrasse 109a, 3001 Bern, Switzerland

Other authors (name only):
1. Tosso Leeb
2. Gaudenz Dolf
3. Martin H. Braunschweig

**Abstract:**
Bovine dilated cardiomyopathy (BDCMP) belongs to the group of progressive degenerative disorders of the myocardium. BDCMP is a severe and hereditary disease and was worldwide observed in cattle of Canadian Holstein origin. The main symptoms of BDCMP are edema of the brisket, congested jugular veins and tachycardia with gallop rhythm. Necropsy findings are cardiomegaly with hypertrophy and ventricle and atrial dilatation. The typical pathological sign is a progressive transmural myocardiofibrosis. The age at onset of BDCMP range usually from two to four years. Pedigree analyses performed in Canada, Japan and Switzerland led to the common ancestral Holstein-Friesian bull ABC Refection Sovereign. An autosomal recessive mode of inheritance was suggested and the BDCMP locus was mapped to the bovine chromosome 18 (BTA18). A BAC contig consisting of 10 INRA BACs and 24 CHORI BACs was constructed by PCR screening. The BAC contig spans the interval between microsatellites MSBDCMP06 and BMS2785, corresponding to a physical distance of about 6.7 Mb. Recently, the region harboring the disease locus was fine mapped to an interval covered by four adjacent INRA BACs. Currently, we are focusing on the confirmation and on the further fine mapping of the interval containing the BDCMP locus.

**Poster 2089**

**Title:** Characterization of a deletion in the cattle Pro-opiomelanocortin gene.

Presenting Author: Heather Deobald, Department of Animal and Poultry Science, College of Agriculture and Bioresources, 51 Campus Drive Saskatoon, Sk, Canada S7N 5A8

Other authors (name only):
1. Sheila M. Schmutz
2. Fiona C. Buchanan

**Abstract:**
Pro-opiomelanocortin (POMC) is a complex gene with four gene products that play roles in the appetite, stress and pigmentation pathways. Previous data suggest that a SNP in POMC (POMC c.288C>T) is associated with shipping and hot carcass weight in cattle. While optimizing this variant using real time PCR an unexpected shift in melting temperatures was observed. Sequencing these animals revealed a 12 bp deletion (POMC c.293_304delTTGGGGGCCGG) that results in four amino acids being removed (Val, Gly, Gly, Ala), and occurs five nucleotides downstream from the original SNP. The deletion starts at the second position of the 98th codon however it does not cause a frame shift. The allele frequency of the deletion was 2% in 386 steers and was only observed with the T allele. No homozygous deletion animals were found. A Mann Whitney U test showed an association with the presence or absence of the deletion and ultrasound rib-eye area at end of backgrounding (P= 0.036) and carcass rib-eye area (P=0.028).

**Poster 2090**

**Title:** A NOVEL HSP90AA1 PSEUDOGENE IN THE OVINE LATXA BREED

Presenting Author: Ane Marcos-Carcavilla, INIA (Dpto. Mejora Genética Animal), Ctra La Coruña Km 7.5, 28040 Madrid (Spain)

Other authors (name only):
1. Carmen González
2. Mario Van Poucke
3. Eva Ugarte
4. Fernando Rendo
5. Magdalena Serrano

**Abstract:**
While analysing the polymorphism of the ovine gene encoding the Hsp90α chaperone (HSP90AA1) in more than 600 animals pertaining to 23 different breeds, we found an HSP90AA1 pseudogene (HSP90AA1 Y) which was presented only in the 10% of the Spanish Latxa breed sheep analysed. Thus, based on the functional gene mRNA sequence, four primer pairs were designed in order to infer the HSP90AA1 Y complete sequence. Additionally, the Universal Fast Walking (UFW) strategy developed by Myrick (2005) was employed to elucidate the pseudogene flanking regions with the aim of inferring its location within the ovine genome. As a result, more than 2.5 Kb of the HSP90AA1 Y sequence has been identified. For the moment, just one variation in the nucleotide sequence has been found between the functional gene and the pseudogene. Furthermore, by comparing the
sequences obtained with the UFW strategy with different databases, preliminary results of the location of this pseudogene have been obtained. Thus, the existence of an HSP90AA1 pseudogene has been described for the first time on sheep as a recent event occurring exclusively within the Latxa breed. Nevertheless, further analysis should be performed in order to assess its biological significance.

**Poster 2091**

**Title:** The quantification of genetic components controlling observed variability to immunisation with a peptide derived from Foot-and-Mouth Disease Virus.

Presenting Author: R. J. Leach., The Roslin Institute, Royal (Dick) School of Veterinary Studies, University of Edinburgh, Roslin Bio centre, Midlothian, EH25 9PS, UK

Other authors (name only):
1. S. Craigmile
2. O. Matika
3. J. L. Williams
4. S. A. Knott
5. E. J. Glass

**Abstract:**

Identifying chromosomal regions which control immune responsiveness may suggest new approaches to disease control in livestock. A number of immune traits, including the response to immunisation with a foot-and-mouth disease virus (FMDV) peptide, have been measured in a Charolais-Holstein F2 and backcross cattle population ($N = 195$). The FMDV peptide comprises two FMDV derived VP1 capsid sequences, which represent the major neutralising antibody epitopes. Considerable variation was observed in the anti FMDV specific IgG1, IgG2 and T cell proliferative responses following immunisation. In order to identify potential genomic regions controlling these traits, a quantitative trait loci (QTL) mapping study was carried out using microsatellite markers covering the bovine genome. Thirty putative QTL were identified with more than half showing over-dominance (heterozygote > homozygote). The majority of the favourable alleles originated from the Charolais breed. This is the first evidence that the immune response to the FMDV peptide is under complex genetic control with multiple genes involved. The results may explain why the peptide only protects a proportion of cattle against viral challenge and why protection does not always correlate with high titres of neutralising antibody. Identifying genes underlying the QTL may help discover novel approaches to improve vaccine efficacy.

**Poster 2092**

**Title:** A Snapshot of Copy Number Variation in the Pig Genome

Presenting Author: João Fadista. University of Aarhus, Blichers Allé 20, P.O. Box 20, DK- 8830, Tjele, Denmark

Other authors (name only):
1. Christian Bendixen
2. Bo Thomsen
3. Lars-Erik Holm

**Abstract:**

Recent studies of mammalian genomes have uncovered the extent of copy number variation (CNV) that contributes to phenotypic diversity, including health and disease status. Here we report the first glimpse of CNVs in the pig genome covering part of the chromosomes 4, 7, 14 and 17 already sequenced and assembled. We used a Nimblegen custom tiling oligonucleotide array with a median probe spacing of 409 bp to screen 12 unrelated Duroc boars that are founders of a large family material. After a strict CNV calling pipeline 40 copy number variable regions (CNVRs) covering all the four chromosomes were identified, with some of the CNVRs overlapping segmental duplications and pig unigenes. This CNV snapshot analysis is the first of its kind and constitutes the groundwork for a better understanding of porcine phenotypes and genotypes with the prospect of identifying important economic traits.

**Poster 2093**

**Title:** A WHOLE-GENOME ASSOCIATION STUDY OF MAJOR DETERMINANTS FOR PARASITIC INFECTION IN ANGUS BREED.

Marcos Vinicius Silva, Bovine Functional Genomics Laboratory, Agricultural Research Service, USDA Beltsville, MD, 20705, USA

Other authors (name only):
1. Tad S. Sonstegard
2. Curt P. Van Tassell
3. Lakshmi K. Matukumalli
4. Steven G. Schroeder
5. Stewart W. Bauck
6. Lou C. Gasbarre

**Abstract:**

Fecal egg count (FEC) data recorded for 410 animals between 1992 and 2003 from an Angus herd were transformed using an extension of the Box-Cox transformation to approach normality. DNA for genetic analysis has been acquired for all animals from the resource population and over 70 sires in the
historic pedigree, in total of 595 animals. Genotyping was performed using Illumina’s BovineSNP50. For GWA analysis, SNP with a call rate (>93%), departure from HW equilibrium (exact test p<0.01), and minor allele frequency below 5 percent were excluded from the final analysis (26,158 markers retained). All the statistical analyses were carried out with scripts in the R environment and Fortran. Empirical p-values were corrected for genome-wide testing and maximization across genetic models, and a genome-wide significance level of 0.05 (two-sided) and Bonferroni corrected level of 0.01 were used. Five genetic models (codominant, dominant, recessive, overdominant and log-additive) were analyzed. The five most significant SNP were localized within a single linkage disequilibrium (LD) block. These findings provide the first evidence of biomarkers that contribute to early disease detection and primary prevention strategies for parasite infection in cattle and suggest new molecular targets for disease-modifying therapies.

Poster 2094
Title: Characterization of two bovine Y chromosome genes, DDX3Y and HSFY

Presenting Author: Wansheng Liu, Department of Dairy and Animal Science, College of Agricultural Sciences, The Pennsylvania State University, 305 Henning, University Park, PA, 16802, USA

Other authors (name only):
1. Aihua Wang
2. Yang Yang
3. Earl Landrito
4. Hiroshi Yasue

Abstract:
The DEAD box polypeptide 3, Y-linked gene (DDX3Y) encodes a putative ATP-dependent RNA helicase. This gene belongs to the DEAD box protein family, characterized by the conserved motif Asp-Glu-Ala-Asp (DEAD). The heat shock transcription factor, Y linked gene (HSFY) belongs to the HSF family, and is characterized by an HSF-type DNA-binding domain. Both DDX3Y and HSFY are candidates for azoospermia factor (AZF) and play an important role in spermatogenesis. Deletion of AZF region leads to infertility in otherwise healthy men. Here, we report the molecular characterization of the bovine (b) DDX3Y and HSFY genes. We found two transcripts for bDDX3Y, which correspond to the long and short transcripts of the human DDX3Y. The two bDDX3Y transcripts are identical except for a 3 bp insertion and an expanded 3’UTR in bDDX3Y-L. The bDDX3Y-S encodes a peptide of 660 amino acids (aa), while the bDDX3Y-L encodes a 661 aa. Both bDDX3Y isoforms contain the conserved motifs of DEAD-box RNA helicases. We identified six transcripts of the bHSFY gene from an adult testis, classified into two groups. Group I contains 3 transcripts with a different size 3’UTR that encode a peptide of 207 aa. Group II contains the remaining 3 transcripts that encode a 417 aa. One of the group II transcripts shows a deletion of 9 bp, resulting in an isoform of 414 aa. All bHSFY isoforms contain the conserved HSF DNA-binding domain. Our preliminary data indicated that multi-copies of the bHSFY gene are present on the bovine Y chromosome. The transcripts of bDDX3Y and bHSFY are predominantly or exclusively expressed in the bovine testis. This project was supported by grants from USDA-CSREES-NRI to WSL (No. 2005-35205-18653).

Poster 2095
Title: Genetic control of swine responses to porcine reproductive and respiratory syndrome virus infection

Joan Lunney, USDA, ARS, BARC, APDL, Building 1040, Room 103, BARC-East, Beltsville, MD 20705 USA

Other authors (name only):
1. Michal WYSOCKI
2. Daniel KUHAR
3. Juan P. STEIBEL
4. Catherine W. ERNST
5. Monte MCCAW
6. RRR ROWLAND

Abstract:
Our goal is use the Swine Protein-Annotated Oligonucleotide Microarray (www.pigoligoarray.org) to identify immune regulatory and protective pathways to uncover genetic components involved in early immune responses during porcine reproductive and respiratory syndrome virus (PRRSV) infection. Animals were divided into three groups: (1) pigs infected with a virulent PRRSV isolate MNW2B; (2) pigs vaccinated using a contemporary PRRS ATP vaccine (Ingelvac®); and (3) control pigs. Tissues [cranial lung, distal lung, tracheobronchial lymph nodes (TBLN)] were collected between days 3-6 post infection/vaccination. Total RNA was labeled using Alexa Fluor® 555 and 647 dyes (Invitrogen) and a microarray loop design applied to compare gene expression between individuals from all three groups. Validation of the pigoligoarrays was performed by analyzing transcriptional profiles for TBLN, Lung and longissimus dorsi muscle from normal or PRRSV infected pigs as confirmed using QPCR for candidate genes. Diagnostic analyses used designed array control features, negatives and perfect
match/mismatch oligonucleotide sets, to assess non-specific hybridization. Final statistical analyses of the PRRSV study are underway. These studies provide important support for our PRRS Host Genetic Consortium studies of neonatal pig resistance to infection that have just begun. Overall, these studies will reveal immune pathways and candidate genes involved in PRRSV resistance.

**Poster 2096**

**Title:** Arachnomelia in cattle: mode of inheritance and initial genetic mapping

**Presenting Author:** Dr. Johannes Buitkamp, Bavarian State Research Center for Agriculture, Institute of Animal Breeding 85580 Grub/Germany

**Other authors (name only):**
1. EMMERLING, R
2. LUNTZ; B
3. SEMMER, J
4. KÜHN, C
5. GÖTZ; K-U

**Abstract:**
Arachnomelia is a congenital disease of cattle. It is characterized by malformed bones of the limbs, back and head. Sporadically, additional findings, like cerebral herniation or hydrocephalus externus and internus, are observed. It causes losses, not only since the calves are not viable, but, more importantly, because sires that carry the mutation are banned from breeding and cows are hurt during delivery. A rapid rise of arachnomelia cases was observed in the years 2005 and 2006 in German and Austrian Simmental cattle. A number of sires with high genetic merit were recognized carriers of the mutation. Therefore, appropriate control measures had to be rapidly developed to avoid further spread of the allele in the population. In a first step breeding management measures were applied. To avoid the final exclusion of important sire lines from breeding, we started the genetic mapping of the disease with the final goal to develop an indirect gene test. Data about the pedigree analysis and population frequency estimation of arachnomelia as well as the genetic mapping are presented.

**Poster 2097**

**Title:** Quantitative trait loci for milk-fat composition in Dutch Holstein Friesians

**Presenting Author:** Anke Schenninkg, Animal Breeding and Genomics Centre, Wageningen University, Wageningen, the Netherlands

**Other authors (name only):**
1. W. Marianne Stoop
2. Marleen H.P.W. Visker
3. Patrick D. Koks
4. Erik Mullaart
5. Johan A.M. van Arendonk
6. Henk Bovenhuis

**Abstract:**
This study is part of the Dutch Milk Genomics Initiative and aims at the characterization of genes involved in milk-fat synthesis and milk-fat metabolism. Previous research has shown substantial genetic variation in milk-fat composition: heritabilities were high (0.42-0.71) for short- and medium-chain fatty acids (C4-C16) and moderate (0.22-0.42) for long-chain fatty acids (C18 and longer). To map QTL affecting milk-fat composition we conducted a whole-genome-scan. The mapping population consisted of 7 half-sib families containing 849 cows in a daughter design. A total of 1379 SNPs were typed covering all 29 autosomes. The phenotypes under study were 56 milk-fat composition traits, including saturated, mono-unsaturated, poly-unsaturated, and conjugated fatty acids, unsaturation indices and fat percentage. A regression interval mapping approach was used to estimate effects and positions of the QTL. QTL for short- and medium-chain fatty acids were detected on BTA6, 14, 19 and 26. QTL for long-chain fatty acids were detected on BTA14, 15, and 16. Our results will enable marker assisted differentiation and marker assisted selection, in order to optimize milk quality and to develop innovative dairy products. Fine mapping of the chromosomal locations is currently in progress.

**Poster 2098**

**Title:** Genomic signatures of artificial selection in U.S. Holstein cows

**Presenting Author:** Tad S. Sonstegard, USDA, ARS, Bovine Functional Genomics Laboratory 10330 Baltimore Ave. BARC-East Bdlg. 200 Rm 2A, Beltsville, MD U.S.A. 20705

**Other authors (name only):**
1. Li Ma
2. John B. Cole
3. George R. Wiggans
4. Curtis P. Van Tassell
5. George Liu
6. Brian D. Mariani
7. Brian A. Crooker
8. Paul M. VanRaden
9. Marcos V. da Silva
10. Yang Da
Abstract:
Selection in Holstein cattle has achieved tremendous phenotypic changes over the past 40 years. However, it is unknown how selection has changed the Holstein genome and how those genome signatures of selection are associated with the phenotypic changes. To categorize genome regions either affected or unaffected by selection, we contrasted genome structure data between 148 contemporary Holsteins and 151 cows from an unselected line of Holsteins bred and maintained at the University of Minnesota since 1964. Marker genotypes from 46,231 SNP were analyzed for allele frequency differences and trait associations using Wright’s Fst test and EPISNPmpi, respectively. Comparison of the most significant marker effects with Fst results revealed strong signatures of selection for production traits on chromosomes 1, 2, 3, 7, 8, 11, and 26. Chr 26 also had significant associations (16 markers; p<1.0E-12) with udder traits that co-localize with strong signatures of selection. Of the 31 SNPs with the most significant effects on both milk and daughter pregnancy rate, 29 had opposite effects on these traits providing strong evidence for the antagonistic effect of selection for production on fertility. These findings identify where selection has affected the genome and assist in searching for genes of large effect.

Poster 2100
Title: Global gene expression for cattle selection lines with low and high feed efficiency
Authors: Yizhou Chen1, Cedric Gondro2, Kim Quinn1, Robert Herd1
Cooperative Research Centre for Beef Genetic Technologies
1New South Wales Department of Primary Industries, Beef Industry Centre, Armidale, NSW 2351, Australia;
2The Institute for Genetics and Bioinformatics, University of New England, Armidale, NSW2351, Australia;
Email: yizhou.chen@dpi.nsw.gov.au

Abstract:
Feed efficiency is an economically important trait in beef cattle and it is affected by several physiological systems. The net feed efficiency measured as net feed intake (NFI) is the difference between the actual feed intake over a test period and its expected feed intake based on its size and growth. The objective of this project was to identify differentially expressed genes between animals with high and low NFI and pathways which contribute to the phenotype by global gene expression profiling using a 24K bovine long oligonucleotide array designed by the Bovine Oligonucleotide Microarray Consortium.

Liver tissue biopsies were taken from the top and bottom 30 tested bulls of Angus cattle divergently selected for low or high NFI and 44 animals were chosen for the microarray experiment. The microarray experiment was designed by ranking the animals based on NFI phenotypes and then pairing the top with the bottom animals using dye-swap. Standard QC measures were performed and arrays were individually normalized using print-tip loess normalization after background correction. Differentially expressed genes were ranked based on the p-values of a moderated t-statistic after fitting a linear model. The differentially expressed genes were assessed with regard to gene ontology, biological function and known genetic pathways.

Poster 2101
Title: Comparative performance and efficiency of five methods used for genome wide selection in dairy cattle using high density SNP data
Presenting Author: Gerhard Moser, Co-operative Research Centre for Innovative Dairy Products-CRC ID, Brisbane, Australia
Other authors (name only):
1. Bruce Tier
2. Mehar Khatkar
3. Ron Crump
4. Herman Raadsma

Abstract:
Three commonly used methods (Least squares regression-LS, Kernel regression-SVM, Bayesian regression- BayesA) and two new approach (Partial least squares-PLS, PLS with SNP selection-PLSjack) were compared in a Genome Wide Selection (GWS) analysis for prediction of genetic merit in dairy cattle. The data included 1945 progeny tested Holstein Friesian sires and genome wide analysis with 7372 SNP. The accuracy (r) of predicting EBV based on SNP genotypes (MBV) was studied by cross-validation. When test subsets were randomly drawn form the data (mirror prediction), accuracies for Australian Profit Rank (APR), an index of 9 component traits, ranged from r=0.69-0.80 and r=0.57-0.68 for protein%. To asses accuracies of MBV for young animals (forward prediction), the data was partitioned in a training set of 1239 bulls born before 1998, and 5 single year cohorts (1998-2002) of young bull teams. For forward prediction a loss in accuracies of 45% to 75% for APR and 15% to 40% for prot% was observed as compared to mirror prediction. In mirror and forward prediction LS, and Bayes A performed worse than SVM or either PLS method. For GWS, PLS with SNP selection performed the best and most consistent across the 5 year cohorts, and was computationally highly efficient.
**Poster 2102**

**Title:** Associations of Melanocortin 1 Receptor with growth and carcass traits in beef cattle.

Presenting Author: Kim L. McLean, Department of Animal & Poultry Science, University of Saskatchewan, 51 Campus Drive, Saskatoon, Saskatchewan, Canada

Other authors (name only): 1. Sheila M. Schmutz

**Abstract:**

Melanocortin 1 Receptor (MC1R) is considered to be the main gene controlling production of eumelanin and phaeomelanin resulting in black or red coat colour of cattle. The recessive red allele e, codes for a nonfunctional receptor which does not bind the agonist alpha-Melanocyte Stimulating Hormone (α-MSH), allowing for production of phaeomelanin or red pigment, whereas the dominant E allele binds α-MSH leading to the production of eumelanin. We hypothesized that black cattle would have more α-MSH bound to MC1R leaving less circulating α-MSH to bind to the appetite suppressing receptor, Melanocortin 4 Receptor. We genotyped 328 crossbred steers of various colours that were purchased at weaning and fed until slaughter. Red cattle (e/e) were found to have a larger rib eye area, shipping weight and hot carcass weight. Black cattle (E/D/E and E/D/e) had increased back fat and required significantly fewer days on feed to reach a target fat level for slaughter than the red cattle. By sorting cattle on a coat colour basis, 21 days on feed were eliminated for black cattle (P < 0.001). As many as 33 days on feed were avoided by genotyping and sorting cattle based on the number of E/D alleles (P= 0.001).

**Poster 2103**

**Title:** Identification of polymorphisms influencing production and functional traits on bovine chromosomes 19 and 29 in Canadian Holstein bulls

Presenting Author: Aparna Prasad, Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, Alberta, Canada, T6G 2P5


**Abstract:**

Detection of polymorphisms influencing economically important traits is a valuable tool for marker-assisted selection (MAS). The objective of the present study was to perform a chromosome-wide scan to detect polymorphisms associated with production and functional traits on bovine chromosomes 19 (BTA19) and 29 (BTA29) in Canadian Holstein bulls (n=322). In total, 505 and 220 single nucleotide polymorphism (SNP) markers on BTA19 and 29, respectively were genotyped. The average resolution of markers on BTA19 and 29 was 1 locus/172 Kb and 1 locus/228 Kb. Production traits studied were milk yield, fat yield, fat%, protein yield, and protein% and functional traits studied were herd life, daughter fertility, milking speed, milking temperament, calving ease and maternal caving ease. Single locus linkage disequilibrium regression model was used to test the association between SNPs and the different traits. In total, 133 SNPs were found to be significantly associated (P<0.01) with these traits. Forty six of them were located in the exon, one in the untranslated region, and three in the coding-synonymous region of the gene, while eighty three of them were not located within any genes. Once confirmed in an independent cattle population, these associations can be used in MAS schemes.

**Poster 2104**

**Title:** Multiplex assay for simultaneous identification of paternity and molecular markers of economic interest in bovines

Presenting Author: KATIA TORRES DE SOUZA, katia@linhagen.com.br


**Abstract:**

Following the world-wide trends, Brazil has invested in the development of more efficient mechanisms of lineage records and genetic improvement of bovines. In the last few decades, the lineage records of bovines have classified an individual animal and decided parentage questions only by using serological typing. This methodology presents limitations and recently has been replaced for DNA markers. LINHAGEN Biotecnologia has developed its own multiplex assay for bovine genetic identification, which includes, besides the 9 (nine) microsatellite markers recommended by ISAG (International Society of Animal Genetics), another 3 (three) extra markers which provides greater accuracy for the test itself.
Now we improved this assay even more by adding another markers to also analyse genetic characteristics of economic interest, such as quality and yield grade and/or the tendency for diseases, and use it for improvement genetic programs. The association of analysis between molecular markers and genetic identification in one test only is new in our country, will represent a significant advance and can provide: 1) bigger knowledge on the genetic characteristics of the national flock and dissemination of the use of molecular markers, hence that the paternity test tends to be mandatory and consequently, will be carried through on a large scale; 2) A large scale economy for the laboratories and breeders, because this assay can provide several analysis simultaneously.

Poster 2105

Title: Prion gene haplotypes of Thai indigenous cattle

Presenting Author: Kalaya BOONYANUWAT1 Chamnan DONGPALEE2 Winit KAMSUNG3

1Beef Cattle Research and Development Group, Division of Animal Husbandry, Department of Livestock Development. Rachathewi, Bangkok, 10400, Thailand. kalayabo@yahoo.com 2Phayao Livestock Research and Breeding Station. Phayao Province, Meung, 56000. Thailand. 3Thepa Livestock Research and Breeding Station. Songkla Province, Thepa, 90450. Thailand.

Abstract: Bovine spongiform encephalopathy (BSE) is a fatal neurological disorder characterized by abnormal deposits of a protease-resistant isoform of the prion protein. PRNP haplotype associated with BSE. The objective of this study was to identify polymorphisms and haplotypes of PRNP gene from octapeptides of the exon 3 region and promoter region through the 3'UTR in a sample of Thai indigenous cattle and compared with network haplotype to interplate BSE resistance and susceptibility. Four hundred and thirty-four individual DNA samples of Thai indigenous cattle were analysed for PRNP haplotype. Sequence analyses of octapeptide in exon 3 region identified 3 haplotypes of PRNP gene which had 1, 3, and 4 repeats of octapeptide. These polymorphisms define PRNP haplotypes that may influence BSE resistance in Thai indigenous cattle.

Poster 2106

Title: Genomic structure and genetic variants of leptin receptor (LEPR) gene in pig

Presenting Author: Kyung-Tai Lee / National Institute of Animal Science, RDA, Suwon, Korea.

Other authors (name only):
1. Hunsik Hwang
2. Kyung-Soo Kang
3. Eung-Woo Park
4. Jae-Hwan Kim
5. Jin-Tae Jeon
6. Seok-Ki Im
7. Tae-Hun Kim

Abstract: Leptin receptor (LEPR) gene is still considered as a positional candidate gene controlling QTL for fatness on chromosome 6 in pig due to its position and physiological role. Four BAC clones were screened by using LEPR-CA microsatellite marker and a segment of 324 kb was sequenced by shotgun strategy. We could define the genomic structure of LEPR gene in pig by comparing the genomic sequence of 324kb with LEPR transcripts found by RACE method. Porcine LEPR gene spanned 189,209bp and exon 1 and 2 were newly found and different from human and other species. Repetitive sequences were estimated to include 45.21% of the contig sequence, of which LINE and SINE were classified as 19.68 and 19.07%. The average G+C content and the number of CpG islands in the contig sequence were estimated to be 40.39% and 17, respectively. We investigated genetic variants such as SNPs within 5' upstream region including exon 1 of LEPR gene by using direct sequencing of 200 pigs from five. Association study for the genetic variants and methylation status on the CpG islands of regulatory region may produce a clue to understand relationship between the expression of LEPR gene and fatness in pig.

Poster 2107

Title: Sequencing-ready physical map of the horse Y chromosome

Presenting Author: …Terje Raudsepp, Department of Veterinary Integrated Biosciences, College of Veterinary Medicine, Texas A & M University, College Station TX 77843, USA

Other authors (name only):
1. Nandina Paria
2. Bhanu P. Chowdhary

Abstract: Horse Y chromosome (ECAY) project aims to identify all Y-linked genes and generate a detailed physical map for the chromosome. ECAY consists of a pseudoautosomal (PAR), male specific (MSY) and a heterochromatic (HC) region. We have constructed a BAC contig map spanning the entire PAR, MSY and a
part of HC. The map consists of 352 BAC clones which are arranged into 4 contigs by STS content analysis of 391 STSs and 52 genes/ESTs. The map demarcates the pseudoautosomal boundary and shows a segmental duplication between PAR and MSY. MSY is divided into single-copy and ampliconic regions. The latter contain multiple copies of known genes (TSPY, CUL4BY, UBE1Y) and several novel ESTs. Single-copy equine SRY is embedded in the middle of ampliconic sequences. Unique to ECAY is the discovery of ESTs in HC. The size of ECAY euchromatin is estimated to be ~13 Mb and the three gaps count for less than 2 Mb. The contig map of the ECAY euchromatin with 71 BACs forming the minimum tiling path is ready for sequencing. Complete information about the organization and function of ECAY is needed to understand the Y-linked component of stallion fertility and study sex chromosome evolution in mammals.

**Poster 2108**

**Title:** Fine mapping of QTL for left-sided displaced abomasum in German Holstein cattle

Presenting Author: Stefanie Mömke, Bünteweg 17P, 30559 Hannover

Other authors (name only):
1. Henner Scholz
2. Klaus Doll
3. Jürgen Rehage
4. Ottmar Distl

**Abstract:**

The displacement of the abomasum is a common disease in dairy cattle. Affected cows often have to be treated surgically and milk performance, as well as productive life are significantly reduced. For this study, samples of 360 animals belonging to 14 half-sib families were analysed. These 14 half-sib families comprised 328 LDA affected cows and 24 unaffected control cows. Five QTL on the bovine chromosomes (BTA) 1, 3, 21, 23, and 24 for left-sided displaced abomasum (LDA) in German Holstein cattle were fine-mapped, using a total of 92 already published or newly developed microsatellite and SNP markers. The peaks of these QTL were located at the markers MNB152 (BTA1, 54.6 cM, Zmean: 3.18), DIK4922 (BTA3, 5.9 cM, Zmean: 3.04), BMS743 (BTA21, 75.3 cM, Zmean: 1.96), IOBT528 (BTA23, 7.9 cM, Zmean: 2.84), and DIK4984 (BTA24, 75.2 cM, Zmean: 1.94), respectively. The QTL on BTA1 and 3 reach genome-wide significance, while the QTL on BTA21, 23, and 24 are chromosome-wide significantly linked with LDA.

Further fine-mapping will be provided by a SNP chip analysis with following association studies.

**Poster 2109**

**Title:** Phylogenetic analysis and genetic variation of population of genus Lepus in Europe.

**Authors:**
Presenting Author: María J. Sanz-Martín. Genetics, Physical Anthropology and Animal Physiology Department. Faculty of Science and Technology. University of the Basque Country.

Other authors (name only):
1. Mikel Iriondo
2. Gonzalo Pérez-Suárez
3. Carmen Manzano
4. Andone Estonba
5. Fernando Palacios

**Abstract:**

Three out of five hare species of genus Lepus in Europe coexist in the Iberian Peninsula, L. europaeus, L. granatensis and L. castroviejoi, and their genetic relationship is uncertain. This work assesses two aims: 1) The phylogenetic relationship of the Iberian hare in a European context 2) The development of an exhaustive genetic map of the Iberian L. europaeus with conservation purposes. To achieve these aims, 295 individuals from all five European species were analysed. A fragment of 400 bp from mtDNA’s control region was sequenced and 17 microsatellite loci were genotyped. Results prove that L. granatensis is more closely related to the European lineage than to the African-Asian one. On the other hand, previous mtDNA studies have shown that the autochthonous populations of Iberian L. europaeus, contrary to the rest of L. europaeus in Europe, are characterized by an ancient introgression by L. timidus. Our results confirm this fact, although we also observed non-introgressed individuals in the northern Iberian Peninsula, likely due to non-controlled restocking from other European regions.

**Poster 2110**

**Title:** DGAT1: a diet-regulated quantitative trait locus that affects hepatic lipid deposition

Presenting Author: Rosane Oliveira, Department of Animal Sciences, University of Illinois, 1201 W Gregory Dr, Urbana IL, 61821

Other authors (name only):
1. Juan J. Loor
2. Heather M. Dann
3. Nicole A. Janovick-Guretzky
4. James K. Drackley
5. Harris A. Lewin

**Abstract:**
A lysine to alanine substitution at codon 232 in the Acyl-CoA:diacylglycerol acyltransferase1 gene (DGAT1 K232A) underpins a quantitative trait locus (QTL) responsible for differences in milk and fat yields in dairy cows. In the present study, dietary energy intake and DGAT1 genotype were evaluated for their effects on transcript levels of hepatic DGAT1 and accumulation of triacylglycerol (TAG) and total lipid in liver. Thirty-five cows fed energy *ad libitum* or restrictively during the prepartum period were genotyped for the DGAT1 K232A mutation. DGAT1 mRNA levels, TAG, and total lipid were determined in biopsied liver samples collected at -14, +1 and +14 days relative to parturition. TAG and lipid concentrations were significantly higher at day +14 in *ad libitum*-fed cows having one or two copies of the DGAT1K allele. DGAT1 genotype had a significant effect on DGAT1 mRNA levels, and an effect of diet on DGAT1 mRNA levels was found at one day postpartum. DGAT1 alleles appear to be responsible for liver TAG and lipid accumulation postpartum in animals overfed during the prepartum period. Nutritional management of cows having the DGAT1K allele can thus be used to reduce economic losses due to metabolic disorders occurring as a result of overfeeding energy prepartum.

**Poster 2111**

**Title:** Transcriptional profiling and biochemical pathways analysis of liver from two breeds of cattle.

Presenting Author: Pawel Lisowski, Department of Molecular Biology, Institute of Genetics and Animal Breeding, Polish Academy of Sciences (IGAB PAS), 05-552 Jastrzebiec, Poland

Other authors (name only):
1. Joanna Goscik
2. Pawel Brym
3. Lech Zwierzchowski

**Abstract:**
We investigated the gene expression profile in the liver of cattle breeds distinctly different in meat and milk production capacity, carcass and meat quality. Studies are based on the assumption that animals differing in meat and milk performance must show different expression profiles of genes that determine these traits. The study was performed in liver tissues from bulls with two different genetic backgrounds, meat and dairy - Hereford and Holstein-Friesian. The animals were at 6 months of age during the intensive growth and muscle development. The 8-K bovine oligo microarrays were used (BLO, Michigan State University). Microarray analyses identified 34 genes of which the expression levels differed at least 2-fold. These genes are involved in 97 biological processes. Hereford animals showed higher expression of L-FAB, GH receptor precursor, STAT1, and STAT3 genes, while in HF animals had higher expression of PKC and ATP synthase gamma chain genes. Combining the microarray results and biological databases information we choose the genes engaged in 24 biochemical pathways for further analysis. These pathways were related to cellular processes, energy metabolism and signal transduction. Our findings provide a description of molecular events accompanying individual development that may underlay the phenotype differences between two cattle breeds.

**Poster 2112**

**Title:** Genetic analysis of local Vietnamese chickens for conservation purposes

Presenting Author: Berthouly Cécile-UMR 1236, AgroParisTech 16 rue Claude Bernard, 75005

Conference registration number of Presenting Author: 0401

Other authors (name only):
1. Nhu Van Thu
2. Hoang Thanh Hai
3. Bertrand Bed’Hom
4. Vu Chi Cuong
5. Etienne Verrier
6. Jean-Charles Maillard
7. Xavier Rognon

**Abstract:**
The Vietnamese H’Mong chicken breed called the Black Chicken because of the black bones, skin and shank, had a strong cultural value. In order to characterise the H’mong chicken from the originating province, namely Ha-Giang, phenotypic and genetic analyses were performed. Thirty communes from the 11 districts of the province were surveyed. A total of 2487 animals were described for 29 phenotypic characters and analysed by Principal Component Analysis (PCA). The two principal components explained 53.6 % of the inertia. All variables describing black colour grouped together and allowed differentiation of three communes among the thirty. From the described animals set, 1082 animals were genotyped using 18 ISAG microsatellite markers. In the Ha-Giang population, $H_{EXP}$ averaged 0.619 and $F_{IS}$ averaged 0.121 and 3.7 % of the genetic diversity was due to differentiation between communes. Bayesian approach did not showed genetic subdivision into subpopulations. In summary, analysis of phenotypes revealed a geographic distribution of Black chickens while the bayesian admixture analysis within the
province did not show genetic differentiation of the Black chicken within the province. Therefore, a reproductive pool of chicken considering only the black phenotype could be used if government wants to create a genetically distinct H’mong breed.

**Poster 2113**

**Title:** Individual assignments to dog breeds using simulated and real data

Presenting Author: Grégoire Leroy
INRA/AgroParisTech Génétique et Diversité Animales
Other authors (name only):
1. Jean-Claude Mériaux
2. Etienne Verrier
3. Xavier Rognon

**Abstract:**
Dog breeds constitute clearly genetically differentiated populations. Thus, breed assignment may represent an interesting tool for clubs and breeders in order to investigate if a dog without genealogy can be assigned to a particular breed. Several Bayesian approaches have been developed during last years, using direct assignment tests, or simulating populations to exclude animals from the populations tested. In a first part, we tested the different approaches on dog breeds by using 21 microsatellite markers. The assignment of 1240 individuals belonging to 63 breeds was tested on a reference set of 2000 dogs of 9 breeds. Using simulation procedures, results showed that all individuals belonging to the same breeds of the reference set were correctly assigned. However, 14% of the animals of other breeds were incorrectly assigned in the 9 breeds at 1% threshold. In a second part, populations were simulated with a genetic structure more or less similar to dog breeds. Our results showed that, as expected, allelic richness constitutes a parameter that influences assignment effectiveness. However heterozygosity (H) by itself has some influence on the results: breeds with a high H level affect incorrectly a larger number of animals when population simulation procedures are used. Effects of other parameters, such as sample size were also tested.

**Poster 2114**

**Title:** Molecular phylogenetics of mtDNA D-loop sequences in Chinese buffalo: implication for origins of domestic buffalo (*Bubalus bubalis*)

Presenting Author: Yi Zhang, Department of Animal Genetics & Breeding, College of Animal Science and Technology, China Agricultural University, Beijing, China, 100094
Other authors (name only):
1. Dongxia Sun
2. Ying Yu
3. Yuan Zhang

**Abstract:**
In this study, to characterize the genetic origin of Chinese swamp buffalo and their phylogenetic relationships with river buffalo and Southeast Asian swamp buffalo, we determined the whole mtDNA D-loop sequences of 100 domestic buffalo geographically covering their main distribution in China and analyzed these sequences in comparison with published data. Neighbor-joining phylogenetic tree revealed two highly divergent mtDNA lineages (A and B) in Chinese swamp buffalo, in which A was predominant (77%) and B could further subdivided into B1 (11%) and B2 (12%). These lineage groups all were demonstrated to have undergone recent population expansion. The divergence estimation between A and B, based on cytochrome b gene sequences, is approximately 650,000 years ago. Moreover, both A and B were found in all Chinese swamp populations with similar proportions (proportion of B = 0.16±0.07). These findings suggest that Chinese buffalo might be derived from a single wild source population. In addition, a combined media-joining network analysis provided evidence that swamp and river buffalo originated from two independent domestication events in China and Indian subcontinent, respectively. The buffalo in South-east Asia, however, have both ancestries, indicating possible genetic mixture of two buffalo types in this region.

**Poster 2115**

**Title:** Investigation of the KIT gene in Nero Siciliano pigs with different coat colour phenotypes

Presenting Author: Enrico D’Alessandro,
Dipartimento di Morfologia, Biochimica, Fisiologia e Produzioni Animali – Sezione di Zootecnica e Nutrizione Animale, University of Messina, 98168 Messina, Italy
Other authors (name only):
1. Luca Fontanesi
2. Emilio Scotti
3. Luigi Liotta
4. Vincenzo Chiofalo
5. Vincenzo Russo

**Abstract:**
The Dominant White locus in pigs is caused by mutations in the *KIT* gene that have been described as
copy number variation and single nucleotide polymorphisms. Nero Siciliano (Black Sicilian) is a local Italian pig breed reared in the Nebrodi mountains of Sicily. The animals are usually completely black but a few show white portions mainly in the face or in the fore legs. We genotyped 97 Nero Siciliano pigs (25 with white portions) for the splice site mutation of the intron 17 of the \textit{KIT} gene that is associated with the Dominant White \textit{I} allele. These animals were also analysed for the presence of the duplication of the same gene. None of the pigs carried the splice site mutation while 7 out of 25 animals with white portions were positive for the duplication test. Then, 20 of the 21 exons of the \textit{KIT} gene (including intronic regions) were sequenced in eight Nero Siciliano pigs with different colour patterns and in one Large White animal. Several new mutations were identified in Nero Siciliano pigs. The results suggest that the presence of white portions in Nero Siciliano pigs might be only in part due to mutations in the \textit{KIT} gene.

**Poster 2116**

**Title:** Single Nucleotide Polymorphism (SNP) density and predicted mapping resolution of complex trait genes in the Thoroughbred horse

Presenting Author: S.C. Blott, Animal Health Trust, Lanwades Park, Kentford, Newmarket, Suffolk, CB8 7UU, UK

Other authors (name only):
1. L.Y. Fox-Clipsham
2. E. Temperton
3. J.E. Swinburne
4. M. Vaudin

**Abstract:**
The recent sequencing of the equine genome and associated single nucleotide polymorphism (SNP) discovery makes genome-wide association studies possible in horses. The required marker density for an effective whole genome scan depends on the extent of linkage disequilibrium across the genome. We have re-sequenced 52 SNPs across a 1 Mb region on ECA 26 in order to assess the extent of fine-scale linkage disequilibrium in the Thoroughbred horse. The median spacing between SNPs is 200 bp, maximum 80 kb and minimum 1 bp. The 52 SNPs have been re-sequenced in 24 unrelated Thoroughbreds and pairwise estimates of $D'$ and $r^2$ between SNPs have been calculated. This allows the extent of linkage disequilibrium for genetic distances between 1 bp to 1 Mb to be assessed. Haplotype phases have been determined for each individual horse and the size of haplotype blocks and number of tagging SNPs per block evaluated. The precision and power of mapping, given the observed patterns of linkage disequilibrium, has been explored using simulation.

The information is useful in determining the SNP density required for whole genome association scans and for follow-up fine mapping in the Thoroughbred.

**Poster 2117**

**Title:** MUTATIONS IN TWO GENES ARE ASSOCIATED WITH INCREASED OVULATION RATE AND LITTER SIZE IN PROGENY OF LACAUNE MEAT SHEEP

Presenting Author: Drouilhet L. INRA Laboratoire de Génétique Cellulaire, 31326 Castanet-Tolosan, France

Other authors (name only):
1. Lercerf F
2. Bodin L
3. Mulsant P

**Abstract:**
A new autosomal major gene affecting ovulation has been evidenced by a statistical approach in the Lacaune population. A full genome scan localized this autosomal prolificacy gene (\textit{FecL}) on sheep chromosome 11. Further investigations allowed the reduction of the localization interval between markers BM17132 and FAM117A, corresponding to one syntenic block of human chromosome 17 of 1.1 megabases, containing 20 genes. This fine mapping is still going on. The expression of those 20 genes was tested by qPCR and no difference was found in homozygous mutants versus wild type, in four tissues (granulosa and theca cells, hypothalamus and pituitary gland). A particular haplotype was associated with the \textit{FecL} mutation, which allowed marker associated classification of animals as carriers or non-carriers of the mutation. In addition, ewes with extreme ovulation rate have been isolated in the Lacaune population and it was hypothesized that another mutation was segregating in the population. A new mutation has been evidenced in the \textit{Bmp15} coding sequence (chromosome X). This mutation is a non conservative substitution (C53Y) that prevents the processing of the protein. Synergic action of the two mutations (in \textit{FecL} and \textit{Bmp15} genes) on both ovulation rate and litter size was demonstrated.

**Poster 2118**

**Title:** Mitochondrial DNA sequence variation in three Sicilian autochthonous horse breeds

Presenting Author: DONATA MARLETTA, DACPA - Sezione di Scienze delle Produzioni Animali. Università degli studi di Catania. Via Valdisavoia, 5 – 95123 Catania. Italy

Other authors (name only):
Matrilineal genetic diversity in Sicilian horses was investigated sequencing the mitochondrial D-loop hypervariable region (397 bp) in Sanfratellano, Sicilian Indigenous and in Sicilian Oriental Purebred, 20 mares per breed. Twenty different haplotypes, with 30 polymorphic sites, were identified: 11 in Sanfratellano, 13 in Sicilian Indigenous, 1 in Sicilian Oriental Purebred.

Genetic distances between haplotypes were estimated using Kimura two-parameters (MEGA 4); phylogenetic tree was constructed using neighbour-joining method. Nucleotide diversity (θ), estimated by Arlequin 3.1, showed that Sicilian Indigenous was the most diverse population (θ = 0.018 ± 0.009) followed by Sanfratellano (0.916 ± 0.041); haplotype diversity stated the same trend. BLAST search showed that all the sequences overlapped with many of the GeneBank database haplotypes, but for 3.

Phylogenetic analysis did not show monophyletic groups for the Sicilian breeds, confirming the multiple origins for the maternal lineages of the domestic horse breeds. The Sicilian Oriental Purebred haplotype, found in 5 Sanfratellano mares, denotes a mixture of these two genetic types. Some haplotypes, identified in Sanfratellano and Sicilian Indigenous, suggest a common origin between these two breeds; furthermore three haplotypes, not shared with other breeds, indicate a possible geographic segregation of these two autochthonous horses.

Supported by ISZS project 14, 03/05/2007

Poster 2119

Title Microarray analysis of skeletal muscle genes in pigs divergent for glycolytic potential.

Presenting Author: Roberta Davoli, Dipartimento di Protezione e Valorizzazione Agroalimentare (DIPROVAL), Sezione di Allevamenti Zootecnici, University of Bologna, 42100 Reggio Emilia, Italy

Other authors (name only):
1. Paolo Zambonelli
2. Silvia Braglia
3. Marco Comella
4. Luca Fontanesi
5. Alessandra Stella
6. Michela Colombo
7. Luca Buttazzoni
8. Vincenzo Russo

Abstract:
Glycolytic potential (GP) is an indicator of the glyogen level in skeletal muscle and this parameter may have a marked effect on technological pork quality. The aim of this research was to identify pig genes involved in the regulation of GP in muscle by microarray analysis. Samples of semimembranosus muscle from eight pigs with extreme and divergent GP were selected among 277 sib-tested Italian Large White (ILW) animals and utilised for RNA extraction. A RNA pool with low GP (4 samples presenting GP<65 μmol lactate equivalent g⁻¹ muscle wet weight) and another RNA pool with high GP (4 samples with GP>145 μmol) were obtained. The Cy3 and Cy5 labelled cDNAs were used to hybridise the 11K Pig Oligo set version 1.0 (Operon-Qiagen). Differentially expressed genes were identified by ANOVA using GLM procedure in SAS. The analyses showed that several genes up-regulated in the pool with low GP were involved in energy metabolism and only few genes coding for transcription factors, glycolytic pathway and cell metabolism were detected in the pool with high GP. Differential gene expression detected by microarray was confirmed for selected genes by quantitative real time on the individual RNA samples of the two pools.

Poster 2120

Title: New mutations in BMP15 and GDF9 genes are not associated with increased prolificacy in Rasa aragonesa sheep breed.

Presenting Author: Albert Martinez-Royo & Unidad de Tecnología en Producción Animal, CITAd, Avda de Montahana 930, 50059 Zaragoza. Spain

Other authors (name only):
1. Laurence Drouilhet
2. Loys Bodin
3. Philippe Mulsant
4. José Folch
5. Jose Luis Alabart
6. Juan José Jurado
7. Jorge H. Calvo

Abstract:
Recently, a new naturally occurring mutation in BMP15 gene in the ovine Rasa aragonesa breed has been described (FecX<sup>+</sup> allele). It has been associated with both increased prolificacy and sterility in heterozygous and homozygous ewes, respectively. Animals from controlled nucleus with high EBVs for prolificacy without the FecX<sup>+</sup> allele have been analysed for BMP15 and GDF9 genes by SSCP technique. Both genes are members of the transforming growth factor beta (TGFβ) superfamily
and play an essential role in mammalian fertility. Animals with different profiles were sequenced and the mutations were studied. A total number of 3 and 4 mutations have been found in \textit{BMP15} and \textit{GDF9}, respectively. The first of these is a leucine deletion in the predicted signal sequence of \textit{BMP15} without any phenotypic effect. Four of the seven polymorphisms are nucleotide changes that do not result in an altered amino acid. The two remaining nucleotide changes occurs in \textit{GDF9} and produce an amino acid change. Both occurs at a position before the furin cleavage site for the mature peptide, so are unlikely to affect the activity of the mature protein.

Poster 2121

Title: A genome scan for QTL affecting milk somatic cell count in the Italian Holstein Friesian cattle breed by selective milk DNA pooling in a daughter design

Presenting Author: Luca Fontanesi, DIPROVAL, Sezione di Allevamenti Zootecnici, University of Bologna, Via F.lli Rosselli 107, 42100 Reggio Emilia, Italy


Abstract:
In dairy cattle, mastitis is the most important health problem causing substantial economic losses. Milk somatic cell count (SCC) is highly correlated with mastitis and hence is used as a surrogate trait for breeding objective. A selective milk DNA pooling strategy in a daughter design was used in the Italian Holstein-Friesian cattle breed to map QTL for milk SCC as a first step towards the application of MAS for this trait. Milk DNA pools from about 200 daughters with high and 200 with low estimated breeding value (EBV) for SCC were constructed for each of six sires with at least 3500 milking daughters each. Sires were genotyped for a panel of 133 dinucleotide microsatellites distributed over all bovine autosomes. DNA pools were genotyped for the markers that were heterozygotes in the sires. Shadow corrected estimates of sire allele frequencies in the pools were compared between high and low pools. Proportion of false positives (PFP) was applied to obtain experimentwise significant levels. Allele substitution effects were calculated for all significant sire-marker trait combinations. Several QTL were identified and their location was compared to the QTL already identified in the same population for milk yield and milk protein percentage.

Poster 2122

Title: Effects of 52 SNPs on growth rate, meat content and selection index in Large White and Landrace boars

Presenting Author: Stanislaw Kaminski, University of Warmia and Mazury, 10-718 Olsztyn, Poland, email: stachel@uwm.edu.pl.

Other authors (name only): 1. Joanna Szyda 2. Hela Help 3. Tomasz Suchocki

Abstract:
306 boars (108 Large White and 198 Landrace) were genotyped for 52 SNPs to find out which of them or their combinations influence growth rate (GR), meat content (MC) and selection index (SI). The effects of SNPs were estimated by using a mixed linear model including random animal effect, fixed effects of SNPs including additive, dominance and pairwise additive-by-additive epistases, year*season of birth, breed and RYR1 genotype. In order to estimate all possible SNP combinations without overparameterising a model a stochastic approach was adopted. 1900 repeats of the model were generated, each containing 5 randomly chosen SNPs. The final estimates of the fixed effects of the model equaled an average out of the replications. The hypothesis of a nonzero effect of SNP was tested by the t test. Among 4300 estimates calculated, many significant (p<0.01) but minor effects (below 1% of a trait mean) were recorded. Outstanding effects (higher than 1 SD) were found in 17, 17 and 10 SNP pairs, for GR, MC and SI, respectively. Loci most frequently involved in epistasis were: H-FABP, MEF2D, LDLRRP1, LDHA, ESR2, GAD2, PRKAG3. The selected SNPs will be further investigated to find out which ones may be applied in MAS.

Poster 2123

Title: Copy number variation and missense mutations in the caprine agouti signaling protein (\textit{ASIP}) gene are present in goat breeds with different coat colour

Presenting Author: Stefania Dall’Olio, DIPROVAL, Sezione di Allevamenti Zootecnici, University of Bologna, Via F.lli Rosselli, 107, 42100 Reggio Emilia, Italy
Abstract:
The Agouti locus encodes the agouti signaling protein (ASIP) that plays a key role in the pigment-type switching from eumelanins to phaeomelanins. In goats, the effect of the Agouti locus on coat colour and colour pattern distribution has been deduced analysing herd book records and segregation data in classical crossbreeding experiments. Here, we sequenced the coding region of the goat ASIP gene in six breeds (Camosciata, Derivata di Siria, Girgentana, Maltese, Murciano-Granadina and Saanen) showing different coat colours and patterns. Five single nucleotide polymorphisms were identified, three of which were missense mutations within exon 4, causing amino acid substitutions in conserved positions of the carboxy-terminal domain of the protein. Allele and genotype frequencies suggested that the missense mutations do not affect coat colour in the investigated goat breeds. Genotyping results and deviation from Hardy-Weinberg equilibrium, as well as allele copy number estimation from sequencing data indicated the presence of copy number variation (CNV) for this gene with high frequency in the Girgentana and Saanen breeds. The obtained results, together with the data already reported by others for the ovine ASIP gene, provide evidence for a recurrent interspecies CNV at this locus.

Poster 2124

Title: Determination of protein and the RNA expression levels of the commonly used housekeeping genes in a neurodegeneration mouse model

Presenting Author: Ana C. Calvo, LAGENBIO, University of Zaragoza. C/ Miguel Servet 177, 50013 Zaragoza, Spain

Other authors (name only):
1. María Moreno-Igoa
2. Raquel Manzano
3. Laura Ordovás
4. Gema Yagüe
5. Sara Olíván
6. María J. Muñoz
7. Pilar Zaragoza
8. Rosario Osta

Abstract:
The choice of housekeeping proteins or genes for internal standards should be made carefully, taking into account the cell and tissue type, the experimental conditions, and the disease state(s) under consideration. GAPDH, β-actin, and β-tubulin are generally considered housekeeping genes and are commonly used as internal standards in a broad range of studies examining levels of gene transcription. We are interested in studying changes in protein and gene expression that may shed light on a neurodegenerative disease mouse model, SOD1<sup>G93A</sup> mice, that bear a G93A mutation in the human gene for superoxide dismutase 1 (SOD1) and therefore results in motor neuron pathologies. As such, we first wanted to assess whether β-actin, GAPDH and β-tubulin could be used as internal protein controls in spinal cord, brain, and skeletal muscle tissues. We next examined the variability of β-actin and GAPDH transcriptional levels, in the same experimental conditions. The results suggest the possibility that alterations in the integrity of microtubules may be early events of the neurodegenerative processes that may finally reflect later alterations in GAPDH metabolism, in terms of altered protein levels that potentially occur in skeletal muscle, one of the most affected tissues.

Poster 2125

Title: Polymorphism and Genomic Organization of the Equine MHC Class II Region

Presenting Author: Donald Miller, Baker Institute, Cornell University, Ithaca, NY, 14853 USA

Other authors (name only):
1. Douglas F. Antczak.
A9, and -A10 haplotypes. Our findings reveal a higher level of polymorphism in the beta genes compared to the alpha genes, and three of the five beta gene loci studied have unique alleles for each of the five ELA haplotypes.

**Poster 2126**

**Title:** Parentage testing in alpacas (*Vicugna pacos*) using semi-automated fluorescent multiplex PCRs with 10 microsatellite markers

Presenting Author: Jose R. Espinoza. Laboratories for Research & Development (LID), Universidad Peruana Cayetano Heredia, Av. Honorio Delgado 430, Lima 31, Peru.

Other authors (name only):
1. Juan Agapito
2. Jorge Rodriguez
3. Teresa Barreto
4. P.J. Boettcher
5. Fernando Garcia

**Abstract:**
A microsatellite multiplex system for parentage determination in alpacas was evaluated in a population of 329 unrelated alpacas from different geographical zones in Peru. All microsatellite markers (n=10), which amplified in two multiplex reactions, were highly polymorphic with a mean of 14.5 alleles per locus (six to 28 alleles per locus) and an average expected heterozygosity (HE) of 0.8185 (range of 0.698–0.946). The total parentage exclusion probability was 0.999456 for excluding a candidate parent from parentage of an arbitrary offspring, given only the genotype of the offspring, and 0.999991 for excluding a candidate parent from parentage of an arbitrary offspring, given the genotype of the offspring and the other parent. In a case test of parentage assignment, the microsatellite panel assigned 105 (from 105 cases) offspring parentage to 15 sires with LOD scores ranging from $2.19 \cdot 10^{13}$ to $1.34 \cdot 10^{15}$ and $\Delta$ values ranging from $2.80 \cdot 10^{12}$ to $1.34 \cdot 10^{15}$ with an estimated pedigree error rate of 13.2%. The performance of this multiplex panel of markers suggests that it will be useful in parentage testing of alpacas.

**Poster 2127**

**Title:** QTL for maternal ability traits in an Iberian x Meishan pig cross.

Presenting Author: Cristina Óvilo, Dpto Mejora Genética Animal, INIA, Ctra Coruña Km 7.5, 28040 Madrid, Spain

Other authors (name only):
1. Jose Luis Noguera
2. Anna Tomás
3. Gloria Muñoz
4. Óscar Ramírez
5. Amandá Fernández
6. María Martínez
7. Meritxell Arqué
8. Carmen Barragán
9. Armand Sánchez
10. Carmen Rodríguez

**Abstract:**
The aim of this study was to investigate chromosomal regions affecting maternal ability traits: litter weight at farrowing (LWF) and at weaning (LWW). For this purpose an F₂ experimental cross between Iberian and Meishan porcine breeds was used. The genomic scan was carried out using 134 markers (microsatellites and SNPs in several genes) covering the 18 porcine autosomes, with 670 and 615 records for LWF and LWW, respectively. Analyses were performed using a repeatability animal model to estimate QTL and additive polygenic effects. For LWF the results showed a genome-wide highly significant QTL ($p<0.1\%$) located on SSC12 (45cM), not reported previously. This QTL present both additive ($0.49\pm0.14$) and dominant ($0.56\pm0.20$) effects, and coincides with the position of growth hormone gene (*GH*). A search of mutations was conducted on this gene for its evaluation as candidate for the QTL. Three missense polymorphisms were genotyped, resulting on three haplotypes segregating in the F₂ population. Results of a marker assisted association analysis showed significant association of *GH* polymorphisms, with a favorable effect of the Iberian allele on LWF ($p=0.13\times10^{-2}$). For LWW a genome-wide significant QTL ($p<0.1\%$) was detected on SSC6, at 78 cM. This QTL shows additive effect ($2.02\pm0.49$).

**Poster 2128**

**Title:** Construction of chicken 50K SNP panel and its application in broiler breeding

Presenting Author: Tun-Ping Yu, DNA LandMarks Inc., St.-Jean-sur-Richelieu, Québec, Canada

Other authors (name only):
1. Ron Okimoto
2. Yinyan Guo
3. Guy Champagne
4. Wing Cheung
5. Albert Paszek
6. Charles Pick
7. Joachim Richert
8. John Hardiman
Abstract:
Cobb-Vantress, Inc. (Cobb) is the world leader in developing broiler breeding stock. Through collaboration with DNA LandMarks (DLM), the Center of Excellence for Genomics of BASF Plant Sciences, we have established a large scale chicken SNP panel for genomic research that enhances genetic progress in economically important traits. With custom-developed bioinformatic tools, a panel of over 60,000 SNPs was identified. Approximately 50,000 SNPs were converted to an Affymetrix custom panel. The concordance rate of SNP genotyping was over 99%. Among the 50K SNPs, 95% were confirmed to be true SNPs and showed segregation in Cobb lines. A highly informative SNP panel evenly covering the chicken genome for each of the Cobb pedigree lines was obtained through a stringent SNP filtering system according to scores of marker informativeness. Linkage Disequilibrium (LD) analysis was performed using Composite Linkage Disequilibrium analysis to gain better understanding of the genetic structure of Cobb pedigree lines. LD results provided key information for experimental design for a whole genome association analysis. Pedigree-based haplotype analysis and decremental search function were used to identify tagged-SNPs within the LD regions. The pedigree line-specific SNP panels are used to study production and animal welfare traits for application in marker assisted selection.

Poster 2129
Title: Gene expression-phenotype associations using eQTL based gene sets.
Presenting Author: L. Janss, Aarhus University, Denmark.
Other authors (name only):
1. V. Obolonkin

Abstract:
In gene expression analysis, gene sets can be used to improve power, biological underpinning and consistency between experiments. Commonly, gene sets are based on bioinformatics data such as pathways or the Gene Ontology. In this study we present a gene expression analysis using gene sets based on eQTL locations for the gene expressions. This can be seen as a “data driven” gene set assignment which can fill gaps in existing knowledge about co-regulated genes. For this analysis, firstly, individual gene expressions were tested for association with the phenotype and a loose selection of primary candidates was made. Subsequently for all these candidates, an eQTL mapping was performed using a Bayesian multi-QTL mapping procedure and the selected expressions were grouped in gene sets according to common eQTL position. In the final stage the so determined gene sets were tested again for association with the phenotype. This procedure identified 3 main gene sets each explaining ~8% of variation, and a handful of gene sets each explaining ~4-6% of variation in the phenotype. Being based on common eQTL position, a genomic location for the putative “master regulator gene” for each gene set is available as well.

Poster 2130
Title: CFA9 association study to identify potential modifier loci for progressive rod-cone degeneration (prcd) in the dog
Presenting Author: Barbara Zangerl, University of Pennsylvania, School of Veterinary Medicine, 3900 Delancey Street, Philadelphia, PA, 19104
Other authors (name only):
1. S.J. Lindauer
2. S.E. Pearce-Kelling
3. G.M. Acland
4. G.D. Aguirre

Abstract:
A single mutation in the PRCD gene, causing progressive rod-cone degeneration (prcd) in the dog, was found to be shared among 27 breeds. The disorder causes retinal degeneration, always resulting in complete blindness; the time of onset and disease progression, although generally breed specific, varies between breeds. Previous breeding experiments established the stable segregation of these variants linked to the original disease. Thus, a modifier likely is segregating in genomic proximity to the prcd locus on CFA9. 11 and 13 animals affected by the fast and slow phenotype, respectively, were compared against each other on the canine Affymetrix SNP chip with respect to a control group comprised of 10 known carriers. An initial 1,807 CFA9 markers in 29 individuals passed minimal quality requirements. PLINK, and t-tests for selected markers, placed the most likely signal for prcd between SNP 6,837,703 and 7,230,119, indeed including the mutation (7,186,710). Surprisingly, additional areas of interest are interspersed throughout CFA9. The strongest signal, 36 Mb from the mutation, suggests involvement of the neuronal amiloride-sensitive cation channel 1 (ACCN1) with disease phenotype, which is currently being investigated. Although preliminary, these data may provide first indicators for a potential prcd modifier.
**Poster 2131**

**Title:** A microarray–based approach for the identification of skeletal muscle genes related to stress in pigs.

Presenting Author: Zambonelli Paolo, DIPROVAL, Sezione Allevamenti Zootecnici, University of Bologna, Via Fratelli Rosselli107, 42100 Reggio Emilia, Italy.

Other authors (name only):
1. Davoli Roberta
2. Fontanesi Luca
3. Stella Alessandra
4. Hedegaard Jakob
5. Hornshøj Henrik
6. Colombo Michela
7. Nanni Costa Leonardo
8. Russo Vincenzo

**Abstract:**
Stress reactions to slaughtering influence muscle metabolism and technological meat quality. The aim of this work was to identify genes involved in stress reactivity in pigs by using DNA microarray analysis. A total of 33 pigs of three breeds: Italian Large White (ILW), Italian Duroc (ID) and Pietrain (PI) were bred on the same farm and slaughtered in a single day on the same abattoir. All ILW and ID pigs had genotype 1843CC at RYR1 locus while PI pigs were CC and CT. At the abattoir the pigs were divided into two groups and only one was physically stressed by running along the raceway before slaughtering. After slaughter, samples of semimembranosus muscle were collected for RNA extraction. Labelled cDNAs were hybridized to 11K pig oligoset version 1.0 (Operon-Quiagen) microarrays using a heart cDNA sample as common reference. The obtained data were analyzed in SAS using a mixed model. Using DAVID Bioinformatic Resources the ontology of differentially expressed genes among stressed and not-stressed pigs was produced for each breed. The results showed that the impact of preslaughter stress on metabolism determines different responses between breeds in term of differentially expressed genes. The differential expression of stress related genes will be validated by qRT-PCR.

**Poster 2132**

**Title:** The lavender plumage colour phenotype in Quail is caused by a large deletion in MLPH gene region

Presenting Author: BED’HOM Bertrand, INRA/AgroParisTech UMR1236 Génétique et Diversité Animales, Domaine de Vilvert, Bâtiment 211, F-78352 Jouy en Josas Cedex, France

Other authors (name only):
1. VAEZ Mohsen
2. FOLLETT Sarah A
3. COVILLE Jean-Luc
4. GOURICHON David
5. TIXIER-BOICHARD Michèle
6. MINVIELLE Francis
7. BURKE Terry

**Abstract:**
The lavender phenotype in Japanese Quail is recessive and autosomal. It produces a dilution of both eumelanin (from black to blue or light grey) and phaeomelanin (from red/brown to buff or beige) in feathers. A similar phenotype is also known in several mammalian species (Mouse, Dog, Cat, Mink, Human) and in Chicken. All these phenotypes are caused by mutations in MLPH (Melanophilin) gene, which is involved in melanosomal transport in melanocytes. Homology between Japanese Quail and Chicken lavender loci has been demonstrated by hybridization between these animals (Minvielle et al., 2002). We have previously established that the mutation in chicken MLPH is a single-point mutation in exon 1, implying amino-acid change in protein (Vaez et al., 2008). By comparison of genomic organization in MLPH region between homozygous lavender and wild-type quails, we describe here that the mutation in Japanese Quail is surprisingly a large deletion encompassing the 3’ end of MLPH gene (11 exons out of 17 are deleted), the entire PRLH (Prolactin Releasing Hormone) gene and the 3’ end of RAB17 (member of RAS oncogene family) gene (5 exons out of 6 are deleted).

**Poster 2133**

**Title:** A genome scan for athletic performance in the thoroughbred

Presenting Author: Keith Durkin, Department of Veterinary Integrative BioSciences, Texas A & M University, College Station TX 77843, USA

Other authors (name only):
1. Terje Raudsepp
2. Loren C. Skow
3. Bhanu P. Chowdhary

**Abstract:**
The primary goal of the thoroughbred industry is to breed and train superior equine athletes capable of excelling on the racetrack. To date research into the genetic underpinnings of athletic ability has been limited in the horse. Advances in equine genomics and the genetics of athletic performance in humans spurred
us to begin investigations into this important trait. We initially mapped 47 candidate genes associated with human athletic performance in the equine genome by radiation hybrid (RH) and fluorescent in situ hybridization (FISH) mapping. Using RH data and later the draft equine sequence we identified microsatellites adjacent to these and other candidate genes, additional microsatellites were added to increase genome coverage producing a final panel of 205 markers. All potential markers were initially screened on a pool of 16 thoroughbreds to ensure they were polymorphic. The panel was genotyped on 161 thoroughbreds; Centimorgans (cM) between microsatellites were determined with CRI-MAP. The animal’s athletic ability was estimated using career winnings which were log transformed to create a linear trait; unraced animals were treated as missing data. The data was analyzed in the program MERLIN using the non-parametric linkage analysis option. None of the genotyped microsatellites showed significant linkage with career earnings.

**Poster 2134**

**Title:** Novel mutations of bovine FASN gene, g.16024A>G and g.16039T>C, effects on the fatty acid composition of Japanese black beef.

Presenting Author: Tsuyoshi Abe/1, Odakurahara, Nishigo-Mura, Fukushima, 9618511, JAPAN

Other authors (name only):
1. Junichi Saburi
2. Hiroyuki Hasebe
3. Tetsuo Nakagawa
4. Satsuki Misumi
5. Toshihiro Nade
6. Hiroaki Nakajima
7. Noriaki Shoji
8. Masato Kobayashi
9. Eiji Kobayashi

**Abstract:**
We constructed the F2 reference family, derived from the cross between Japanese black and Limousin, and detected significant QTL for fatty acid composition of back fat, intermuscular fat, and intramuscular fat at 60 to 70cM region on BTA19. We then focused and analyzed bovine fatty acid synthase (FASN) gene as a candidate for this QTL. Eight novel and 4 known mutations were detected in the coding sequence of the FASN gene between two breeds that were used to produce our F2 population. Among these mutations, 6 were nonsynonymous, and in particular, g.16024A>G and g.16039T>C, detected in exon 34, which determine amino acid substitution of threonine (T) to alanine (A) and tryptophan (W) to arginine (R), respectively, were clearly separated in the 2 parental breeds. Two haplotypes comprising these 2 mutations (TW and AR) were segregated in F2 individuals, and had a significant effect on the fatty acid composition of three adipose tissues above. The TW haplotype was associated with increasing C18:0 and C18:1 content and MUFA/SFA, and decreasing C14:0, C14:1, C16:0, and C16:1 contents, respectively. Furthermore, we observed a markedly higher frequency of the TW haplotype in Japanese Black (0.67) than in Holstein (0.17), Angus (0.02), and Hereford (0.07).

**Poster 2135**

**Title:** Comparative Study on Gene Expression of DNA Methyltransferases in Gallus gallus and Mus Musculus

Presenting Author: Taeko Isokane, Graduate School of Biosphere Science, Hiroshima University, 1-4-4, Kagamiyama, Higashi-Hiroshima, JAPAN; 739-8528

Other authors (name only):
1. MOHAMED E. GHANEM.
2. NORIKO HIRAIWA
3. TAKASHI SHIMOGIRI
4. HIROSHI YASUE
5. MASAHIDE NISHIBORI

**Abstract:**
In mammals, genomic imprinting leads to the monoallelic expression of specific genes. Commencing with genomic imprinting, DNA methylation is an essential process in the course of development, including X-chromosome inactivation. Five DNA methyltransferase genes (Dnmts) are involved in the methylation in mammalian species. However, avian species are reported to have no methylation in their genomes, which indicates that little transcription occurs from the genes orthologous to mammalian Dnmts. In the present study, the amounts of transcripts from 4 Dnmts (no ortholog of Dnmt3L in chicken) were determined in chicken 4 organs (Brain, Liver, Kidney, Testis) using real-time RT PCR through developmental stages, and compared with those of corresponding mouse organs. In the course of development, the amount of Dnmt1 mRNA was consistently lower in chicken organs than in the corresponding mouse organs; those of Dnmt2 and Dnmt3b mRNAs were low in chicken as well as in mouse organs; that of Dnmt3a mRNA was generally higher in chicken than in mouse. These findings indicate that the 4 Dnmts in chicken may contribute to the DNA methylation, though the methylation has not been reported so far; or that they have different roles other than methylation in chicken development.
**Poster 2137**

**Title:** Isolation and characterization of 27 novel microsatellites for individual identification and parentage testing in Humboldt penguin (*Spheniscus humboldti*)

Presenting Author: Kei Onga, Department of Life Science, Faculty of Agriculture, Meiji University, Higashimita, Tama-ku, Kawasaki-city, Kanagawa 214-8571 Japan

Other authors (name only):
1. Teruaki Tozaki
2. Michio Fukuda
3. Akihiko Ohta

**Abstract:**

The Humboldt penguin is one of the most endangered species in the wild. To conserve the species, captive breeding has become important at zoological organizations. High levels of genetic diversity are also essential for species survival. Therefore, appropriate management of individual pedigrees is required. Microsatellites are widely used in parentage testing and population genetic studies in various species. However, no microsatellites for parentage testing have been reported in the Humboldt penguin. We constructed a genomic library enriched with (CA)n repeats. Thirty-one novel microsatellites were efficiently isolated from the library, and were genotyped in 24 Humboldt penguin individuals. Of these, 27 microsatellites were polymorphic. Heterozygosity ranged from 0.042 to 0.917, with an average of 0.514, while the number of alleles ranged from 2 to 13, with an average of 5.3. Power of discrimination (PD) and probability of exclusion (PE) ranged from 0.080 to 0.970 and 0.020 to 0.740, respectively, resulting in total PD and PE values of $1 - 1.80 \times 10^{-19}$ and 0.9999994, respectively. Application of these novel microsatellites will be a powerful tool for individual identification and completion of pedigree records in setting up captive breeding plans.

**Poster 2138**

**Title:** Allelic Analysis of Thoroughbred Horses Using Fourteen Microsatellite Markers

Presenting Author: SHIN-WOOK KANG & Korea Racing Authority, 685, Juam-dong, Gwacheon-si, Gyeonggi-do, South Korea, 427-711

Other authors (name only):
1. Sun-Young Lee
2. In-Su Cho
3. Man-Bae Hu
4. Sung-Jean Kim

**Abstract:**

Microsatellite are repetitive DNA sequence that are randomly distributed throughout vertebrate genome. In this study, we experimented the total 1,660 thoroughbred horses. Genomic DNAs were prepared from hair roots samples. Genomic DNAs from Thoroughbred horses were extracted using MagExtractor System MFX-2000 according to the manufacturer's protocols. Muti-plex PCR was performed to respectively amplify alleles using horse DNA kit. We analyzed a number of alleles, allelic frequencies, observed(Ho) and expected(He) heterozygosity, probability of exclusion(PE), total PE for parentage test.

The number of alleles of the markers was varied between 4 and 10 with an average number of alleles of 6.6. The heterozygosity were ranged from 0.527 to 0.855 and mean expected heterozygosity was 0.706. Observed PIC was from 0.494 (CA425) to 0.817 (ASB2) and mean PIC was 0.659. The PE was observed from 0.314(CA425) to 0.674(ASB2) and total PE of all markers was 0.9999. These results verify that fourteen microsatellite markers used in this study are very powerful for parentage test and individual identification of thoroughbred horse in Korea.

**Poster 2139**

**Title:** Fine mapping of ovine microphthalmia

Presenting Author: Adrian Brunner, Institute of Genetics, University of Berne, Switzerland

Other authors (name only):
1. Jens Tetens
2. Michaela Drögemüller
3. Tosso Leeb
4. Cord Drögemüller

**Abstract:**

The ovine microphthalmia (OMO) is characterized by bilateral abnormal small eyes in newborn lambs. This congenital anomaly is inherited as a monogenic autosomal recessive trait with complete penetrance and is predominantly found in the Texel breed. We assigned the OMO locus by linkage mapping to a 12.3 cM interval on OAR23. This chromosome region corresponds to segments of HSA18 and BTA24. We constructed a BAC contig covering the region of interest based on sheep-human BLASTN sequence alignments. For fine mapping purposes we scanned the bovine genome sequence for additional microsatellites and comparatively anchored ovine BAC end sequences (BES). PCR primers were designed from the anchored ovine BES and SNPs were developed in targeted regions by re-sequencing these PCR products. Eight new polymorphic
microsatellites and 206 SNPs were identified using this approach. These markers cover the entire critical interval with marker spacing between 5 and 200 kb. Genotyping of 117 affected lambs and 163 healthy relatives belonging to 60 sheep families lead to a reduction of the critical interval to approximately 8 Mb on the virtual OAR23 map.

Poster 2140
Title: Association of PPARGC1A with growth in a Charolais x German Holstein F2 resource population

Presenting Author: Annett Eberlein, Research Institute for the Biology of Farm Animals (FBN), 18196 Dummerstorf, Germany

Abstract:
The birth weight of calves is an important economic trait and strongly associated with dystocia, stillbirth and hypothermia, the most frequent causes for perinatal deaths. Birth weight is also used as an initial reference point for monitoring the development of an individual animal, because it represents the result of the most dynamic process of prenatal growth and development. Several studies reported overlapping QTL for birth weight, calving traits and stillbirths on bovine chromosome 6 (BTA6) in different breeds. On this account QTL detection for birth weight was performed in F2 individuals generated by embryonic transfer as well as their P0 and F1 ancestors of a Charolais x German Holstein resource population. The analysis shows a genomewise significant QTL (p<0.05) for birth weight on BTA6. This is in agreement with a previously reported QTL for stillbirth in the German Holstein population. Within this region the PPARGC1A gene (peroxysome proliferator-activated receptor γ coactivator 1α) is located, a potential candidate gene, because of its function in energy, fat and glucose metabolism as a basis for growth and development. SNPs in the PPARGC1A and other candidate genes located in the targeted chromosomal region were tested for association with prenatal growth.

Poster 2141
Title: Bovine chromosome 18: Mapping of quantitative trait loci for somatic cell score and udder conformation traits

Presenting Author: Bodo Brand, Research Institute for the Biology of Farm Animals (FBN), 18196 Dummerstorf, Germany

Other authors (name only):
1. Manfred Schwerin
2. Christa Kühn

Abstract:
To improve udder health and resistance against clinical mastitis (CM) many studies have attempted to identify chromosomal regions, genes and polymorphisms that exert influence on udder health in order to include this information in upcoming breeding strategies. So far QTL for CM or somatic cell score (QTL_{SCS}) have been detected on all autosomes. On bovine chromosome 18 (BTA18) several studies detected QTL_{SCS} in the telomeric region where also a QTL for mastitis incidence was found. To further characterise this QTL_{SCS}-region on BTA18 QTL detection in 6 large half-sib families in the German Holstein population was performed. Merging information from published linkage-maps, RH-maps, human and bovine sequence-assemblies as well as own linkage- and RH-mapping results enabled definition of a refined marker order and highlighted discrepancies in the bovine NCBI sequence-assembly Build 3.1. In order to gather information about the functional background of the QTL_{SCS} QTL analysis for SCS and functional traits with potential impact on SCS were performed. First results show that experimentwise significant QTL for SCS and udder conformation traits segregate in the telomeric region of BTA18. To check the value of the QTL_{SCS} information an exemplary marker assisted selection of animals with high and low susceptibility to mastitis was performed.

Poster 2142
Title: Hypotrichosis in a Charolais x Holstein cross is not associated with allelic variants in the bovine SILV gene

Presenting Author: Christa Kühn, Research Institute for the Biology of Farm Animals (FBN), 18196 Dummerstorf, Germany

Other authors (name only):
1. Rosemarie Weikard

Abstract:
A hypotrichosis phenotype, also known as rat-tail syndrome, is segregating in a Charolais x German Holstein F2- resource population as well as in backcrosses from the F2 individuals to a purebred German Holstein sire. The phenotype is characterised by curly, sparse hair in pigmented regions of the skin, whereas unpigmented hair is unaffected. Recently,
there had been reports that mutations in the bovine SILV gene that harbours polymorphisms associated with Charolais coat colour dilution (Dc) might also be the functional background of the hypotrichosis phenotype. Sequencing the entire coding region as well as the promoter of the SILV gene of seven individuals with different dilution, background coat colour and rat-tail-phenotypes did not reveal any polymorphism in total concordance with the expression of the hypotrichosis phenotype. For the bovine SILV gene, multiple splice variants have been described. However, sequencing SILV transcripts in skin from individuals with different dilution and hypotrichosis phenotypes did not reveal any indication on splice variation being responsible for the defect in hair structure. Thus, we conclude that segregation of hypotrichosis is independent from the SILV dilution locus. A refined model of inheritance of hypotrichosis in the Charolais x German Holstein F2 population will be presented.

Poster 2143

Title: Homozygosity mapping of arachnomelia in Brown cattle

Presenting Author: Tosso Leeb, Institute of Genetics, University of Berne, Switzerland

Other authors (name only):
1. Marco Rossi
2. Arcangelo Gentile
3. Stefania Testoni
4. Hannes Jörg
5. Marie-Louise Glowatzki-Mullis
6. Gerald Stranzinger
7. Cord Drögemüller

Abstract:
Fifteen calves affected with arachnomelia were sampled in the Swiss and Italian Brown cattle population. Analysis of the pedigree data confirmed monogenic autosomal recessive inheritance and revealed that all affected individuals traced back to a single acknowledged carrier founder sire. Eight affected offspring and 10 available relatives were genotyped for a panel of 240 microsatellites spanning the 29 bovine autosomes. Homozygosity mapping showed homozygosity of all cases at two adjacent microsatellite markers on BTA5. For fine mapping purposes 14 additional BTA5 microsatellites were genotyped in all available cases and indicated shared homozygous haplotypes across 20 Mb among affected individuals. A second pedigree with 23 acknowledged carriers among 34 sires in total related to the founder were genotyped for these markers. Linkage and haplotype analysis showed that the arachnomelia mutation could be assigned to a 7.2 Mb candidate region (LOD score >6). The linked markers can be used for indirect determination of the arachnomelia genotype in offspring of related carrier sires. This bovine chromosome segment corresponds to an ortholog segment of HSA12q containing more than 140 annotated genes and loci. The only obvious functional candidate gene of this region GDF11 is probably not responsible for this fatal bovine disorder.

Poster 2144

Title: Genotyping methodology and frequencies of the myostatin F94L mutation in Australian cattle breeds.

Presenting Author: Dianne M Vankan, University of Queensland, Animal Genetics Laboratory School of Veterinary Science, St Lucia, Qld 4072 Australia

Other authors (name only):
1. Sean Corley
2. Deanne Wayne
3. Marina Fortes

Abstract:
The F94L SNP (g.433C>A) in the growth differentiation factor 8 (GDF8) gene, commonly known as the myostatin (MSTN) gene, has recently been shown to be functionally associated with increased muscle mass and carcass yield in cattle. Since the F94L mutation is not, like other MSTN mutations, associated with reduced fertility and dystocia, it is a candidate for introgression into other breeds to improve retail beef yield, and development of an accurate test for genotyping this mutation is warranted. Accurate genotyping of the F94L SNP by primer extension is not possible so genotyping has been achieved using PCR-RFLP. We found that variations in the efficiency of cleavage of PCR amplicons by Taq1 compromised the accuracy of genotyping, resulting in an overestimation of the frequency of the mutant A allele. We developed two real-time PCR assays that both accurately genotype the F94L mutation in cattle based on allele-specific and high resolution melt analysis. The frequency of the F94L mutation was determined in 960 animals from 14 breeds of cattle. The mutation was present in Simmental (0.8%), Piedmontese (2%), Droughtmaster (4%) and Limousin (94.2%) but not found in Salers, Angus, Poll Hereford, Hereford, Gelbvieh, Charolais, Jersey, Brahman, Holstein and Maine Anjou.
Poster 2145

Title: Gene expression profiling in pig liver tissue following administration of Clenbuterol-HCL

Presenting Author: Qiuyue Liu, Department of Biochemistry and Molecular Biology, China Agricultural University, Beijing 100193, China. Xiaoxiang Hu, Department of Biochemistry and Molecular Biology, China Agricultural University, Beijing 100193, China. Ning Li, State Key Laboratory for Agrobiotechnology, China Agricultural University, Beijing 100193, China.


Abstract:
Clenbuterol-HCL can greatly reduce fat deposition and increase skeletal muscle growth at high dosages. In order to advance our understanding of the fundamental molecular mechanisms, Genechips, real-time PCR, two-dimensional electrophoresis and mass spectra were used to study the differential gene expression profiles of liver tissues among the pigs with/without Clenbuterol-HCL. Four Affymetrix Porcine Genome Genechips were used for global evaluation of the gene expression. Totally 692 probe sets presented on Microarrays were identified differentially expressed with 337 upregulated and 355 downregulated (False discovery rate adjusted p-values < 0.05). Gene ontology functional annotation analysis was applied to the above genes, and the most significantly overrepresented categories were deemed the organismal physiological process, enzymes of lipid metabolism, receptors of membrane, and mediators involved in positive regulation of signal transduction. Moreover, pathway mining analysis revealed that physiological pathways such as MAPK, Wnt, cell adhesion molecules as well as signaling pathway from G-protein families, etc have been regulated remarkably when Clenbuterol-HCL was feed. Results obtained in this study deepen our understanding in the function mechanism of β-adrenergic receptor agonists on animal models. Differentially expressed genes identified will be the candidates for further investigations on the molecular mechanisms involved in Clenbuterol-HCL of reducing adipose accumulation.

Poster 2146

Functional analysis of the equine kappa casein (CSN3) gene promoter

1Hobor Sebastijan, 1Kunej Tanja, 2Majdić Gregor and 1Dovc Peter
1Department of Animal Science, Biotechnical Faculty, University of Ljubljana, Groblje 3, 1230 Domzale, Slovenia
2Veterinary Faculty, University of Ljubljana, Gerbiceva 69, 1000 Ljubljana, Slovenia

Kappa casein gene (CSN3) is the fourth member of the casein gene cluster in mammals, which is evolutionary not related with the other three casein genes. However, the structure of its promoter allows the spatially and temporally coordinated expression with the other three casein genes. In our study we sequenced 2286bp of the equine CSN3 promoter and performed in silico search for potential transcription factor binding sites. Thirty two potential TFBSs were found by manual search within the 2286 bp region of the CSN3 gene promoter and additional ten were identified by bioinformatic tools. Fifteen SNPs were discovered by sequencing of the promoter and 12 of them were located within the identified TFBSs, potentially interfering with TF binding. We found SNPs in putative binding sites for NF1, C/EBP and TFID, which are all presumed to be involved in expression of casein genes. Sequence alignment of the core CSN3 gene promoter (1234bp) of nine species (sheep, goat, cow, zebra, donkey, horse, human, chimp, macaque), revealed 22 completely conserved sequence stretches, six to fourteen bp in length and seven of them harbored potential transcription factor binding sites (TRFBs). Since many of these potential TFBSs may not be functional, it is possible that many of the discovered SNPs are not associated with TF binding. Electrophoretic mobility shift assay (EMSA) using synthetic DNA fragments representing potential equine CSN3 gene promoter elements has been performed. Furthermore, reporter gene constructs containing different parts of the equine CSN3 promoter were constructed and tested in the heterologous (bovine and murine) cell system.

Poster 2147

Title: Identification of the causative mutation of congenital pseudo-myotonia in Chianina cattle

Authors:
Presenting Author: Michaela Drögemüller, Institute of Genetics, University of Berne, Switzerland
Conference registration number of Presenting Author: 17378

5. Cord Drögemüller

**Abstract:**
Congenital pseudo-myotonia (PMT) in Chianina cattle is a muscle function disorder which is mainly characterized by an exercise-induced muscle contracture which prevents animals from performing muscular activities more intense than a simple walk at a slow pace. Mutations in the human ATP2A1 gene, encoding a fast-twitch skeletal-muscle Ca$^{2+}$ ATPase (SERCA1), cause Brody myopathy, a very similar rare autosomal recessive disorder characterized by exercise-induced muscle cramps and impaired muscle relaxation. The analysis of the collected Chianina pedigree data suggested monogenic autosomal recessive inheritance and revealed that all PMT affected individuals traced back to a single founder sire. A family with 16 PMT affected cattle was genotyped with two bovine ATP2A1 gene flanking microsatellites. Linkage analysis within this family showed that the PMT mutation could be assigned to the ATP2A1 gene region on BTA25 (LOD score >3). Subsequent DNA sequencing of the 16 PMT affected calves revealed a missense mutation (c.491G>A) leading to a p.Arg164His substitution in exon 6 of ATP2A1. Arg164 in bovine ATP2A1 is located within the functional important N-terminal actuator domain and is a highly conserved residue. Genotyping 112 unaffected unrelated Chianina animals did not reveal a single individual homozygous for mutation and indicated a carrier frequency of >0.10.

---

**Poster 2148**

**Title:** Genetic Characterization of Korean Cattle (Hanwoo) using SNP Markers

Songlan Li$^1$, Kyung-Tak Do$^1$, Jong-Joo Kim$^2$, Du-Hak Yoon$^3$, Hyeong-Koon Myeong$^2$ and Kwan Suk Kim$^1$

$^1$Department of Animal Science, Chungbuk National University, Cheongju, Korea.
$^2$School of Biotechnology, Yeungnam University, Gyeongsan, Gyeongbuk, Korea
$^3$Animal Genomics & Bioinformatics Division, National Institute of Animal Science, Suwon, Korea

Korea government has introduced genetic testing to identify the origin of beef and to assure safety and quality of domestic “Hanwoo” beef products. To develop reliable and accurate markers, we used to single nucleotide polymorphisms (SNP) based on the information of NCBI bovine SNP database. The SNP sequence data developed at the BCM-HGSC were used for DNA sequencing of Hanwoo, Hereford and Limousin. One hundred four SNP sequences spanning 27.5 kb were randomly selected and analyzed by direct sequencing of 24 individuals for each breed. A total of 270 SNPs were identified from the sequence comparisons, among which 22 SNPs were further genotyped for the 960 animals with Hanwoo or other foreign breed origin. Thirteen SNP markers showed significant genotypic differences between Hanwoo and other beef breeds, which may enable designing of robust and accurate high-throughput genotyping assays for Hanwoo meat identification.

---

**Poster 2149**

**Title:** Identification of new pig cSNPs using polymorphisms between Korean native pig and Genbank registered sequences.

Presenting Author Kwangha Park, Address: Laboratory of Mammalian Genomics, Department of Animal Biotechnology, Konkuk University, Seoul 143-701, South Korea.

Other authors (name only)  
1. Vijaya Ramu Dirisala  
2. Younshin Oh  
3. Hojun Choi  
4. Kyungtae Lee  
5. Jinhoi Kim  
6. Hoontaek Lee  
7. Kunho Suh  
8. Chankyu Park

**Abstract**
Sequences from the clones of full-length enriched cDNA libraries serve as valuable resources for functional genomic studies. We have analyzed 1,970 high quality chromatograms (Phred value ≥ 30) that were obtained from sequencing the 5’ends of brainstem, liver, neocortex and spleen clones derived from full-length enriched cDNA libraries from Korean native pigs. In addition, 50,000 pig EST sequence trace files were obtained from Genbank and combined with our sequencing information to facilitate SNP identification in silico. The process generated 8,118 contigs containing 678 putative cSNPs. Of these, 33 putative cSNPs were randomly selected for confirmatory analysis and validated using 20 pigs from four different breeds (Duroc, Landrace, Yorkshire, Korean native pig). 20 of the 33 putative cSNPs were confirmed with a confirmation percentage of 61%, which was similar to that reported by other studies. We also identified 15 new cSNPs from the validation process, which were not detected by our in silico analysis. Our study shows that analyzing genetically diverse breeds pig breeds such as the Korean native pig could serve as a useful strategy for generating a large number of cSNPs.
**Poster 2150**

**Title:** Regulatory SNPs and fat related traits in cattle. Implications of the bovine g.763G>C FASN SNP on milk fat content.

Presenting Author: Laura Ordovas, Laboratorio de Genetica Bioquimica, Facultad de Veterinaria, Miguel Servet 177, 50013 Zaragoza, Spain

Other authors (name only):
1. Laura Ordovas
2. Rosa Roy
3. Sandra Pampin
4. Pilar Zaragoza
5. Arianne Sanz
6. Jose Carlos Rodriguez-Rey
7. Clementina Rodellar

**Abstract:**
Recent surveys have estimated that between 22% and 35% of SNPs located in regulatory regions (rSNPs) have functional consequences. The accumulation of several rSNPs could constitute the basis for the different gene expression profiles that define traits in a species-, breed- or individual specific manner. The aim of our research is to identify and functionally characterize rSNPs in candidate genes for fat related traits in cattle with the ultimate goal of obtaining genetic profiles that could define them. As result, we previously reported the identification of a Fatty Acid Synthase (FASN) SNP, g.763G>C, that was significantly associated with milk-fat content in dairy cattle. The SNP is included in a GC-rich region that may constitute a cis element for Sp transcription factors (TFs). In this work, the functional characterization of the g.763G>C SNP is presented. Reporter assays in 3T3L1, HepG2 and MCF-7 cells showed that the SNP affected the bovine FASN promoter activity. Supershift assays (EMSA) demonstrated that the Sp1/Sp3 TF-binding ability of the sequence was also altered by the SNP in these cellular lines. In bovine lactating mammary gland, EMSA assays showed that the specific binding of Sp3 may account for the association of the g.763G>C SNP with milk-fat content.

**Poster 2151**

**Title:** "Discarded SNPs to be used for identification even showing intermediate frequencies"

Presenting Author: Clementina Rodellar, Laboratorio de Genetica Bioquimica, Facultad de Veterinaria, Miguel Servet 177, 50013 Zaragoza, Spain

Other authors (name only):
1. Sanz

**Poster 2152**

**Title:** High exclusion probability in ovine genealogical control through 18plex and 20plex PCR

J.A. Bouzada1,2, T. Mayoral2, J.M. Lozano1,2, B. Ossorio1,2, M. Martínez-Galiana1,2, M. Estévez1,2, E. Anadón, y C. Gómez-Tejedor2

1 Sanidad Animal y Servicios Ganaderos, S.A. (TRAGSEGA). Dirección de estudios y consultoría. C/ Conde de Peñalver 84, 28006 Madrid
2 Laboratorio Central de Veterinaria. Ctra. de Algete, Km. 8. 28110 Algete MADRID

**Abstract:** Genealogical control is one of the weaknesses in the Selection and Breeding Programs in sheep breed mainly due to the reproduction systems, based in the use of large number of females with several males more than artificial insemination. It is very important to develop a system allowing certify animals genealogy as well as the unequivocally identification of them throughout their life. Central Veterinary Laboratory Algete (LCV) in collaboration with TRAGSEGA has developed a high-exclusion
capacity, *multiplex* PCRs system which amplify, in two reactions 18 (BM1818, CSRD247, ETH152, D551, HSC, ILST87, INRA005, INRA006, INRA023, INRA049, INRA063, INRA172, MAF65, McM42, McM527, OarCP49, OarFCB20, SPS113 y SPS0115) and 20 (BMS1967, BP28, BP33, CDS, CSRD2111B, CSRD2115B, DU330122, ILSTS05, ILSTS39A, ILSTS62A, INRA081, MAF209, MMP9, OarAE119, OarFCB128, OarJMP11, TCRVB6, TGLA53, THRA, UMJM30) microsatellite markers, respectively. All of them have been selected from the list proposed by the International Society for Animal Genetics (ISAG) for the 2007-08 Ovine Comparison Test and used for studies in this specie. Primers were adapted to achieve a final configuration that allowed all markers to be analysed together. Robotic procedures were implemented to minimize the risk of genotyping errors and parentage verification mistakes. The proposed system has been designed to process a very large number of samples in a short space of time ensuring a perfect reliability and traceability of the obtained results. The main set of markers is used routinely for ovine parentage verification at LCV and reached a exclusion probability of 99,9999930%. It further provides for a panel composed of 20 additional markers to be used in cases where there is a need for greater exclusion capacity.

**Poster 2153**

**Title:** Effect of a SNP in the promoter region of the lactoferrin gene upon ovine milk somatic cell counts

**Presenting Author:** Maria Rosário Marques, Escola Superior de Biotecnologia, Universidade Católica Portuguesa, Rua Dr. António Bernardino de Almeida, P-4200-072 Porto, Portugal

**Other authors (name only):**
1. Vanessa Ralha
2. Carlos Carmona Belo
3. Maria Manuela Pintado
4. Francisco Xavier Malcata

**Abstract:**
Lactoferrin (LF) is an iron-binding glycoprotein, which is naturally found in milk. Its concentration in milk increases dramatically in response to udder infection, which may be an indication of its role upon prevention of/against mastitis. The occurrence of subclinical mastites has been related to increased levels of somatic cell counts (SCC) in milk. The objective of this effort was to find single nucleotide polymorphisms (SNP) in the promoter region of the LF gene, and ascertain their effects upon SCC in ewes’ milk.

Polymerase chain reaction – single strand conformation polymorphism analysis (PCR-SSCP) was carried out in a DNA fragment of the promoter of the LF gene. Three SSCP patterns were detected and sequenced, and a SNP (A/G) was found at position 73 (GenBank sequence nº AF091651). Seventy-six Lacaune crossbred sheep were genotyped. The frequencies of the three SSCP patterns, which correspond to genotypes AA, AG and GG, were 71.1%, 27.6% and 1.3%, respectively. Mixed model data analysis revealed that AA ewes tended to have significantly lower SCC in milk than AG ewes (P<0.10), but no differences were observed on milk production. The aforementioned SNP disrupts a putative transcription factor binding site in the promoter of the LF gene – hence suggesting its involvement in the regulation of LF expression levels.

**Poster 2154**

**Title:** Expression profile of the MRF4 gene in muscle during postnatal growth of pigs.

**Presenting Author:** Mariusz Pierzchala, IGAB PAS, 05-552 Jastrzebiec. Poland

**Other authors (name only):**
1. Paweł Urbański
2. Paweł Lisowski
3. Joanna Wyszyńska-Koko
4. Marian Kamyczek
5. Marian Różycki

**Abstract:**
The MRF4 (herculin) is a member of the basic-helix-loop-helix family of proteins. The basic Helix-Loop-Helix (bHLH) proteins are transcription factors that play important roles during the cells development of various tissues. The myogenic factor MRF4 gene is expressed in a unique pattern defined by period and place within the skeletal muscle. The goal of our study was to analyse the MRF4 gene expression profile during postnatal growth pigs. As the study material the 5 different pig breeds were chosen: Polish Large White, Polish Landrace, Duroc, Pietrain and Pulawska (polish native pig breeds). Two muscle probes - m. longissimus dorsi and m. semimembranosus were collected from pigs at slaughter at the age of 60, 120 and 180 days. Quantification of transcripts were performed by Real-time PCR technique. The MRF4 gene transcript level was standardised to three reference genes such as: B-ACT, GAPDH, and B2M. The results show significant variation in the level of MRF4, related to breed and the age of slaughter. The highest difference were observed between Duroc, Pietrain and Polish Landrace or Pulawska. This confirm such presumption that selection of pig breeds would result in variation of metabolic pathways of myogenic factor (MRF4) regulation.
Poster 2155

**Title:** Polymorphism of the ADIPOR1 gene and its transcription level in two skeletal muscles during postnatal development in five pig breeds.

Stachowiak M.¹, Flisikowski K.², Fries R.², Switonski M.¹

¹ Department of Genetics and Animal Breeding, University of Life Sciences in Poznan, Poland; ² Chair of Animal Breeding, Technical University of Munich, Freising, Germany

**Abstract:**

The adiponectin receptor 1 gene (ADIPORE1) can be considered as a candidate for the pig fatness and growth. The aim of the study was searching for its polymorphism as well as transcription level in two skeletal muscles (m. longissimus dorsi, m. semimembranosus). Five breeds were analysed: Polish Large White, Polish Landrace, Duroc, Pietrain and Pulawska. Tissue samples were collected post mortem at the age of: 60, 90, 120, 150, 180 and 210 days.

Transcription start site and a nucleotide sequence of the 5'UTR region was established. The transcript of the porcine ADIPOR1 gene contains 190 bp of the 5'-UTR sequence (GenBank, EU556496) and its sequence is similar to a previously reported ADIPOR1 transcript in human with the identity of 85%. In this fragment the Kozak consensus sequence occurs at the position 188–194 bp. Altogether, eleven SNPs were detected in exon 5, intron 6 and 3'UTR. Genotyping performed for a silent SNP in exon 5 (759 G>A) and two SNPs in 3’ UTR (+129 A>C and +536 A>G) revealed interbreed variability in terms of allele frequencies. Differences between breeds in the transcript level occurred mainly at early stages of the postnatal development (60, 90 and 120 days).

Poster 2156

**Title:** Single Nucleotide Polymorphism (SNP) discovery from the horse Y chromosome (ECAy)

Presenting Author: Maria do Mar Oom, University of Lisbon, Faculty of Sciences, DBA/CBA C2-3º Piso, Campo Grande, 1749-016 LISBOA, PORTUGAL

Other authors (name only):
1. Sandra Afonso
2. Bhanu P. Chowdhary
3. Terje Raudsepp

**Abstract:**

Mammalian Y chromosome is male specific, constitutively haploid and inherited exclusively through patrilineages. Polymorphic markers derived from male specific region of Y (MSY) are, thus, ideal tools to trace recent evolutionary events in the populations of horses and other equids. Previous attempts to identify SNP or microsatellite polymorphism from ECAy have not been successful. Taking advantage of the recently developed BAC contig map of the entire ECAy we conducted a systematic search for SNPs both from MSY and the pseudoautosomal region (PAR). BAC end sequences were used to generate 54 STSs (sequence-tagged sites) which collectively covered ~30 kb. All STSs were sequenced from: i) a DNA pool of 39 male horses and ponies representing 11 diverge breeds; ii) Bravo – the DNA donor for CHORI BAC library; iii) 1 tuva horse; iv) 1 Mongolian horse; v) 2 Equus przewalskii, and, vi) 1 donkey. Preliminary sequence analysis has already detected a few SNPs from the ampliconic region of MSY and from PAR.

Poster 2157

**Title:** MHC typing in swine: SLA class I and class II-allele distribution in Austrian Large White, Landrace, and Pietrain breeding stocks

Presenting Author: Sabine E. Hammer, Institute of Immunology, University of Veterinary Medicine Vienna, Veterinärläkplatz 1, A-1210 Vienna, Austria

Other authors (name only):
1. Julia Deutsch
2. Barbara C. Rütgen
3. Sandra Groiß
4. Wilhelm Gerner
5. Simone Müller,
6. Armin Saalmüller

**Abstract:**

MHC (major histocompatibility complex) genes encode cell surface glycoproteins presenting antigenic peptides to T cells. They are highly polymorphic suggesting that diversity in MHC genes is a good measure of population fitness. Resource herds of swine leukocyte antigen (SLA)-characterized pigs are valuable large animal models for biomedical research in terms of immune responses, disease resistance, and production traits. Furthermore detailed knowledge about the distribution of MHC genes in a population enabled a more target-oriented design of vaccines and the development of specific reagents for studying correlates of protection like tetramers. This study represents the initial characterization of founder haplotypes of purebred Large White, Landrace, and Pietrain populations. The respective founder SLA-haplotypes were detected by a reverse transcription-polymerase chain reaction (RT-PCR)-based SLA typing method to clone and DNA sequence the putative alleles at four SLA class Ia loci, designated as SLA-1, SLA-2, SLA-3, and SLA-6 and four SLA
class II loci, SLA-DQA1, SLA-DQB1, SLA-DRA1, and SLA-DRB1. The obtained sequences are used to establish rapid SLA typing assays based on sequence-specific PCR primers (PCR-SSP) to customize allele-specific PCR screening assays for large-scale MHC typing of Austrian pig populations and on a next level European pig herds.

Poster 2159

Title: Genetic variation in pro-melanin-concentrating hormone affects carcass tenderness and fat levels in Bos taurus cattle.

Authors: Sarah C. Helgeson & Sheila M. Schmutz
Department of Animal and Poultry Science, University of Saskatchewan, Saskatoon, Canada

Abstract:
Pro-melanin-concentrating hormone (PMCH) had previously been shown to affect appetite and metabolism in rodent species. PMCH was mapped to BTA5 (Stone et al. 2002) and QTL to backfat were reported for this chromosomal region (Casas et al. 2000, Li et al. 2004). The objective of this research was to examine whether there were similar effects in cattle. Sequencing of Bos taurus PMCH from several prevalent North American beef breeds revealed an adenosine-to-thymine single nucleotide polymorphism (SNP) upstream of the transcriptional start site. This SNP was subsequently shown to significantly affect both average fat and grade fat levels in two populations of beef cattle. One of these populations had tenderness data. Consumer taste panel evaluations indicated that the thymine allele contributed to decreased steak tenderness, as measured by a consumer taste panel and mechanical shear force, in the one population where these data had been measured. In silico analysis of the promoter region indicated that the presence of the thymine allele introduces a binding site for a transcriptional repressor, E4BP4. It is believed that the introduction of the thymine allele leads to downregulation of PMCH, which would increase physiological metabolic rates, resulting in tougher steaks and decreased fat production.

Poster 2160

Title: Using a 1.5K ovine SNP array to expand the sheep linkage map

Jillian F. Maddox¹, Allan M. Crawford², Roxann Ingersoll³, James Kijas⁴, Brian P. Dalrymple⁵, Ian R.W. Evans⁶, Frank W. Nicholas⁷, John McEwan⁸, Tracey van Stijn⁹, Herman Raadsma⁹ and the International Sheep Genomics Consortium⁹.

¹Department of Veterinary Science, University of Melbourne, Victoria 3010, Australia
²AgResearch, Invermay Agricultural Centre, Mosgiel, New Zealand
³Johns Hopkins University, Institute of Genetic Medicine, Baltimore MD, USA
⁴CSIRO Livestock Industries, Brisbane, Queensland 4067, Australia
⁵irwevans@bigpond.net.au
⁶Reprogen, University of Sydney, New South Wales 2006, Australia
⁷http://www.sheephapmap.org/isgc_participants.htm

Abstract
The sheep linkage reference map has been developed by genotyping a 3-generation, full-sibling pedigree, the International Mapping Flock (IMF), produced by AgResearch that consists of 127 sheep in 9 families. Version 4.7 of the map, released in December 2006, comprised 1,425 markers representing 1,381 loci and spanned approximately 3,600 cM. The majority of mapped markers (87%) were microsatellites, and 51% were derived from markers initially developed for cattle. The International Sheep Genomics Consortium (ISGC) recently identified 6,022 ovine SNPs from a targeted BAC end resequencing project and public ESTs. Of these, 1,536 SNPs were selected for genotyping, based on their predicted chromosome positions, with Illumina’s BeadArray platform. SNPs were selected to represent every ovine chromosome (OAR) with an average of 56 SNPs per chromosome ranging from one SNP on OARY to 151 SNPs on OAR1. The 1.5K array was used to genotype the IMF so that the SNPs could be integrated into the linkage map. The CRI-MAP and MultiMap programs were modified so that they could better handle a larger volume of genotyping data. Construction of version 5 of the linkage map is currently underway and the new map will contain approximately 2,530 markers including 1,113 SNPs from the array.

Poster 2161

Title: Expression of the chicken β2-crystallin gene: Occurrence of sense and antisense RNAs in testis.

Presenting Author: Takeshi SHIMOGIRI, Korimoto 1-21-24, Kagoshima, 890-0065, JAPAN

Other authors (name only):
1. Junshi KAWAKAMI
2. Akiko MATSUMOTO
3. Masahide NISHIBORI
4. Shin OKAMOTO
5. Yoshizane MAEDA
6. Hiroshi YASUE

Abstract:
Crystallin, the major eye lens protein in birds, is homologous to the urea cycle enzyme argininosuccinate lyase (ASL). In chicken, there are two δ-crystallins, denoted δ1 and δ2. Of them, δ2-crystallin gene (CRYD2) is the chicken ASL ortholog because only δ2 has retained the ASL activity. Birds are uricotelic in terms of nitrogen excretion and apparently do not require the urea cycle for nitrogen excretion. Here, to examine how and why the urea cycle genes have been maintained in chicken, we examined the CRYD2 expression in chicken tissues by RT-PCR. We detected the CRYD2 fragment in various chicken tissues using cDNA prepared with random hexamers and PCR. These results revealed the presence of the CRYD2 mRNA in all tissues and two additional transcripts in testis. Chicken BLAT search indicated that exon/intron structure was different amongst the CRYD2 mRNA and two testis-specific transcripts. Additionally, we found that two testis-specific transcripts were natural antisense RNAs of the CRYD2 gene: the testis 1st-strand cDNA prepared with gene-specific forward primer produced the two PCR products corresponding to the testis-specific transcripts by the subsequent PCR reaction, while the testis 1st-strand cDNA prepared with gene-specific reverse primer produced the PCR products from the CRYD2 mRNA.

Poster 2162
Title: Linkage disequilibrium mapping of QTL on chromosome 7 affecting female fertility and twinning rate in Israeli Holstein cattle
Presenting Author: Ron M., The Volcani Research Center, P.O.B. 6, Bet-Dagan 50-250, Israel
Other authors (name only):
1. Glick G
2. Golik M.
3. Shirag A
4. Seroussi E
5. Weller J.I

Abstract:
Two segregating quantitative trait loci (QTL) affecting female fertility, defined as: 100/(number of inseminations to conception), were detected on BTA7 in the Israeli Holstein population by a daughter design analysis. One QTL was located near the centromere, and the other in the vicinity of 40 Mbp from the centromere. In order to confirm and fine map these QTL, linkage disequilibrium between SNPs and the putative QTL was analyzed by the regression of the genetic evaluations of 650 bulls on their marker genotypes. DNA samples were genotyped for 206 SNPs located within 43 Mbp from the centromere, using IPLEX technology. Two markers, located near position 42 Mbp had significant effects on female fertility (comparison-wise p < 10^-4), and a marker near position 40 Mbp had a significant effect on twinning rate (comparison-wise p < 10^-5). Thus the QTL near position 40 Mbp was confirmed, but not the QTL near the centromere. The allelic substitution effects on female fertility and twinning rate were 1.05% and 0.71%, respectively. Previous results have shown a negative genetic correlation between twinning rate and fertility, and this was confirmed for all eight SNPs with nominally significant effects on both traits.
Poster 2164

**Title:** Evaluation of the genetic structure of the Brazilian Fila dog with microsatellites

Presenting Author: Aníbal Souza Felipe da Silva, Rua Rio Grande do Sul, n.646, apto.703, Barro Preto – Belo Horizonte – MG / CEP: 30170-110

Other authors (name only):
1. Marcelo Yukio Kuabara
2. Eduardo Geraldo Alves Coelho
3. Daniel Cardoso de Carvalho
4. Cláudia Salviano Teixeira
5. Denise Aparecida Andrade Oliveira

**Abstract:**
The Fila dog is the most important molosser breed in Brazil for its guard abilities, loyalty and its historical origin associated to the development of the Brazilian society and culture. But there is a lack of studies about its genetic structure. Using the ISAG recommended markers of 21 microsatellites, 26 non-related dogs were genotyped. All microsatellites were polymorphic, with a number of alleles ranging around 3 (AHTk253, INU030, REN162C04) and 8 (AHT121). Allele frequencies, inbreeding index (F), effective number of alleles (Ne), observed (Ho) and expected heterozygosity (He), polymorphic information content (PIC), parentage exclusion probability when both parents are typed (PE-2) and matching probability of two unrelated individuals (MPR), as well as Hardy-Weinberg genetic equilibrium were estimated. Mean Ho was 0.598 (ranging from 0.385 at AHTk211 and AHTH130 to 0.808 at REN105L03. Two loci (AHTh171 and INU005) were out of HW equilibrium (p<0.01). The PIC value for the set of 21 microsatellites was 0.56. The PE-2 reached 0.9999. This study contributed to elucidate the genetic structure of the Brazilian Fila dog breed and to orientate conserving programs, in order to avoid high levels of endogamy, specially when associated to reproductive biotechnologies. Thanks: GenLab/UFMG Veterinary School and CAFIB Fila Brasileiro breeders.

Poster 2165

**Title:** Characterization of PrP gene polymorphism in South Croatian sheep populations

Presenting Author: Vlatka Cubric Curik, Faculty of Agriculture, University of Zagreb, Department of Animal Science, Svetosimunska 25, 10000 Zagreb.

Other authors (name only):
1. Marcelo Yukio Kuabara
2. Eduardo Geraldo Alves Coelho
3. Daniel Cardoso de Carvalho
4. Cláudia Salviano Teixeira
5. Denise Aparecida Andrade Oliveira

**Abstract:**
Polymorphism at codons 136, 154 and 171 in the ovine prion gene (PrP) influence the progress and incidence of scrapie and bovine spongiform encephalopathy (BSE). The European Union legislation promotes breeding programs that favour genetic resistance to scrapie. In Croatia, EU Candidate country, the genetic polymorphism in sheep populations has never been studied. Here, we present first results of the polymorphism at codons 136, 154 and 171 in six Croatian sheep breeds/populations (Cres Island sheep, Rab Island sheep, Pag Island sheep, Brac Island sheep, Dalmatian Pramenka and Dubrovnik Ruda), each sampled over more than 50 individuals. The preliminary results (estimates) have shown that three haplotypes; ARQ (> 50%), ARR and AHQ were the most common haplotypes in the Croatian meta-population, although, variability among breeds/populations have been observed. The estimated frequency of VRQ haplotype has been very low. Still, at some breeds/populations the frequency of “non-resistant” genotypes (ARQ/VRQ) was not negligible (> 5%) indicating that genetic resistance for polymorphism at codons 136, 154 and 171 is probably not the reason why scrapie has never been reported in Croatia.

Poster 2166

**Title:** Gene expression profiles in CNS of naturally scrapie-affected sheep

Presenting Author: Inmaculada Martín-Burriel, Laboratorio de Genética Bioquímica. Universidad de Zaragoza. Miguel Servet 177. 50013 Zaragoza (Spain)

Other authors (name only):
1. Carmen Serrano
2. Jaber Lyahyai
3. Rosa Bolea
4. Eva Monleón
5. Frank Harders
6. Ane Marcos-Carcavilla
7. Juan J. Badiola
8. Alex Bossers
9. Pilar Zaragoza

**Abstract:**
Transmissible spongiform encephalopathies (TSEs) are fatal neurodegenerative disorders caused by prions. Naturally scrapie-infected sheep can be considered as a natural animal model of these diseases. The underlying mechanisms of scrapie pathogenesis are still poorly understood. The identification of genes with differential expression in
CNS (central nervous system) of infected animals might provide clues to clarify the molecular mechanisms that lead to neuronal loss and to identify molecular biomarkers that might be the basis for new diagnostic tests. We present here an initial expression profiles of 18 genes involved in the regulation of apoptosis and in the response of cells to stress (HSPs) using Real Time RT-PCR. This study has been carried out in four CNS regions that display different lesion degrees. We have detected the up-regulation of pro-apoptotic genes and down-regulation of anti-apoptotic genes in brain stem (the most affected area). However, we have also observed the induction of antiapoptotic genes suggesting that neuroprotective mechanisms could counteract the effect of pro-apoptotic stimuli. Moreover, we report here an initial study on the transcriptional differences in cerebellum using cDNA microarray hybridizations. Small sequences of GMPS, RPL32, ACTG1 and ATP6AP2 genes align with up-expressed clones. By contrast, GNB2L1, HSPA8, RPS3 and FN1 genes have similarity with down-regulated sequences.

**Poster 2167**

**Title: The pig as a biomedical model for melanoma: identification of MITF as a potential predisposing gene**

Presenting Author: Emmanuelle BOURNEUF 1, 2

1 INRA DGA, UMR 314 Laboratoire de Radiobiologie et Etude du Génome, Jouy-en-Josas, France

2 CEA, DSV, iRCM, SREIT, LREG, Jouy-en-Josas, France

Abstract:
The MeLiM strain is a porcine model of cutaneous melanoma characterized by the spontaneous development of tumours in utero or during the first weeks of life. Clinical and histopathological observations have shown a high similarity between porcine and human lesions. A cross between MeLiM and healthy pigs was used to perform a QTL analysis, that mapped several susceptibility regions to swine chromosomes 1 (SSC1), SSC2, SSC13, SSC15 and SSC17. The most significant QTL (p<0.01), located on SSC13, is partly orthologous to HSA 3p14, and harbours the MITF gene, a master regulator of melanocyte biology. MITF amplification has been shown in human melanoma, and thus represents a particularly interesting candidate gene for melanoma predisposition in our model. We used BAC-end sequences to detect new microsatellites within the SSC13 QTL, and genotyped 15 on our backcross pedigree. The results show a refinement of the QTL peak above MITF, and an association between a close marker and melanoma development. We are currently analysing a panel of SNPs spread over 1Mbp around the gene to confirm this association. Ongoing functional studies have already shown a subtle regulation of MITF isoforms in melanocytes and tumors.

**Poster 2168**

**Title: Mapping and expression studies of the DMRT1 gene in domestic pig**

Presenting Author: Ewa Slota, Department of Immuno-and Cytogenetics of Animals, National Research Institute of Animal Production, Krakowska 1, 32-083 Balice n. Kraków, Poland

Abstract:
*Doublesex and mab-3 related transcription factor 1 (DMRT1)* acts during animal sexual development and -in the majority of vertebrates- its function is limited to the upstream or downstream testis regulators. Porcine radiation hybrid panel (IMpRH) was used for physical mapping. PCR was carried out using two primer pairs located in 5'-flanking (AF426435) and 5'-UTR region (AF216651) resulting in PCR fragments of 399 and 122 bp respectively. PCR signals of hybrid clones were scored, evaluated with the IMpRH mapping tools (http://imprh.toulouse.inra.fr). For both examined sequences the closest linked marker is SW1621 (LOD score 5.66) on SSC1 (RH Map 4442 centiRay). Expression of swine DMRT1 was analyzed by RT-PCR using dT-primers for RT and PCR-primers located on different exons (AF216651). Total RNA was isolated from various tissues. As a positive control all the cDNA samples were amplified with beta-actin primers (AJ312193). The highest DMRT1 expression was detected in testis cDNA and followed unexpectedly by ovaries, kidney, heart and muscle. No amplicons were detectable in brain, liver, lung and spleen. The present study of porcine DMRT1 gene confirms its autosomal status and its chromosomal homology to humans, but reveals differences in expression pattern in adult tissues which is not limited only to testis.
**Poster 2169**

**Title:** The Ile-442-Met substitution in NCAPG as a positional candidate for bovine carcass weight QTL (CW-2) on chromosome 6


**Abstract:**
We previously mapped carcass and body weight QTL (CW-2) on BTA 6 using Japanese Black Sire A’s half-sib family. The CW-2 QTL was replicated in Sire B that harbored the same Q haplotype as Sire A. We also mapped a body or carcass weight QTL in Sire C that harbored a different Q haplotype, and Japanese Brown Sires D and E that shared another Q haplotype, in the overlapped regions with CW-2. Assuming that these QTL were the same, we searched for a shared Q region using 39 microsatellite markers spanning the CW-2 region (38-55 cM), and performed linkage disequilibrium mapping using maternal alleles of Sire C and Sire D offspring. An 810-kb region was shared among the three Q haplotypes and associated with body and carcass weight. By adding 42 SNP markers, the region was refined to 660 kb containing 4 genes. The SNP changing Ile-442 to Met in NCAPG (chromosome condensation protein G) was heterozygous in all the 5 sires and highly associated with carcass weight (p < 1.2 x 10^-11) in a large Japanese Black population (187 steers selected from the 4.7% extremes of 7990 steers). This SNP provides a CW-2 marker as a positional candidate QTN.

**Poster 2170**

**Title:** Biomarker development for pork quality using SELDI-TOF proteomics technology

Presenting Author: Henri C.M. Heuven


**Abstract:**
Development of meat quality relates to rate and status of the degradation of the muscle proteome after slaughter. Therefore, protein breakdown quantified using SELDI-TOF might identify good predictors of meat quality. SELDI-TOF technology, with its high throughput capability, allows for faster analysis of many samples permitting its use in explorative research of biomarkers. Knowledge regarding the progress of proteolysis on the entire proteome of muscle tissue is limited and breed-specific differences have received little attention. The objectives therefore were to investigate breakdown profiles of the m. longissimus dorsi proteome and the repeatability of the proteomic measurements. Meat quality and proteome profiles (in duplicate) of 5 Yorkshire and 5 Duroc pigs and proteome profiles were determined at slaughter and after 1, 2, 3, 7, and 10 days of ageing. The proteome showed four degradation profiles for different peptides. Association analysis suggested several biomarkers for drip loss and shearforce, but none for cooking loss. The repeatability between duplicated SELDI-TOF measurements ranged from 0.38 to 0.82 depending on the matrix and on the relative amount of the protein in the sample. For biomarkers development, samples should be taken at least in duplicate to minimize chance of obtaining false positive markers.

**Poster 2171**

**Title:** Selecting cattle for host resistance to tick (Rhipicephalus microplus) infestation – natural and human-directed selection

Presenting Author: Nicholas N Jonsson


**Abstract:**
Two taurine populations of cattle with high resistance to tick infestation exist in Australia: the Adaptaur, selected by CSIRO from Hereford and Shorthorn cattle for resistance to ticks, and the Kimberley Shorthorn, a nearly feral descendent of original cattle introduced into Australia over 100 years ago and selected naturally to survive with ticks and without insecticide. We genotyped 58 Adaptaur and 94 Kimberley Shorthorns from tick-infested locations, 67 Shorthorn, 65 Hereford from non-infested areas, and 60 Brahman cattle, using 9 microsatellite markers from our parentage testing panel and 9 markers for...
known genes within the MHC. Mean allele number in the MHC differed: Adaptas 5.6±0.34, Herefords 7.3±0.91, Shorthorn 9.2±0.12, Brahman 11.1±1.3 and Kimberley Shorthorn 12.6±1.6. Linkage disequilibrium was high in the MHC in all breeds, particularly for CYP21 and DRB3. PIC values for both these loci in the Kimberley Shorthorn were 0.90 compared with overall mean PIC of 0.68. Our results suggest that directed selection for high host resistance led to a population with low diversity and likely high susceptibility to other diseases (Adaptaurs), while natural selection and gene flow from other breeds led to cattle with high resistance and high levels of variability in the MHC (Kimberley Shorthorn).

Poster 2172

Title: Characterization of ORF and LTRs of Porcine Endogenous Retroviruses in pigs.

Presenting Author: Kie-Chul Jung, Division of Animal Science and Resources, Chungnam National University, Daejeon 305-764, Korea

Other authors (name only):
1. W. Y. Jung
2. E. W. Park
3. J. T. Jeon
4. C. Moran

Abstract:
Porcine endogenous retroviruses are one of serious problem regarding xenotransplantation because PERVs can infect human cells in vitro. Previously, PERVs have 30 to 50 copies in the pig genome and are classified as PERV-A, -B and –C based on the receptor-binding domain of the ENV gene. The PERV positive BAC clones were screened by PCR from the Korean native pig BAC library and 11 non-redundant BAC clones were selected for full-length PERV sequencing. Also, sequences from PERV flanking regions and chromosomal locations were investigated. The ORF analysis showed that only one PERV has complete ORF structure and the others have premature stop codons and INDEL mutations in gag, pol and env genes, indicating they are non-functional. Phylogenetic analysis of PERV LTRs was also performed and the preliminary results indicated that some of the LTR sequences are distinct from others. The results presented here can be a valuable information for generating PERV negative pigs which can be used as the xenotransplantation donor.

Poster 2173

Title: Polymorphisms in MYOD Gene Family Affect Carcass Traits in Cattle

Presenting Author: Mohammad Shamsul Alam Bhuiyan, Division of Animal Science and Resources, Chungnam National University, Daejeon 305-764, Korea;

Other authors (name only):
1. Seong Lan Yu
2. Yong Min Cho
3. Eung Woo Park
4. Jun Heon Lee

Abstract:
Myogenic differentiation genes (MYOD) are skeletal muscle specific transcription factors that encode highly conserved basic helix-loop-helix proteins. The MYOD family consists of four related genes; MYOD1 (MYF3), MYOG (MYF4, myogenin), MYF5 and MYF6. These genes play a key role in growth and muscle development in farm animals and are therefore considered as candidate genes for meat production and quality related traits. Polymorphisms were detected and confirmed by PCR using 17 primer pairs, sequencing and restriction fragment length polymorphism (RFLP) method. Fourteen SNPs (4 from MYOD1, 4 from MYOG, 3 from MYF5 and 3 from MYF6) were identified both in intron and exon of these genes, whereas 2 of them were nonsense mutations. Allele frequencies for g.783G>A and g.1274A>G mutations in MYOD1; g.511G>C and g.1111C>G in MYOG; g.1911A>G in MYF5 and g.183T>C in MYF6 varied significantly among the 7 different cattle breeds. The g.783G>A genotype of MYOD1 gene was significantly associated with carcass weight (P<0.05) and eye muscle area (P<0.001) in Korean cattle population. The g.1111C>G and g.1911A>G SNPs in MYOG and MYF5 genes, respectively, had significant association with carcass weight (P<0.01) and lean weight (P<0.01). Moreover, the g.1111C>G polymorphism in MYOG gene was associated with back fat thickness (P<0.001). In conclusion, the SNPs g.783G>A, g.1111C>G and g.1911A>G of MYOD gene family may be useful for marker assisted selection on carcass and meat quality traits in cattle.

Poster 2174

Title: Distinctive patterns of IGF2 and IGF2R expression in Bos indicus, Bos taurus and hybrid fetuses

Presenting Author: Carolyn Fitzsimmons, University of Adelaide, School of Agriculture, Food, & Wine. Roseworthy Campus, Roseworthy, SA 5371 Australia

Other authors (name only):
1. Radmilla Feldman
2. Sarah Truran
3. Zbigniew Kruk
4. David Lines
5. David Rutley
6. Stefan Hiendleder

Abstract:
We have investigated transcript-specific IGF2 and IGF2R expression in brain, cotyledon, liver and semitendinosus muscle of B. indicus, B. taurus, and reciprocal cross fetuses (n=40). Real-time qPCR data were analysed (GLM, SPSS) including fixed effects dam breed, sire breed, fetal sex, and covariate dam weight (i.e., four factors) and 3 housekeeping genes. Significant (P<0.05) effects on IGF2 transcripts were similar within each fetal tissue. Muscle IGF2 exons 2, 6, and 10, were affected by dam weight, and exon 10 by fetus sex. Liver IGF2 exon 10 was affected by the interaction of sire breed × dam weight. Cotyledon IGF2 exon 6 was affected by the interaction of dam breed × dam weight × fetus sex; exon 10 by an interaction between all four factors. No effects were significant on exon 10 expression in brain. The expression of IGF2R in brain, liver, and cotyledon, but not muscle, displayed dissimilar patterns than IGF2. Interactions between sire breed × fetal sex, and dam breed × dam weight, were detected in brain. None of the factors affected IGF2R expression levels in cotyledon or liver. Dam weight again affected IGF2R in muscle. These data demonstrate the complexity of imprinted gene expression in fetal development.

Poster 2175

Title: Partial duplication of the PRLR and SPEF2 genes at the late feathering locus in chicken

Presenting Author: Martin G. Elferink, P.O. Box 338, Wageningen, The Netherlands

Other authors (name only):
1. Amélie A.A. Vallée
2. Annemieke P. Jungerius
3. Richard P.M.A. Crooijmans
4. Martien A.M. Groenen

Abstract:
One of the loci responsible of feather development in chickens is called K. The K allele is associated with the late feathering phenotype that causes a retard in emergence of primary and secondary flight feathers. The k+ allele is associated with the early feathering phenotype resulting in the earliest emergence of feathers. The K locus is located on the Z chromosome and can be used to produce phenotypes that allow distinction between the sexes of chicks at hatch. This sexing method based on differences in rate of feather growth provides a convenient and inexpensive approach.

Previous studies on the organization of the K allele concluded the integration of endogenous retrovirus 21 into one of two large homologous segments located on the Z chromosome of late feathering chickens. In this study, the K locus was investigated with quantitative PCR by examining copy number variations in a total of fourteen markers surrounding the ev21 integration site. The results showed a duplication at the K allele and sequence analysis of the breakpoint junction indicated a tandem duplication of 176,324 basepairs. The tandem duplication of this region results in the partial duplication of two genes; the prolactin receptor and the gene encoding sperm flagellar protein 2.

Poster 2176

Using image analysis in a Jersey – Limousin double backcross to identify candidate genes for fat deposition traits in cattle

A.R. Egarr1,2, W.S. Pitchford1,2, M.J. Bottema3 and C.D.K. Bottema1,2
1Discipline of Agricultural and Animal Science, The University of Adelaide, Roseworthy, S.A. 5371
2CRC for Beef Genetic Technologies
3School of Informatics and Engineering, Flinders University, Bedford Park, S.A. 5042

Internal, intramuscular and subcutaneous fat deposition traits were mapped previously in a Limousin-Jersey double backcross cattle herd and several quantitative trait loci (QTL) identified. However, in order to quantify seam fat as a measure of intramuscular fat and to better describe intramuscular or marbling fat, a method was developed to analyse photographs of the L. dorsi muscle from the QTL mapping progeny. Using Adobe Photoshop® and Matlab® software designed for the photographic image analysis, seam fat areas were quantified and new marbling parameters were determined. QTL were then genetically mapped from the image analysis data. Two significant QTL of large effect were found on BTA2 and BTA19 for seam fat. Myostatin was an obvious candidate gene for the seam fat QTL on BTA2. Subsequent genotype analyses of the myostatin gene indicated that the Limousin F94L myostatin DNA variant was responsible for this QTL. For the seam fat QTL on BTA19, several candidate genes have been selected for sequencing to identify polymorphisms for association studies. In addition, the seam fat and marbling parameter measurements are being incorporated into fat deposition models to provide more informative and specific QTL for adipose partitioning.
Poster 2177

Title: Sequence of mitochondrial gene 12SrRNA in Emys orbicularis orbicularis population from Poland

Presenting Author: Beata Prusak, Institute of Genetics and Animal Breeding, Polish Academy of Sciences, Jastrzębiec, Poland

Other authors (name only):
1. Bartłomiej Najbar,
2. Sławomir Mitrus,
3. Tomasz Grzybowski,
4. Adam Hryniewicz,
5. Grzegorz Górecki,
6. Robert Bochen,

Abstract:
From the last morph metric observations conducted in Europe it results that Polish population of pond turtle is the last one relatively plentiful group of nominative subspecies of Emys orbicularis orbicularis (Linnaeus 1758) with presumably two ecological lines existing (“eastern” and “western”) differing morphologically and genetically. According the fact that pond turtle is morphologically varied, Emys orbicularis is probably the most divided vertebrate species in North Paleartic with several genetic lines existing in Europe. Aim of the research is to determine the genetic diversity of pond turtle within and between populations from different regions of Poland, to determine reference sequences of mtDNA for populations existing in Poland and to verify hypothesis about distinctiveness of domestic eastern and western E. orbicularis populations. From the research, it results that pond turtles from western Poland are genetically closer related to Spanish population than to Polish eastern one. The material covered buccal swabs possessed from turtles from four main regions of the population living-grounds in Poland: Wlodawa Lake District, Radom environs, Mazurian Lake District and Wielkopolska-Kujavian Lowland. Polymorphism of mitochondrial 12SrRNA gene was analyzed. Results presented are first step of wider research project covering coding and non-coding regions of pond turtle’ mitochondrial genome.

Poster 2178

Title: Disease gene identification in dogs: use of SNP arrays and candidate genes

Presenting Author: Alan N Wilton, School of Biotechnology and Biomolecular Sciences, Ramaciotti Centre for Gene Function Analysis, University of New South Wales, 2052 Sydney, Australia

Other authors (name only):
1. Jeremy R Shearman

Abstract:
We have identified mutations for a storage disease (NCL) which causes nerve degeneration and a neutrophil dysfunction (TNS) that results in high susceptibility to infections. TNS (Trapped Neutrophil Syndrome) is common in border collies on all continents. Both diseases have homologous human diseases. High throughput SNP arrays now make it easier to locate disease genes in inbred dog populations. Whole genome analysis on affected dogs and controls (around 10 of each) could be used in association studies and homozygosity analysis. Diseases where the mutations are identical-by-descent will be homozygous in the region of the disease gene in affecteds. The method is effective in cases where we have already identified the mutation. The application to cerebellar abiotrophy that causes ataxia in kelpies. seems simple in theory but is proving difficult in practice. Reliability of genotype calls and errors in assigning disease phenotypes are possible reasons for the lack of significant associations. A combination of homozygosity analysis and comparative genomics could also be used to identify disease genes from a single affected dog by eliminating large regions of the genome as candidate regions. We are applying this approach to a rare second type of NCL in broder collies.

Poster 2179

Title: Polymorphism and mapping of the porcine inhibitor of differentiation genes, ID1, ID2, ID3 and ID4

Antonín Stratil1, Jitka Filkuková1, Pavel Horák1, Mario Van Poucke2, Heinz Bartenschlager3, Luc J. Peelman2, Hermann Geldermann3

1Institute of Animal Physiology and Genetics, AS CR, Libechov, Czech Republic
2Faculty of Veterinary Medicine, Ghent University, Merelbeke, Belgium
3Department of Animal Breeding and Biotechnology, University of Hohenheim, Stuttgart, Germany

Abstract:
ID proteins are important members of signaling pathways involved in development, cell cycle and tumorigenesis. They are considered negative regulators of MRFs and the responsible genes are therefore candidates for carcass and meat quality traits. Specific PCR primers for the four porcine ID genes (ID1, ID2, ID3 and ID4) were designed and the PCR fragments were comparatively sequenced (Meishan and Pietrain) to verify the genes and identify polymorphisms. The genes were mapped using
radiation hybrid (IMpRH) panel and by linkage analysis in the Hohenheim Meishan x Pietrain F$_2$ family. The results of RH mapping were: ID1 – SSC17, closest marker ENDO3 (21 cR; LOD=11.11); ID2 – SSC3, closest marker SW2532 (20 cR; LOD=13.88); ID3 – SSC6, closest marker SW709 (8.0 cR; LOD=16.10); ID4 – SSC7, closest marker SW1369 (65 cR; LOD=4.64). By linkage analysis only ID3 and ID4 genes were mapped (in ID1 and ID2 no suitable polymorphism was found). ID3 was mapped to SSC6 (…FABP3 – 3.5 cM – ID3 – 14.2 cM – S0146….) and ID4 to SSC7 (…S0064 – 27.0 cM – ID4 – 12.1 cM – CYP21A2….). The ID4 gene maps to chromosome region with numerous QTL for carcass and meat quality traits. For this locus and for ID3 significant associations between genotypes and a series of traits were found (GLM procedure). (Supported by the Czech Science Foundation 523/06/1302)

Poster 2180

Title: GENETIC RELATIONSHIP BETWEEN ANDALUSIAN AND AMERICAN HORSE BREEDS

Presenting Author: Alexandra Gómez Tarazona
BIOTECGEN Calle 103A N 21-49 Bogotá. Colombia

Other authors (name only):
1. Cecilia Penedo
2. Fernando Ariza Botero
3. Andres Vera
4. Manuela Jay

Abstract:
Microsatellite polymorphisms were used to assess both the genetic diversity and the relationship between Spanish, South American and American Paso Fino horse breeds. Allelic characterization was made with 35 molecular microsatellite markers in order to do further population genetics studies. An amount of 387 animals unrelated registered representing each breed; Andalusian, Colombian horse, Peruvian and Paso Fino American horse breeds were analyzed. American horse populations displayed a relatively high level of genetic variation as estimated by allelic diversity and heterozygosity, whereas the Spanish breed showed reduced levels of genetic diversity. Only six percent of the total genetic variability could be attributed to differences among breeds (F$_{ST}$=0.06). American breeds clearly formed a separate cluster from the Andalusian breed. In addition, the molecular variance analysis indicated that 5.4% of variance could be explained by differences between Andalusian and New world breeds.
Abstract:
A genome scan was performed to identify quantitative trait loci (QTL) for short and medium chain FA*. Seven half-sib families with in total 849 cows and their 7 sires were genotyped for 1,379 SNP. Phenotypes of the cows were pre-corrected for systematic effects. The daughter design was used for a QTL program with multi-marker interval mapping in a weighted across-family QTL regression. Thresholds and confidence intervals were set using 10,000 permutations and 1,000 bootstraps. The genetic map measured 2,831 cM and had an average information content of 0.83. The results showed significant QTL (P<0.05) for 15 traits at 4 distinct chromosomal regions. Two of these QTL regions mapped to the approximate locations of the DGAT1 and the SCD1 gene. Correcting phenotypes for these two major genes resulted in complete disappearance of the linkage previously found in the two regions. Furthermore, suggestive QTL (P<0.05) were found for 28 traits. The identified QTL may be a first step in understanding physiology of milk fat synthesis, as well as providing the basis for the development of a genomic selection tool to decrease the proportion of the unfavorable fatty acids c14:0 and c16:0.

* c4:0, c6:0, c8:0, c10:0, c12:0, c14:0, c16:0, c10:1, c12:1, c14:1, c16:1, c10index, c12index, c14index, c16index

Poster 2183

Title: Identification of differentially expressed genes in adipocyte differentiation using Affymetrix Bovine Genome Array

Presenting Author: Seong-Lan Yu, Department of Animal Science, College of Agriculture and Life Science, Chungnam national university, 220, Gung Dong, Yuseong Gu, Daejeon 305-764, South Korea.

Other authors (name only):
1. Sang Mi Lee.
2. Man Jong Kang
3. Hang Jin Jeong
4. Byung Chan Sang
5. Seung Heui Ryoo
6. Jun Heon Lee

Abstract:
Adipocytes are differentiated from preadipocytes and have large capacity for storing fats inside cells. In cattle, intramuscular fat (IMF) content is one of the major determinants for meat quality and also highly affects market prices, especially in Japan and Korea. In order to profiling differentially expressed genes between intramuscular fibroblast-like cells (preadipocyte cells) and their differentiated adipocytes, we have been established intramuscular fibroblast-like cells from M. longissimus thoracis in Korean cattle (Hanwoo). The differentially expressed genes were selected by comparing these two cells using commercially available 23k Affymetrix Bovine Genome Array. The data analyses indicated that 206 array elements were up or down regulated. Of these, 38 and 57 known genes were up and down regulated, respectively, in adipocytes using both 2-fold difference and Welch’s T test as cut-off points. The differentially expressed genes identified in this study can be used as good candidate markers for improving meat quality traits, especially IMF content, in cattle.

Poster 2184

Title: Fine-mapping of a pea comb locus on chicken chromosome 1

Presenting Author: Shuji Sato, National Livestock Breeding Center, Nishigo, Fukushima, 961-8511, Japan

Other authors (name only):
1. Shinichi Sato
2. Tsuyoshi Otake
3. Junichi Saburi
4. Hiroko Hashimoto
5. Eiji Kobayashi

Abstract:
Although the Japanese game fowl (Shamo) is characterized by the pea comb (P/P or P/p), chickens with the single comb (p/p) often appear due to the presence of carrier (P/p). Our objective of this study was to narrow the genome region affecting the comb type to facilitate candidate gene identification. We used two populations for this purpose. First, an F2 resource population was generated by crossing Shamo and White Plymouth Rock (single comb, p/p). The number of F2 offspring was 265 from F1 intercross between eight males and 57 females. The comb type was segregating with the ratio of pea : single = 3:1 in F2 generation and we mapped a pea comb locus to a chromosomal region on Gallus gallus chromosome 1, flanked by microsatellite markers MCW0019, MCW0112 and ABR521. The second population was five generations population derived from cross between Shamo and Rhode Island Red (single comb, p/p), which was genotyped for additional polymorphic SNP and microsatellite markers within this region developed from Chicken draft sequences. To close some gaps in Chicken draft sequences, we constructed a BAC contig and sequenced it by the shot-gun sequencing. Chickens selected from some pedigree in these populations are grouped according to the inheritance of P or p haplotype at this locus constructed by additional markers. Finally, this locus was fine-mapped to roughly 50kb based on the association of the haplotypes and comb types. Chicken draft sequences suggest that sex determining region Y
–box 5 (SOX5) gene is the most likely candidate gene for the pea comb locus.

**Poster 2185**

**Title:** Functional analysis of the c.1892+19 T>C SNP of the PPARGC1A gene which is associated with variation in milk fat yield

Presenting Author: Rainer Fürbass, Research Institute for the Biology of Farm Animals (FBN), 18196 Dummerstorf, Germany

Other authors (name only):
1. Christa Kühn
2. Rosemarie Weikard

**Abstract:**
Recently, we identified the bovine peroxysome proliferator-activated receptor-γ coactivator-1α (PPARGC1A) gene as a plausible positional and functional candidate gene for a previously described QTL for milk fat yield on BTA6. A significant association between a SNP within intron 9 of PPARGC1A, c.1892+19 T>C, and milk fat yield was observed in a major dairy cattle population, indicating that the PPARGC1A gene could be involved in genetic variation underlying the QTL for milk fat synthesis on BTA6. A possible functional role of this SNP for PPARGC1A expression was investigated by electrophoretic mobility shift assays (EMSA) using allele-specific probes and nuclear protein extracts from various embryonic and adult bovine tissues. Probes including the C-allele of the SNP yielded a specific binding complex with all protein extracts. In contrast, no complex was observed with T-allele-specific probes. Hence, the C-allele represents a new binding site for a transcription factor. This finding supports the hypothesis, that the trait-associated SNP may have a functional effect. By competition and supershift experiments it could be demonstrated that the sequence motif is a target of the transcription factor Sp1. Currently, possible effects of the SNP on gene expression are being investigated by reporter gene analyses in various cell culture models.

**Poster 2186**

**Title:** Expression of antimicrobial peptides in different localisations of the bovine udder

Presenting Author: Jens Tetens, Institute of Animal Breeding and Husbandry, Olshausenstr. 40, D-24098 Kiel

Other authors (name only):
1. Judith Friedrich
2. Manfred Schwerin

**Abstract:**
Antimicrobial peptides (AMP) are key effectors of innate immunity and play an important role in local host defence. In order to enhance the knowledge about the role of bovine AMPs in the protection against udder infection we analysed the gene expression level of the six bovine AMPs LAP, TAP, DEFB1 (EBD), BNBD4/10, and S100A7 (psoriasin) in five localisations of the bovine udder, namely the streak canal, the rosette of Fürstenberg, the cisterna, the udder lymph node, and udder parenchyma. We analysed samples from nine healthy cows applying a quantitative real-time PCR approach. Consistent with the role of AMPs in local host defence, predominantly taking place at the sites of invasion, we found no or marginal expression of all genes in parenchyma. The highest expression levels for LAP and TAP were detected in the lymph node with remarkably high values for LAP. The neutrophil ß-defensins BNBD4/10 were predominantly expressed in the lymph node with BNBD10 also showing remarkably expression in the streak canal and cisterna. The S100A7 (psoriasin) gene which has been shown to be active against E. coli was found highly expressed in the streak canal and the cisterna, indicating a major role in the defence against coliform mastitis.

**Poster 2187**

**Title:** Characterization of BLV (Bovine leukemia virus) envelope gene in Holstein from Korea

Presenting Author: H. J. Jeong, Department of Animal Science, College of Agriculture and Life Science, Chungnam National University, 220, Gung Dong, Yuseong Gu, Daejeon 305-764, South Korea.

Other authors (name only):
1. S. L. Yu.
2. S. H. Ryoo
3. C. H. Do
4. B. C. Sang

**Abstract:**
Bovine leukemia virus (BLV) is an oncogenic virus predominantly present in cattle. It belongs to Delta-retrovirus in the family Retroviridae. The genomic structure is similar to human T-cell leukemia virus. BLV has envelope proteins which contain functional domains that determine its infectivity. We conducted a nested polymerase chain reaction (PCR) for amplification of the envelope gene which can be act for the serological status in the host. To examine the distribution of BLV variants, sequencing of the PCR product was performed. The nucleotide sequences were compared with other previously characterized BLV sequence from different serological areas. We
compared sequences of Korea Holstein population and BLV env sequences available at GenBank. The phylogenetic tree showed three different clusters and most samples from Korea population formed one cluster similar in USA Holstein sequences. Further research about BLV give some clues in order to make general strategy for improving the disease.

Poster 2188

Title: Associations between polymorphisms in the cholecystokinin B receptor gene and behavioural traits in beef cattle

Presenting Author: K. Glenske, Department of Animal Breeding and Genetics, Justus Liebig University, Giessen, Germany

Other authors (name only):
1. M. Gauly
2. G. Erhardt

Abstract:
The importance of temperament and docility in successful management of beef cattle is already recognized as well as its genetic background. One candidate gene is the cholecystokinin B receptor (CCKBR), which plays a role in regulation of anxiety. The molecular analysis of two regions of bovine CCKBR, including exon III - exon IV and the 3' UTR, resulted in two SNPs. The SNP in intron 3 (C/T), which belongs to a binding site for a transcription factor, was genotyped by a PCR-RFLP with BseNI; the SNP (C/T) in the 3'UTR by a tetra-primer ARMS-PCR.

Behavioural traits related to temperament were scored in 962 calves (born in 1998-2001) from progeny groups of German Simmental (n = 8) and German Angus beef cattle (n = 6) in three standardized tests at different ages (tethering test, 3 weeks; weighing test, 3 month; separation- and restraint-test, 7-8 month). Behaviour scores were given to the animals. The association analysis was done for each SNP with a mixed linear model within breeds, including the four years and genotype as fixed effects, age at test as covariate and sire as random effect. No significant associations between the SNPs and the behavioural scores could be demonstrated.

Poster 2189

Title: QTL mapping of Japanese Black cattle using a family structure combining seven paternal half-sib families.

Presenting Author: TOSHIO WATANABE, Shirakawa Institute of Animal Genetics, 1 Odakurahara Odakura Nishigo Fukushima, Japan

Other authors (name only):
1. MARIO SEKIYA
2. YOICHI SATO
3. HIROKI NAKAJIMA
4. SHUNJI MARUTA
5. TOMOHITO ITOH
6. AKIKO TAKASUGA
7. YOSHIKAZU SUGIMOTO

Abstract:
For QTL analyses of livestock, it often takes long time to collect enough number of half-sib offspring to achieve reasonable statistical power. In this study, we examined plausibility to map QTL of a grandsire by combining several small families to construct a large three-generation family of Japanese Black cattle. We combined genotype data of seven half-sib families, where all the sires are sons of grandsire A, containing 44-103 offspring (426 in total) and genotyped with 185-436 microsatellites. Transmission of grandsire A’s haplotype to a sire was traced using 263 microsatellites. Three QTL-models were assumed: superior QTL allele is present (1) on the first haplotype of the grandsire, (2) on the second haplotype, and (3) on both. Probabilities to succeed the grandsire’s haplotype were linearly regressed on phenotypic values of 3rd generation offspring. Sixteen QTL were detected for beef marbling standard score, carcass weight, longissimus muscle area, and rib thickness (P<0.05, chromosome-wise significance level). Of sixteen, seven were newly mapped and nine were overlapped with the regions previously mapped in Japanese Black. We ensured that marbling QTL on BTA19 segregated within A’s 400 offspring. Combining small families will provide more opportunities to map novel QTL in livestock.

Poster 2190

Title: Two novel single nucleotide polymorphism markers of MC1R gene in Thai Native White Lamphun cattle

SUPAMIT MEKCHAY1,2
1Department of Animal Science, Faculty of Agriculture, Chiang Mai University, Chiang Mai, 50200 Thailand, 2Center of Agricultural Biotechnology, Faculty of Agriculture, Chiang Mai University, Chiang Mai, 50200 Thailand

Abstract
Melanocortin receptor 1 (MC1R) gene is an important gene in the regulation of pigment synthesis and responsible for coat color in mammals. The polymorphism of MC1R gene in Bos taurus has been characterized as four alleles (E2, E*, E3 and e). In this study was carried out to identify single nucleotide polymorphism (SNP) of MC1R gene in Thai native
cattle (*Bos indicus*). The coding region of *MC1R* gene in Thai native cattle namely White Lamphun cattle (white coat color) was sequenced and compared with Holstein and Charolais breeds. The *MC1R* sequence of White Lamphun cattle has high homology with the *Bos taurus* (99% identity). Four polymorphic sites were found in *MC1R* gene of the White Lamphun at position 296, 416, 663 and 725 bp of open reading frame. Out of these, three SNPs were identified as missense mutation, consisting of (1) a single base substitution (T296C) resulting in an amino acid change from leucine to proline (E allele) (2) a single base substitution (C416T) leading to an amino acid change from alanine to valine and (3) a single base substitution (A725C) leading to an amino acid change from asparagine to threonine. Moreover, a nonsynonymous mutation was located at position A663C of bovine *MC1R* coding region. Based on this observation, two novel SNPs at position 416 and 663) were found only White Lamphun cattle breed. This result indicated that the *MC1R* gene of White Lamphun cattle breed was E allele and these two novel SNPs may be used as allele specific markers for the White Lamphun cattle breed.

**Poster 2191**

**Title:** Expression of Non-Classical MHC Class I Genes by Bovine Embryos

CJ Davies¹, A Wilhelm¹, BR Sessions¹, P Parasar¹, Y Li², EE Schmidt², KL White¹

¹Department of Animal, Dairy and Veterinary Sciences & Center for Integrated BioSystems, Utah State University, Logan, UT, USA; ²Veterinary Molecular Biology, Montana State University, Bozeman, MT, USA.

**Abstract:** In human pregnancy trophoblast cells express the non-classical MHC class I (MHC-I) proteins HLA-G and HLA-E, which are important immunoregulatory factors. Previously, we showed that during the third trimester bovine trophoblast cells expressed four non-classical MHC-I genes, BoLA-NC1 through BoLA-NC4 (Davies et al. AJRI 55:188-200, 2006). Thus far, no one has studied expression of bovine non-classical MHC-I genes early in pregnancy. We have cultured bovine in vitro fertilized embryos to gestational age 30-37 days in either DMEM/F12 or bES medium plus 15% serum and used these embryos for analysis of MHC-I gene expression. Following RNA extraction from individual or pooled embryos with TRIzol Plus (Invitrogen), RNA was reverse transcribed with Superscript III (Invitrogen), cDNA amplified by PCR with MHC-I primers that amplify alleles encoded at all classical and non-classical MHC-I loci, cloned in the pCR4 vector (Invitrogen), and 40-48 clones sequenced by cycle sequencing using BigDye V3.1 and an ABI PRISM 3730 DNA Analyzer (Applied Biosystems). Embryos expressed both classical and non-classical MHC-I genes. Expression of non-classical MHC-I genes suggests that bovine trophoblast cells express these genes around the time of placental attachment, which occurs about day 30 of pregnancy. Expression of classical MHC-I genes could either be by trophoblast cells, which are the predominant cell type in these embryos, or by endothelial or other cells. For one pool of four day 37 embryos 33% of the pCR4 clones contained BoLA-NC1 sequences. BoLA-NC1 transcripts exhibit differential splicing involving intron 4 and exon 5, which encodes the transmembrane domain. The splicing patterns are similar to patterns seen with HLA-G. Consequently, BoLA-NC1 may be a functional homolog of HLA-G.

**Poster 2192**

**Title:** Predicting genetic merit for production and health traits in dairy cattle using genome wide selection and high density SNP screens

Presenting Author: Herman W. RAADSMA, Reprogen, The University of Sydney, Australia

Other authors (name only):
1. G. Moser
2. R. Crump
3. K.R. Zenger
4. M.S. Khatkar
5. J.A. Cavanagh
6. R.J. Hawken
7. J.Solkner
8. B.Tier

**Abstract:** A genome wide selection (GWS) platform was developed for prediction of genetic merit in dairy cattle. The critical components of the GWS platform included a genome wide SNP analysis assay representing between 15,000 and 70,000 SNP’s and 1945 progeny tested Holstein Friesian sires with EBV for 42 lactation performance traits. Two methods for complexity reduction and feature selection were used - Partial Least Squares (PLS) and regression using a genetic algorithm (GAR), to find optimal solutions of EBVs against SNP information. Extensive internal cross validation was used to find the best predictive models followed by external validation without direct use of pedigree information or SNP location. Derived Molecular Breeding Values (MBV) using either all or a fraction of the available SNP information were shown to have high predictive value for genetic merit (r=0.65-0.87 with EBV) when test subsets were randomly drawn form the data (mirror prediction). A general loss in accuracy was seen when genetic merit
was predicted in test co-horts of young bulls which were not part of the training set. However MBV was more accurate in predicting genetic merit than pedigree based EBV alone at time of birth for young bulls. PLS was computationally more efficient than GAR.

Poster 2193

**Title:** Positional cloning of the canine hairless mutation reveals an essential role for the **FOXI3** gene in ectodermal development

**Presenting Author:** Cord Drögemüller, Institute of Genetics, University of Bern, Switzerland

**Other authors (name only):**
1. Elinor Karlsson
2. Marjo K. Hyytönen
3. Michele Perloski
4. Gaudenz Dolf
5. Hannes Lohi
6. Kerstin Lindblad-Toh
7. Tosso Leeb

**Abstract:**
Chinese Crested, Mexican and Peruvian hairless dogs are characterized by a very sparse hair coat and deficient and abnormally shaped teeth, a phenotype summarized under the classification canine ectodermal dysplasia (CED). CED is inherited as a monogenic, autosomal, semi-dominant trait. Heterozygous dogs exhibit the CED phenotype and homozygous mutant animals die during embryonic development. We applied a whole-genome SNP association mapping approach using DNA from 20 hairless and 20 coated Chinese Crested dogs and the canine Affymetrix v2 SNP genotyping microarray. This analysis identified a single region of strong association on CFA17. This region contains a three SNP haplotype with complete phenotype concordance. Subsequent linkage disequilibrium fine mapping across three breeds narrowed the critical interval to 102 kb. This interval contains only one gene, a previously uncharacterized member of the forkhead box transcription factor family (**FOXI3**). Whole mount in situ hybridization of mouse embryos demonstrated a specific expression of Foxi3 in hair and whisker follicles and in developing teeth. Mutation analysis revealed a frameshift mutation within the **FOXI3** coding sequence that co-segregates with the CED phenotype in all three investigated dog breeds. Thus, using a two-stage, genome-wide mapping approach, we have efficiently identified a novel gene involved in ectodermal development.

Poster 2194

**Title:** High resolution chicken radiation hybrid maps using the Illumina™ technology

**Presenting Author:** Alain Vignal, UMR génétique Cellulaire, INRA Chemin de Borde-Rouge, Auzeville BP 52627, 31326 CASTANET-TOLOSAN CEDEX, FRANCE

**Other authors (name only):**
1. Caroline Hourcade
2. Thomas Faraut
3. Sophie Leroux
4. Marine Douaud
5. Suzanne Bardes
6. Frédérique Pitel
7. Mario Foglio
8. Ivo Gut
9. Mireille Morisson

**Abstract:**
The chicken genome sequence assembly is composed of more than 1 Gb of sequence covering almost all chromosomes, with the exception of a few microchromosomes. To confirm the existing assembly and to study the missing microchromosomes, we are developing high density radiation hybrid (RH) maps. Markers from the ChickRH database and 9212 markers genotyped on an Illumina platform were used and the construction of framework maps was done using the Carthagene software. Alignment of the maps to the sequence indicates a good agreement with only a few minor discrepancies, such as small inversions of a very limited number of markers, which can suggest an ordering error either in the sequence or in the RH maps. The resolution of the framework maps is 26 kb/cR with an average spacing of 410 kb between markers. Additional linkage groups could correspond to microchromosomes with very little genomic sequence information. These results indicate a good quality of the chicken sequence and confirm the possibility of using the Illumina™ technology for RH mapping, despite the fact that it was primarily designed for SNP genotyping.

Poster 2195

**Title** **PGK1** endogenous gene shows expression differences in porcine tissues.

**Presenting Author:** Ana Isabel Fernández. Dpto. Mejora Genética Animal. INIA. Ctra. Coruña Km7.5 28040 Madrid. Spain

**Other authors (name only):**
1. Amanda Fernández-Rodríguez
2. Almudena Fernández
3. M. Carmen Barragán
4. M. Carmen Rodríguez
5. Cristina Óvilo

Abstract:
Gene expression is quantified by normalization to the level of one or more reference genes. However different conditions might influence their expression, which could lead to errors in the interpretation of results. The Phosphoglycerate kinase 1 (PGK1) gene codes for a protein that participates in glucose metabolism and it is usually considered as reference gene. We conducted a gene expression analysis to identify hepatic gene expression patterns in male and female Iberian pigs fed with high and low feeding levels using microarrays. This study showed that PGK1 expression was conditional on feeding level. Here we have performed the validation of this result by qPCR in a larger number of liver samples (n=17) using as endogenous GADPH and BM2 genes and geNorm software. Statistical analysis model of the expression differences by qPCR included the sex, feeding level and their interaction effects. Statistical results show that PGK1 liver gene expression depends on sex (p=0.0003; females=1.5x males) and feeding level (p=0.0251; high= 1.23x low), and the interaction sex x feeding level is also significant (p=0.0215). In addition, we have detected similar effects in porcine psoas major muscle. These results would invalidate PGK1 as reference gene for these tissues in our pig population.

Poster 2196

Title: Detection of signature of selection among 14 cattle European and African populations in genomic regions associated to trypanotolerance

Presenting Author: Guiguibaza Kossigan DAYO, CIRDES, BP454 Bobo Dioulasso, Burkina Faso

Other authors (name only):
1. Sophie THEVENON
2. Katayoun MOAZAMI-GOUDARZI
3. Catherine DENIS
4. Andre EGGEN
5. Mathieu GAUTIER

Abstract:
Breeding of indigenous African taurine cattle tolerant to African trypanosomoses remains a straightforward alternative to control these diseases. Previous studies have identified QTL regions underlying trypanotolerance in experimental cross. In order to validate and refine these QTL in different cattle populations, we looked for signature of selection in surrounding regions using a panel of 92 microsatellite markers genotyped on 20-50 individuals belonging to 14 different cattle breeds. Breeds were chosen according to their history and their known status regarding tolerance to trypanosomoses. Hence, 4 West African trypanotolerant taurine breeds and 10 European and African trypanosusceptible cattle breeds were chosen. Two FST based tests and tests based on heterozygosity, were considered to analyse the data. Among 92 microsatellite loci, two were significantly less variable and lied out in heterozygosity based tests when comparing tolerant and susceptible populations. These results suggest that these loci are likely under selection or linked to local adaptative genes. Results of the two FST based tests were also presented and we discussed differences obtained with all methods. From a practical perspective, the functional testing of the polymorphisms of genes mapped in outliers regions may be used to determine how their genetic variation modulates susceptibility to trypanosomoses.

Poster 2197

Title: TG and DGAT have no influence on intramuscular fat content in Bavarian Simmental bulls

Presenting Author: Susanne Kämmerer, Bavarian State Research Center for Agriculture, Department Quality Assurance and Analysis, 85586 Grub/Germany

Other authors (name only):
1. Buitkamp, Johannes
2. Geuder, Ulrich
3. Schuster, Manfred
4. Götz, Kay-Uwe

Abstract:
Research has demonstrated that single nucleotide polymorphisms in the thyroglobulin gene (TG) and in the gene encoding diacylglycerol O-acyltransferase (DGAT 1) are associated with intramuscular fat content (IMF) or rather with the assessed trait marbling in cattle. IMF beeing an important beef quality trait, genotyping of breeding bulls is widely used. Nevertheless, it is unknown whether these associations are useful for typical breeds and production conditions in Germany. In the present study the associations of an SNP in the 5' promotor region of TG and a lysine/alanine polymorphism of DGAT1 with the fat content of m. long. dorsi in 100 Bavarian Simmental bulls was investigated. Bulls were chosen from 2.110 animals raised under standardized conditions of the Bavarian progeny test on station. Variance components and breeding values for IMF were estimated with DMU and PEST, respectively. Two groups were selected, each consisting of 50 bulls with the highest (Ø 0.92%) and lowest (Ø -0.60%) IMF breeding values, respectively. Neither of the markers showed significant differences in allele- or genotype-frequencies in both groups.
These results suggest that the TG and DGAT1 markers are no efficient predictors of IMF content in Bavarian Simmental bulls.

Poster 2198

Title: Maternal dietary methionine supplementation modulates foetal hepatic expression profiles

Presenting Author: Klaus Wimmers, Research Institute for the Biology of Farm Animals (FBN), Research Unit ’Molecular Biology’, 18196 Dummerstorf, Germany

Other authors (name only):
1. Eduard Murani
2. Wirawan Nuchchanart
3. Siriluck Ponsuksili

Abstract:
Impact of maternal nutrition during pregnancy on gene expression profiles and phenotypes of offspring was shown experimentally and in epidemiological studies in model animals and humans. The aim of the study is demonstrating of foetal programming in pigs, listing of genes sensitive to foetal programming and evidencing and quantifying of the role of DNA-methylation in this phenomenon. Therefore, 36 sows were fed methionine supplemented (‘MET’) vs. control (‘CON’) gestation diets. Liver samples of fetuses of three developmental stages (d35, d63, d91 p.c.) were subsequently monitored for diet-dependent differential gene expression by using Affymetrix microarrays. Statistical and bioinformatics evaluation of the expression profiles revealed largest effect of gestation diets at late foetal stages (91 dpc) when more than 1600 transcripts were differentially regulated. Functional annotation clustering highlighted the GO terms biological quality, cellular and anatomical structure morphogenesis, and negative regulation of metabolic processes. Interestingly, members of the IGF axis were found being diet-dependent regulated that play a key role in prenatal growth and are sensitive to regulation by DNA-methylation. Quantification of the DNA methylation at these loci will shed light on the role of this epigenetic mechanism as the molecular basis of genotype-environment (diet) interactions.

Poster 2199

Title: Frequency of IGF2 mutation in Polish breeds of pigs.

Presenting Author: Barbara Rejduch, National Research Institute of Animal Production Department of Immuno and Cytogenetics, ul. Krakowska 1, 32-083 Balice, Poland

Other authors (name only):
1. Maria Oczkowski
2. Katarzyna Walinowicz
3. Magdalena Rogoz
4. Marian Różycki

Abstract:
IGF2 is one of the gene that control muscle development. Using assay based on Real Time PCR for IGF2 genotyping, we evaluated frequency of IGF2 mutation in Polish breeds and assessed the usefulness of selection for this marker in Polish pig industry. In total 400 gilts and 35 wild boars (WB) were genotyped. Pigs represented seven breeds: Puławska(Pł), Landrace (L), Large White (LW), Duroc (D), Hampshire (H), Pietrain (P) and Line 990 (L.990). Previous reports shown that breeds, subjected to strong selection for lean meat have low frequency of G allele. Our study, performed on much higher number of animals, are in a great agreement with earlier findings. No G allele was observed in Duroc. In L. 990, Hampshire and Pietrain frequency of G allele did not exceed 11 %. Higher frequency (approximately 25%) of G allele was noted in Landrace and Large White. The highest frequency of G allele (36%) was observed in Puławska. Similarly to previous studies, no A allele was observed in wild boar.

Poster 2200

Title: Characterization of the ovine multicopy ribosomal protein SA gene family

Presenting Author: Mario Van Poucke Heidestraat 19 B-9820 Merelbeke Belgium

Other authors (name only):
1. Alice Van den Broeke
2. Ane Marcos-Carcavilla
3. Karine Hugot
4. Alex Van Zeveren
5. Luc Peelman

Abstract:
Transmissible spongiform encephalopathies (TSEs) are fatal, neurodegenerative diseases occurring in man (e.g. Creutzfeldt-Jakob) and animals (e.g. BSE and scrapie). They are caused by the accumulation of PrPSc, the alternatively folded isoform of the cellular protein PrPC, which is encoded by the PRNP gene. Several other genes are thought to play a role in TSE pathogenesis. This is the case for the ribosomal protein SA gene (RPSA), previously named 37-kDa laminin receptor precursor (LRP)/67-kDa laminin receptor (LR), which not only acts as a receptor for
Poster 2202

Title: Association in heavy pigs of CTSF and CAST polymorphisms with productive and meat quality traits

Presenting Author: Estefânia Alves, INIA, Deptº Mejora Genética Animal, 28040 Madrid - Spain

Abstract:
The porcine ACTC1 gene (actin alpha cardiac muscle 1) belongs to the alfa actin gene family. Alfa actins are a major constituent of the contractile apparatus and are involved in various types of cell motility. A partial cDNA (~1200 bp) of the porcine ACTC1 gene was identified in the subtracted foetal hind limb muscle cDNA library (44 days of gestation; using adult biceps femoris cDNA as the driver). Specific PCR-primers were designed for amplification and comparative sequencing (Meishan and Pietrain breeds) of corresponding segments of the gene. The whole gene...

Poster 2201

Title: The role of MHC class II genes in nematode infection of lambs.

Abstract:
Current methods for controlling nematodes are threatened by the evolution of drug resistance. Selective breeding of nematode-resistant animals represents an attractive alternative. In the UK, essentially all grazing sheep are exposed to a mixed, predominantly Teladorsagia circumcincta infection. Faecal nematode egg counts are widely used to classify lambs but the use of genetic markers would simplify the identification of superior animals. One genetic marker is the DRB1 locus in the class II region of the major histocompatibility complex. The G2 allele has been found to be associated with resistance in both Scottish Blackface and Suffolk lambs. The mechanism of gene action appears to be the IgE-mediated prevention of nematode establishment and possibly survival. Targets of the IgE response have been identified by 2-dimensional gel electrophoresis and genetic marker is the association in heavy pigs of CTSF and CAST polymorphisms with productive and meat quality traits. The IgE response followed by tandem mass spectrometry. Individuals with different MHC alleles respond differentially to the infection of lambs.

Poster 2203

Title: The porcine ACTC1 gene – structure, polymorphism, mapping and expression

Abstract:
The porcine ACTC1 gene (actin alpha cardiac muscle 1) belongs to the alfa actin gene family. Alfa actins are a major constituent of the contractile apparatus and are involved in various types of cell motility. A partial cDNA (~1200 bp) of the porcine ACTC1 gene was identified in the subtracted foetal hind limb muscle cDNA library (44 days of gestation; using adult biceps femoris cDNA as the driver). Specific PCR-primers were designed for amplification and comparative sequencing (Meishan and Pietrain breeds) of corresponding segments of the gene. The whole gene...
structure was completed using primer-walking approach applied on a porcine genomic BAC clone isolated with the ACTC1 primers. The gene is approx. 8 kb long and consists of 7 exons spanning 4163 bp, 1134 bp of which is the coding sequence. A G>A substitution was identified in intron 1. The gene was mapped using radiation hybrid panel (IMpRH) on chromosome 1 with SW65 as closest marker (41 cR; LOD = 7.73). By real-time RT-PCR, the gene was proven to be significantly overexpressed in porcine foetal skeletal muscles compared to adult ones. The ACTC1 gene is expected to be involved in muscle development and growth in pigs. (Supported by the Czech Science Foundation 523/06/1302)

**Poster 2204**

**Title: Determination of IFNG, IL2, IL4, and GATA3 single nucleotide polymorphisms in New Hampshire and White Leghorn chicken and analysis of association with worm burden after artificial infection with Heterakis gallinarum**

Presenting Author: G. Lühken, Department of Animal Breeding and Genetics, Justus Liebig University, Giessen, Germany

Other authors (name only):
1. M. Gauly
2. S. Weigend
3. H. Brandt
4. G. Erhardt

**Abstract:**

There is much evidence in chicken, similarly to mammalian species, that antihelmintic worm reactions are driven by an increase of type 2 cytokines and a decline of type 1 cytokines. The 5'-flanking region of the type 1 cytokine genes IFNG and IL2, the type 2 cytokine gene IL4 and the type 2 immune response directing transcription factor gene GATA3 were analysed for the occurrence of polymorphisms in the two layer lines New Hampshire and White Leghorn. Novel single nucleotide polymorphisms (SNPs) were identified in IL4 and GATA3. For each of the four genes, a SNP was genotyped by PCR-RFLP analysis in samples of New Hampshire (n = 146) and White Leghorn (n = 263) chicken where the worm burden had been determined eight weeks after artificial infection with 100 embryonated Heterakis gallinarum eggs. White Leghorn showed a significantly higher number of worms when compared with New Hampshire. Genotyping results of IFNG and GATA3 in New Hampshire were not used for statistical analysis because the two SNPs were nearly monomorphic in this line. Restricted maximum likelihood analysis with fixed effects of line, sex and genotype revealed no association of any of the tested SNP with the worm burden.

**Poster 2205**

**Title: Genetic characterization of the Bracco Italiano (Italian Hound) breed: first results on 22 STRs from the ISAG Canine Comparison Test**

ROBERTA CIAMPOLINI1, FRANCESCA CECCH1, ASSUNTA BRAMANTE2, FABIO CASETTI3, BRUNO DE SANCTIS4, SILVANO PRESCIUTTINI1

1Centro Interuniversitario di Ricerca e di Consulenza sulla Genetica del Cane, Università di Pisa, Viale delle Piagge 2, 56124 Pisa, Italy; 2 PhD Student Dip. Prod.i Anim., Università di Pisa Viale delle Piagge 2, 56124 Pisa, Italy; 3 Società Amatori Bracco Italiano (SABI), Via Dosso 7, 26856 Mirabello di Senna Lodigiana, Italy; 4 ASL Viterbo Italy.

**Abstract:**

The Società Amatori Bracco Italiano is financing a project aimed at investigating the demographic, genetic and genealogical structure of this old breed. We show an assessment of the genetic variability for 22 STRs typed in a sample of 33 unrelated Italian hounds (‘Bracchi’) and a sample of 43 dogs from other breeds (‘Other dogs’). Individual codes of all subjects were traced in the breed database (20,499 animals), so that the coefficient of relationship was < 0.2 with any other animal. Three multiplexes were worked out, which allowed analyzing 22 STR markers from the panels recommended for the 2006 and 2008 ISAG canine comparison test. Allele size in bp was determined using the comparison-test reference samples as anchor values. Data were analyzed using ARLEQUIN 3.11. The number of different alleles ranged 3 to 8 (mean 5.2) in the “Bracchi”, compared with 6-13 (mean 9.5) in the “Other dogs”, whereas the expected heterozygosity ranged 0.41-0.81 (mean 0.65) compared with 0.67-0.89 (mean 0.82). Interestingly, the difference between the observed and expected heterozygosities was lower in the “Bracchi” than in the “Other dogs” (0.65 - 0.58 = 0.07 vs. 0.82 - 0.62 = 0.20), despite the absolute values were lower in the “Bracchi”.

**Poster 2206**

**Title: MicroRNA profiles of the fetal pig during skeletal muscle development**

Presenting Author: Tara G. McDaneld, USDA/ARS Meat Animal Research Center, Clay Center, NE, 68901, USA
Other authors:
1. Ralph T. Wiedmann
2. Jeremy R Miles
3. Tim P.L. Smith

Abstract:
MicroRNA (miR) are a class of small RNAs that regulate gene expression. To evaluate the role of miR in skeletal muscle growth of the fetal pig, global miR expression was measured in skeletal muscle at three stages of growth including: primary fiber development (d 60), secondary fiber development (d 90), and preparturition (d 105). Adult muscle was also evaluated to compare muscle at a mature stage of growth (adult) to rapidly growing muscle (fetal). MicroRNA clone libraries identified a total of 95 miR matching sequences in the database with 3 sequences representing novel miR. Muscle-specific miR-206 was the highest expressed miR at all time points evaluated. Additionally, muscle-specific miR-1 was moderately expressed throughout fetal development with highest expression at d 105. Differential expression of miR at different stages of fetal development was also observed. MiR-432 was moderately expressed during early fetal development at d 60 compared to d 90 and d 105, while miR-424 and miR-126 expression increased during d 90 and d 105, respectively. These data are the first to evaluate miR expression at specific stages of fetal skeletal muscle growth in swine and identify novel miR that may play a vital role in skeletal muscle development and growth.

Poster 2208
Title: Nutrigenomics studies in sheep using a custom microarray platform

Presenting Author: Sgorlon Sandy, Via delle Scienze, 208 – 33100 Udine, ITALY

Other authors (name only):
1. Ferrari Alberto
2. Asquini Elisa
3. Dreos Renè
4. Stefanon Bruno

Abstract:
The aim of the research was to investigate the modifications of gene expression profiles in blood cells, after dietary supplementation with Echinacea angustifolia (EA) or Andrographis panicolata (AP) (INDENA, Milan) to sheep under ACTH challenge. Experimental protocol was approved by local laws and regulations. Twenty four sheep at maintenance were allotted to 4 groups: CTR- (negative control, without ACTH and supplementation), CTR+ (positive control, with ACTH and without supplementation), EA (ACTH and 3 mg/kg liveweight of EA) and AP (ACTH and 1 mg/kg liveweight of AP). ACTH was injected for 3 subsequent days and blood was sampled before (T0) and after 3 (T3) and 51 (T51) hours from the first injection. RNA extracted samples were pooled together within group and time of sampling. A custom oligoarray was synthesized using 24384 35-40mer probes designed from 12194 UniGenes (NCBI) on a CombiMatrix 90K platform. Cy5 labelled samples were hybridized on the chip. Data analysis of microarray allowed to identify a set of genes which were up or down regulated as a consequence of ACTH treatment. Functional classification showed that most of the genes behave to immune response and translation categories. EA and AP administration also showed an activity on these pathways regulation.
**Poster 2209**

**Title:** Linkage disequilibrium in a Korean beef cattle (Hanwoo) population

Presenting Author: Jong-Joo Kim, School of Biotechnology, Yeungnam University, Gyeongsan, 412449, Korea

Other authors (name only):
1. Yun-Mi Lee
2. Chan-Myung Han
3. Xioping Li
4. Song-Lan Li
5. Kwan-Suk Kim
6. Robert Schnabel
7. Jerry Taylor

**Abstract:**
The purpose of this study was to evaluate linkage disequilibrium (LD) in a Korean beef cattle (Hanwoo) population. Forty eight steers under paternal half-sib families and eight sires, each with six steers, were selected and sampled for DNA extraction in the population of Jeong-Ub Hanwoo association. Using the Illumina chips with 55K SNPs, 38,023 SNPs were determined as useful SNPs in Hanwoo, which was based on minor allele frequency of 0.05 and call rate of 90%. The SNPs were distributed across autosomal chromosomes with maximum (minimum) number of 2372 (659) on BTA1 (BTA28). Linkage phases or haplotypes were determined using fastPHASE (v 1.2.) program, and LDs between syntenic markers using only maternal chromosomes (haplotypes) were measured with the \( r^2 \) method. When the two markers were within 10 kbp, the average \( r^2 \) value was 0.52, decreased down to 0.2 for the pair of markers within 40 kbp, and down to 0.1 within 200 kbp. When the pair of markers was located longer than 5000 kbp (5cM), the average \( r^2 \) value was in equilibrium at 0.025. These results will provide useful information in implementing fine-mapping QTL for production traits in Hanwoo.

**Poster 2211**

**Title:** Molecular genetic dissection of bone mineral traits and osteochondrosis in porcine limbs

Presenting Author: Chirawath Phatsara, Institute of Animal Science, University of Bonn, Endenicher Allee 15, Bonn GERMANY

Other authors (name only):
1. Nopawan Rangkasenee
2. Watchara Laenoi
3. Elisabeth Jonas
4. Dawit Tesfaye
5. Heinz Juengst
6. Ernst Tholen
7. Helga Sauerwein
8. Karl Schellander

**Abstract:**
Osteochondrosis (OC) is regarded as the main cause of leg weakness in pigs. In this study, the OC lesion at articular cartilage was observed using a histological scoring method in 310 F$_2$ Duroc X Pietrain crossbred population. As well as bone mineral content and density (BMC and BMD, respectively) were also observed using a dual-energy x-ray absorptiometry technique. The histological scoring results revealed the most prevalent location was the knee joint (distal femur). There were no correlation between bone mineral traits (BMD and BMC) and performance traits, except slaughter weight. Five functional candidate genes (MGP, MMP3, TGFβ1, COL2A1, and COL10A1) for Osteochondrosis and bone mineral traits were screened for polymorphisms and further analysed for an association to the traits. Ten SNPs...
were identified in the candidate genes and genotyped by multiplex SNPs base extension method. An association analysis revealed polymorphisms of TGFβ1 and MGP were significantly associated with OC lesion at knee and hip joints, respectively. Polymorphisms of MGP, MMP3 and COL2A1 were found associated with BMD, while the association between MGP, MMP3 and TGFβ1 polymorphisms and BMC were also detected. Haplotype analysis of five candidate genes showed no association to the traits of interest.

Poster 2212

Title: Genetic diversity and admixture among Chinese cashmere goats

Ran DI
1 CAAS-ILRI Joint Laboratory on Livestock and Forage Genetic Resources, Institute of Animal Science, Chinese Academy of Agricultural Sciences (CAAS), No. 2, Yuan Ming Yuan Xi Lu, Haidian District, Beijing 100193, China

Other authors (name only):
1. Jianlin HAN
2. Yuehui MA

Abstract: Cashmere goats occur in northern China and Mongolia and are used to produce high quality natural fibre for the textile industry. In this study, 14 ISAG/FAO goat microsatellite markers were used to assess genetic diversity and admixture among nine Chinese cashmere goat breeds. Two Iranian and one West African goat populations were genotyped for comparative purposes. The results showed that the Chinese cashmere goats were much more genetically diverse (Nei’s unbiased gene diversity = 0.540-0.6520; allelic richness = 3.904-5.956) than the West African population (0.599 and 4.901), but lesser than the Iranian populations (0.6535-0.699 and 6.340-6.566); backing the hypothesis that goats were domesticated in the Middle East, from where they dispersed to Asia and Africa. Bayesian clustering approach grouped the Chinese cashmere goats into three separate genetic entities that were consistent with their geographic distribution and breeding history. The Tibetan Hegu goats were genetically distinct and formed the first cluster. The Liaoning cashmere goats from northeastern China together with the Xinjiang and Shanbei cashmere goats formed the second, while the remaining five breeds (Alashan, Erlangshan, Wuzhumuqin, Caidamu and Hexi) found in northwestern China formed the third cluster. These results could form the basis for genetic improvement of cashmere trait.

Poster 2213

Title: The first genetic linkage map for the saltwater crocodile (Crocodylus porosus)

LEE MILES1, SALLY ISBERG1,2, PAULINE DALZELL2, TRAVIS GLENN1,3, PETER THOMSON1, DANIEL PETERSON6, XUEYAN SHAN6 AND CHRIS MORAN1

1 Faculty of Veterinary Science, University of Sydney, NSW 2006, Australia.
2 Porosus Pty Ltd, PO Box 86, Palmerston, NT 0831, Australia.
3 South Eastern Area Laboratory Services, Randwick, NSW 2031, Australia.
4 Savannah River Ecology Laboratory, University of Georgia, P.O. Drawer E, Aiken, SC 29802, USA.
5 Department of Environmental Health Science, University of Georgia, Athens, GA 30602, USA.
6 Mississippi Genome Exploration Laboratory, Mississippi State University, Mississippi State, MS 39762, USA.

Abstract: Genetic maps currently exist for many commercially as well as evolutionary important species. In this study we presents the first genetic-linkage map for the Saltwater Crocodile (Crocodylus porosus), and indeed the first for the Class Reptilia. We constructed male, female and sex-averaged linkage maps for C. porosus using a total of 189 microsatellite DNA markers typed for between 83 and 480 individuals from between five and ten families obtained from Darwin Crocodile Farm, NT Australia. We identified 12 linkage groups (LG) with LG sizes ranging from two to 53 loci. The overall map consists of 161 loci, while 28 markers still remain unlinked. LG are currently being physically anchored to chromosomes (2n=34) using Fluorescent In Situ Hybridisation (FISH) methods. The current linkage map confirms that female crocodiles have extraordinarily higher rates of recombination than males, with overall map lengths of 1636.5 cM and 275.2 cM respectively. Next to the Atlantic salmon, this uncommonly large difference in recombination rates is the greatest reported for a vertebrate species. This first genetic map lays the framework for further mapping analyses, with QTL mapping of economically and evolutionarily important traits currently underway in the saltwater crocodile.

Poster 2214

Title: Intra-specific sequence variability of tumor necrosis factor-alpha gene (TNFα) in domestic horse breeds

Francisco EI1,3, Kalemkerian PB1,3, Liron JP1, Giovambattista G1, Echeverría MG2, Peral García P1, Diaz S1
Abstract:
Tumour Necrosis Factor alpha (TNFα) is a cytokine that plays a fundamental role in inflammation and immune responses. In horses the coding sequence of TNFα has been mapped within the MHC Class III region in chromosome 20, and SNPs in the promoter region have been previously reported in equine breeds and species. In order to preliminarily investigate the genetic variation within the TNFα gene in domestic horses, we analyse the genomic full-length sequences in fourteen animals from four horse breeds (Argentine Creole, Arab, Spanish Pure Breed and Argentine Silla). Oligonucleotide primers were designed based on the reported TNF horse genomic sequence, to amplify by PCR four overlapping fragments comprising ~2.6 kb within the gene. Each PCR product was sequenced, analyzed and the TNF full-length sequence assembled on each animal by using the algorithms of the Vector NTI and DNAMAN programs. A total of nine SNPs were observed, seven of them located in non-coding regions and two in coding regions. In addition, inter-specific variability was observed when compared to TNF reported sequences from other mammalian species; this variability corresponds to the presence of horse specific amino-acid motifs. Further studies are needed in order to investigate the functional specificity of those motifs and the significance of the observed intra- and inter-specific polymorphisms both in gene expression levels as in susceptibility to various horse diseases.

Poster 2215
Title: Association Study between DLA polymorphism and susceptibility to chronic superficial keratitis in German Shepherd dogs.

It, Verónica¹; Laura S Barrientos¹; Gustavo Zapata²; Silvina Díaz², Guillermo Giovambattista³.¹I.GEVET, CCT La Plata, FCV-UNLP-CONICET, 60 y 118 S/N, CC 296, 1900 La Plata, Argentina. ²Clinical Hospital, Faculty of Veterinary Science, University of La Plata, Argentina.

Abstract:
Chronic Superficial Keratitis (CSK) is a progressive, inflammatory and potentially blinding disease of the canine bilateral cornea. The German shepherd is the most commonly affected breed. CSK have been described as an immune-mediated disease. The increased MHC class II expression may play a part in perpetuating the corneal inflammation seen in the disease. Given the previously reported association between the CSK and up-regulation of major histocompatibility class II antigen expression, the primary goal of this work is to determine the association between the Dog Leukocyte Antigen system (DLA) and the disease in German Shepherd dogs. Blood samples were collected from twelve dogs with pannus and twelve control dogs. The polymorphism of DLA-DQA, -DQB and -DRB promoters, and four DLA linked microsatellites was genotyped. Genetic diversity was calculated for each locus, and association between genetic markers and presence/absence of pannus was evaluated. Preliminary results showed that: (i) the allele DLA-DQA1-DQA*p1a is fixed in both subpopulations; (ii) two alleles of the DLA-DRB1 promoter (DLA-DRB1*p1 and DLA-DRB1*p2) were detected with similar gene frequencies in both groups; (iii) we have only detect one (DLA-DQB*7) out of four DLA-DQB alleles reported in German Shepherd, as well as, the allele DLA-DQA*p1; (iv) three putative new DLA-DQB promoter variants were detected in our sample; (v) not significant differences were observed between illness and control group for the microsatellites FH2200 and FH2202; (vi) significant differences between pannus and control for the loci FH2054 and FH2975 (pFH2054 = 0.00744; pFH2975 =0.003214) were observed; (vii) a relative risk of 6.25 and 2.75 for the alleles FH2054*152 and FH2579*320, respectively, were calculated; and (viii) an odds ratio of 9 and 3.4 for these alleles, respectively, were estimated.

Poster 2216
Title: Genetic variability in Hunting dog breeds using microsatellite markers and mitochondrial DNA polymorphisms.

Presenting Author: Giulia Pertica, Dipartimento di Scienze Animali - Sezione di Zootecnica Veterinaria - Università degli Studi di Milano, 20133 Milano – Italy

Other authors (name only):
1 M.G. Strillacci 2 S.P. Marelli 3 P. Valiati 4 B. Gandolfi 5 M.C. Cozzi 6 M. Polli 7 M. Longeri 8 L. Guidobono Cavalchini

Abstract:
In order to define genetic relationships of Hunting dog breeds, 258 unrelated dogs of 13 breeds (pointing, retrievers and spaniels) were analyzed using ten microsatellite markers. Allele frequencies, average
number of alleles per locus, Hardy-Weinberg equilibrium were determined by GENEPOP. Observed proportion of heterozygotes (H_o), gene diversity within sample (H_s), overall gene diversity (H_t) were calculated using FSTAT. Factorial Correspondence Analysis (FCA) was carried out using the GENETIX. Individual genetic distances (D_DAS) were estimated and individual dendrogram was constructed using POPULATION. Individual breed assignment was carried out by STRUCTURE.

Genetic variations at the 694 bp fragment of the mtDNA D-loop were evaluated. We analysed 5 samples for each breeds. Sequences obtained were aligned with the complete canine mtDNA sequence (GenBank#NC_002008) using the CLUSTALX. Haplotype diversity (Hd), nucleotide diversity (p), average number of nucleotide differences (k) and Fu’s Fs statistic were calculated using DnaSP. Genetic distances were estimated by the Kimura-two parameter method and a phylogenetic tree was constructed using UPGMA algorithm. Median-joining network and mismatch analysis were calculated in order to investigate haplotypes relationship using NETWORK. The proportion of allele sharing combined with mtDNA analyses resulted in a clear clustering of the analyzed breeds.

Poster 2217

Title: Horse morphological judgment follow natural selection criteria?

Presenting Author: Jose Luis Vega-Pla. Laboratorio de Investigación Aplicada. Aparatado Oficial Suc. 2, 14071-Cordoba (Spain)

Other authors (name only):
1. Miguel Alcaide
2. Alvaro de Santiago
3. Constantino Alvarado
4. Juan José Negro

Abstract:
The heterozygosity theory attempts to identify the essence of genetic quality. It leads to the following general prediction: Traits of all kinds that are used by females when judging males reach their extreme expression in males with the greatest average heterozygosity. (Brown, 1997). More specifically, if the heterozygosity theory is correct, male symmetry expression, therefore male mating success in a large population, will positively correlate with degree of individual heterozygosity. The present study checks on the hypothesis that judge criteria in morphological competitions of Spanish Pure Breed horses (Andalusian) are linked to the heterozygosity theory. The inbreeding coefficient over the entire pedigree of each animal was estimated in two hundred champion stallions and eighty-four mares that became one of the three first positions in different morphological competitions. Also Heterozygosity and Fis statistic was calculated from twelve microsatellites typing. Results were compared with those obtained from a random sample population.

Poster 2218

Title: MicroRNA expression in differentiating porcine myoblast cells

Presenting Author: Mathilde Nielsen, University of Aarhus, Department of Genetics and Biotechnology, Denmark

Other authors (name only):
1. Bo Thomsen

Abstract:
MicroRNA contributes to regulation of gene expression by binding to complementary sites in protein-coding mRNA, thereby modulating the translation process. MicroRNAs control many biological processes such as myogenesis, and several muscle-specific microRNAs are known to be involved in diverse aspects of muscle function such as myoblast proliferation, differentiation, contractility and stress responsiveness. We are interested in identifying microRNAs that regulate muscle growth and differentiation. Thus, small RNA libraries were constructed from pig skeletal muscles and sequencing analysis identified several microRNAs including miR-133 and miR-206. The expression pattern of miR-133 and miR-206 were examined during differentiation of cultured myoblast cells into myotubes using stem-loop primers for reverse transcription followed by TaqMan PCR analysis. The data showed that the levels of mature miR-133 and miR-206 were low in the undifferentiated proliferating cells and that induction of myogenic differentiation pathway led to abruptly increased expression levels. This result is in agreement with the observation that miR-133 and miR-206 are among the most abundant microRNAs in skeletal muscle tissue and suggests a critical role of these microRNAs in expression of muscle-specific proteins.

Poster 2219

Title: Method of accurately detecting copy number variations in porcine KIT copies by oligonucleotide ligation assay

Presenting Author: Chae-Kyoung, Yoo, Division of Applied Life Science, Gyeongsang National University, Jinju 660-701, Korea

Other authors
1. Bo-Yeong, Seo
Abstract
Previous studies have used pyrosequencing, minisequencing, real-time PCR, invader assays and other techniques to detect copy number variations (CNVs). However, the higher a genome's copy number, the more difficult it is to resolve the copies; so an accurate method of measuring CNVs and assigning genotypes is needed. PCR followed by a quantitative oligonucleotide ligation assay (qOLA) was developed for quantifying CNVs. The assay's accuracy and precision were evaluated for porcine KIT, which was selected as a model locus. Using a method that combined qOLA and another pyrosequencing to quantitatively analyze KIT copies with spliced forms, we confirmed the segregation of KIT alleles in 145 F₁ animals with pedigree information; and verified the correct assignment of genotypes. In a diagnostic test on 100 randomly sampled commercial pigs, the genotypes obtained by grouping observations on a scatter plot agreed perfectly with those obtained by clustering using the nearest centroid sorting method implemented in PROC FASTCLUS of the SAS package. A test on 159 large white pigs noted only two discrepancies between genotypes assigned by the two clustering methods (98.7% agreement), confirming that the quantitative ligation assay established here enables genotyping through the accurate measurement of high KIT copy numbers (>4 per diploid genome).

Poster 2220
Title: Reconfirming and minimizing a QTL for intramuscular fat content by polymorphism of pig PDE4B in an Iberian × Landrace population

Presenting Author: Jae-Hwan, Kim, Address: Division of Applied Life Science, Gyeongsang National University, Jinju 660-701, Korea

Other authors
1. Cristina, Ovilo
2. Eung-Woo, Park
3. Almudena, Fernández
4. Jun-Heon, Lee
5. Jung-Gyu, Lee
6. Jin-Tae, Jeon

Abstract
Six transcription factors (POU5F1, NANOG, SOX2, LIN28, KLF4 and MYC) play important roles in the maintenance of embryonic stem cell pluripotency. We recently isolated cat embryonic stem (ES)-like cells from cat blastocysts generated in vivo. To identify genetic markers for characterizing cat ES-like cells, we have determined these six genes' coding sequences (CDSs). The sequence identity of these genes with orthologs of humans and mice was 76-100% and 58-99%, respectively, at the amino acid level. Among these, the expression of POU5F1 and NANOG mRNA was investigated in ES-like cells, blastocysts, fibroblast feeder cells and six adult tissues by RT-PCR. Transcripts of POU5F1 and NANOG were detected at a high level in ES-like cells and blastocysts. However, these two genes were undetectable in cat fibroblast feeder cells and six adult tissues. We also examined ES-like cells by immunocytochemistry, and demonstrated that POU5F1 and NANOG are present at a high level in cat
ES-like cells, and are undetectable in cat fibroblast feeder cells. These results confirm that cat ES-like cells can be successfully isolated from blastocysts produced in vivo. Moreover, the expression of POU5F1 and NANOG can be used as a biomarker for the characterization of cat ES-like cells; and SOX2, LIN28, KLF4 and MYC are necessary for additional analyses using cat ES-like cells.

Poster 2222

Title: A comparison of discriminating power between 14 STR and 60 SNP makers for individual cattle identification

Presenting Author: Hyun-Tae, Lim, Division of Applied Life Science, Gyeongsang National University, Jinju 660-701, Korea

Other authors
1. Jay You
2. Han-O, Park
3. Jung-Gyu, Lee
4. Jin-Tae, Jeon

Abstract
Short tandem repeat (STR) and single nucleotide polymorphism (SNP) typing methods have been widely used for cattle paternity and identity testing. Eleven STR markers used in the StockMarks® of Applied Biosystems, and three additional STR markers (BMS1747, BM4305 and BL1009), were selected and prepared in a multiplex PCR. Sixty SNP markers applied in the iPEX™ assay for the cattle paternity test of SEQUENOM were used. Genotyping data were collected from 480 Korean cattle. The discriminating efficiency of the two types of markers was compared by calculating He, PIC, F-statistics and probability of homolog using the MSA, CERVUS, FSTAT, GENEPOP and API-CALC programs. The probabilities of homolog for the STR markers were estimated as 3.43×10⁻²⁷, 4.18×10⁻¹⁹ and 3.98×10⁻⁸ in random mating, half-sib mating and full-sib mating populations, respectively. For the SNP markers, the probabilities were estimated at 2.09×10⁻²⁴, 4.69×10⁻²⁰ and 8.02×10⁻¹², respectively. For cattle individual identification, the STR markers would be about 1000 times more efficient than the SNP markers in a random mating population, and the SNP markers would be more effective than the STR markers in half-sib mating and full-sib mating populations. This indicates that a high number of makers is more informative in populations with blood relationships, although an STR markers has more alleles than an SNP markers in general.

Poster 2223

Title: High-Resolution RH Mapping of a Potential QTL Region on Sheep Chromosome 9 (OAR9)

Presenting Author: Tracy Hadfield, Utah State University, Old Main Hill 4700 Logan UTAH 84325 USA

Other authors (name only):
1. JAMES E. MILLER
2. CHUNHUA WU
3. NOELLE E. COCKETT

Abstract:
Gastrointestinal parasites have a profound detrimental effect on sheep production. Molecular markers associated with parasite resistance in sheep could be useful for genetic selection of superior animals. As a first step in identifying suitable markers, a genome-wide QTL scan has been implemented using DNA samples from 42 grandparents, 3 F1 sires, 97 F1 dams, and 195 F2 offspring, for which fecal egg count (FEC) for Haemonchus contortus were measured after natural and experimental challenges. Selective genotyping of the upper and lower 20% of the lambs for FEC after natural challenge identified a potential QTL region on ovine chromosome 9 (OAR9) between markers BMS108 and MCM63 within a 3.5 cM interval. In order to refine the QTL region, a high-resolution radiation hybrid (RH) map of OAR9 was constructed using the USUoRH5000 panel. Using a logarithm of odds (LOD) threshold of 6.0, a single linkage group was constructed and aligned with the genetic linkage map of this chromosome. The resulting RH map contains assignments for 28 microsatellite sequences, 21 BAC-end sequences and 8 genes. These experiments allowed resolution of the parasite resistance QTL region to an 8.0 Mb interval at position 90.0 Mb on OAR9.

Poster 2224

Title: DIAGNOSTIC OF EYE HEREDITARY DISEASES IN SLOUGHIS

Presenting Author: MICHAELA PŘIBÁŇOVA1, Ceskomoravská společnost chovatelů a.s., Laboratory of Immunogenetics, 252 09 Hradistko 123, Czech Republic

Other authors (name only):
1. PAVEL HORÁK
2. DANIELA SCHRÖFFELOVÁ
3. RENATA BECHYOVÁ
4. EVA ŠKODOVÁ
1, Czech Moravian Breeders Corporation, Laboratory of Immunogenetics, Hradištko, Czech Republic
**Abstract:**
Progressive retinal atrophy (PRA) causing blindness occurs in many breeds of dog in different forms. Sloughis suffer from late onset PRA form (**rcd1a**). Small population in the Czech Republic is based on Sloughis imported from Algeria, later breeders used for mating dogs from France, Germany, Italy and Morocco.

In this study, mutation test, causing PRA in Sloughis (8-bp insertion in exon 21 of the **PDE6B** gene, Dekomien et al, 2000) was used for testing Czech population of Sloughis. We have tested a total sum of 42 individuals, 41 Sloughis (30.4% of all living dogs registered in the Czech studbook) from 13 different litters (86.7% of all registered litters) and 100% imported dogs and 1 Erdelterrier, clinically affected with unknown PRA form. For comparison of results, samples of Affected and Carrier (heterozygous) Sloughis, obtained from Germany were used. All tested dogs was homozygous Normal, so we can claim, Czech Sloughi’s population is nearly clear from **rcd1a** mutation in contrast to German and American population.

We used the same test for clinically PRA affected Erdelterrier. In this dog breed PRA-causing mutation is unknown. There was observed no presence of **rcd1a** mutation in this sample, so we can except it as a cause of PRA in Erdelterriers.

(This work was supported by IRP IAPG project no. AVOZ50450515)

**Poster 2225**

**Title: Immunological Parameters of the Chicken Lines selected for the response in Graft-versus-host reaction in ovo**

Presenting Author: S Mashima, Graduate school of Science and Technology, Niigata University, Niigata, Japan

Other authors (name only):
1. Ko-zaburo Yamamoto
2. Ikuo Okada
3. Toru Abo

**Abstract:**
Chicken lines selected for the high and low competence in the response of the graft-versus-host reaction in ovo are valuable models for avian immunogenetics. Although **HA** and **LA** lines had been developed from a single colony and share the same **B** haplotype, **HA** is high responder in the reaction, whereas the latter exhibits lower response in the reaction.

Previously, **HA** and **LA** line showed significant differences in susceptibility to Marek’s disease (MD). The higher mortality in the **HA**, as compared with the **LA**, had been confirmed by virus challenge and transplantation of MD-transformed cell line.

In this study, we characterized cellular immunological parameters such as constitution of **T cell** subsets, mitogenic response to lectins or allogenic cells in these lines.

**Poster 2226**

**Title: Equine individual identification and parentage verification with 18plex and 22plex PCR**

Presenting Author: J.A. Bouzada, LCV, Algete, Spain

Other authors (name only):
1. J.A. Bouzada
2. J.M. Lozano
3. T. Mayoral
4. M.R. Maya
5. A. Trigo
6. E. Anadón
7. C. Gómez-Tejedor

**Abstract:**
Multiplex PCR is a powerful technique typically used in genotyping applications where the simultaneous analysis of multiple markers is required. For parentage testing and individual identification **multiplex PCR** amplifying DNA microsatellites is routinely used. In this study we propose a main set of 18 markers (AHT4, AHT5, ASB2, ASB17, ASB23, CA425, HSM1, HSM2, HSM3, HSM6, HSM7, HTG4, HTG6, HTG7, HTG10, LEX3, LEX33 y VHL20) and an additional set of 22 markers, (AHT29, AHT39, HSM8, LEX22, LEX27, TKY19, TKY279, TKY287, TKY294, TKY297, TKY301, TKY312, TKY321, TKY325, TKY333, TKY337, TKY341, TKY343, TKY344, TKY374, TKY394 y UCDEQ405) from the 2005-2006 International Society of Animal Genetics (ISAG) Comparison Test. The markers of each panel can be co-amplified simultaneously and meet the needs for parentage testing an individual identification in horses. Some changes in the habitual primers were required to achieve a final configuration that allowed all markers to be analysed together. Robotic procedures were implemented to minimize the risk of genotyping errors and parentage verification mistakes.

DNA from whole blood was extracted using BioSprint 96 Blood Kit (Qiagen) and each panel of markers was simultaneously amplified using the QIAGEN Multiplex PCR kit. PCR products were analyzed in a single run on an ABI PRISM® 3100 Genetic sequencer.
Microsatellites included in the proposal panels were highly polymorphic. The mean allele number is 6.61 in the main and 7.00 in the additional panel. The cumulative exclusion probability reached 99.9999526% using the main set and 99.9999977% using the additional one. The main panel is used routinely for equine parentage verification at LCV, and the additional one, as a complementary set of markers in more complex paternity queries involving related sires or when exclusion for one only marker is observed in the main panel.

DNA markers used in this study can be used in numerous fields of animal breeding, e.g. examination of genetic structure of populations, estimation of populations inbreeding, maintenance of autochthonous populations, estimation of genetic distance between populations and breeds, as well as planning of crossing programs.

**Poster 2227**

**Title: Detection of QTL on BTA6 for carcass traits in a Korean beef cattle (Hanwoo) population**

Presenting Author: Yun-Mi Lee, School of Biotechnology, Yeungnam University, Gyeongsan, 412449, Korea

Other authors (name only):
1. Chan-Myung Han
2. Yun-Seok Lee
3. Xioping Li
4. Song-Lan Li
5. Kwan-Suk Kim
6. Jeong-Sou Yeo
7. Jong-Joo Kim

**Abstract:**
The purpose of this study was to detect QTL for carcass traits in a Korean beef cattle (Hanwoo) population. Four hundred and fifteen steers were produced from 25 grand-sires under the national progeny-testing program in National Livestock Research Institute, Korea. The traits analyzed in this study were final weight at 24 month, carcass weight, backfat thickness, average daily gain, longissimus dorsi muscle area, marbling score. DNAs were extracted and eight microsatellite markers were chosen on BTA6, genotyped and linkage-mapped using Cri-map. The used markers and map locations (cM) were IL90 (0), BMS2508 (15), BMS518 (34), BM4621 (63), BM2460 (81), BM8124 (117), IL35 (144), and BMC4203 (175). MQREML program was used to detect QTL, and threshold at the 5% chromosome-wise level was determined using chi-square distribution with one degree of freedom. Three QTL were detected at the 5% chromosome-wise level; for average daily gain at 33 cM, for final weight at 63 cM, and for backfat thickness at 0 cM, respectively. These results will provide useful information to find potential candidate genes relevant to the traits that are located near the detected QTL.

**Poster 2228**

**Title: A First-Generation Ovine Whole-genome Radiation Hybrid Map using the USUoRH5000 Panel**

Chunhua Wu, Koh Nomura, Tom Goldammer, Jens Tetens, Cord Drögemüller, Tracy S. Hadfield, Brian P. Dalrymple, Jill F. Maddox and Noelle E. Cockett

Presenting Author: Noelle E. Cockett, Department of Animal, Dairy and Veterinary Sciences, Utah State University, 4900 Old Main Hill, Logan, Utah 84322-4900 USA

**Abstract:**
Whole-genome radiation hybrid (RH) mapping is a method for producing high resolution maps which can be used for integrating linkage and physical maps within a species and for determining gene order. Comparative maps can be created by incorporating genes/ESTs into the RH map that are common across species. An ovine whole-genome 5000 rad RH panel, referred to as USUoRH5000, was used to construct a first-generation ovine RH map that contains 918 type I and 388 type II markers clustered into 95 RH linkage groups distributed over all 26 ovine autosomes. The RH map has good agreement with existing ovine linkage and physical maps. The mapped loci were distributed, on average, every 18.42 cR (~ 2.63 Mb). By identifying the locations of mapped loci within the whole genome sequences of human and other closely related mammalian species, ovine–mammalian comparative maps were developed. Large blocks of synteny exist between sheep and other mammalian species, but previously unreported micro-rearrangements have been identified in this study. The USUoRH5000 panel will be an important tool for assembly of the ovine whole genome sequence and refinement of the virtual sheep genome, as well as aiding in the identification and positional cloning of genes influencing traits in sheep.

**Poster 2229**

**Title: A 6540 bp deletion in the LAMA3 gene is associated with hereditary junctional epidermolysis bullosa (JEB) in the American Saddlebred Horse**

Presenting Author: Kathryn T. Graves, Department of Veterinary Science, University of Kentucky, Lexington KY USA 40546

Other authors (name only):
1. Pamela J. Henney
Abstract:
Laminin 5 is a heterotrimeric basement membrane protein integral to the structure and function of the dermal-epidermal junction. It consists of three glycoprotein subunits; the α3, β3 and γ2 chains, encoded by the LAMA3, LAMB3 AND LAMC2 genes respectively. A mutation in any of these genes results in the condition known as hereditary junctional epidermolysis bullosa (JEB). A 6540bp deletion spanning exons 24 to 27 was found in the LAMA3 gene in American Saddlebred foals born with the skin blistering disease epitheliogenesis imperfecta. The deletion confirms that this autosomal recessive condition in the American Saddlebred Horse can indeed be classified as junctional epidermolysis bullosa and corresponds to Herlitz-JEB in humans. A diagnostic test was developed and 9 of 175 randomly selected American Saddlebred foals from the 2007 foal crop were found to be carriers of the mutation (gene frequency 0.026).

Poster 2230

Title: Whole genome association study for protein yield in German Braunvieh

Authors:
Presenting Author: Hermann Schwarzenbacher, Lehrstuhl für Tierzucht, Technische Universität München, Hochfeldweg 1, 85354 Freising-Weihenstephan, Germany

Other authors (name only):
1. Krzysztof Flisikowski
2. Christine Wurmser
3. Franz Seefried
4. Peter Lichtner
5. Thomas Meitinger
6. Ruedi Fries

Abstract:
Selectice genotyping was applied to 140 bulls with highest estimated breeding values (EBVs) for protein yield and to 148 bulls with lowest EBVs among 973 progeny tested Braunvieh bulls. Animals were genotyped using the Illumina BovineSNP BeadChip which comprises 54001 SNPs. PLINK software v1.02 and the R environment for programming and statistical computing were used for the analysis of genotyping data. SNP and animals with call rates <90% were omitted. Other exclusion criteria were HWE P-values ≤0.001 and a minor allele frequency <5%, resulting in 36821 SNPs in 287 animals to be included in the association study by allele regression. Comparison of observed with expected test statistics under the null hypothesis indicated substantial stratification. However, accounting for pedigree information and identical by state clustering successfully controlled for population substructure. Several significant SNPs will be followed up by fine-mapping and candidate gene analyses.

Poster 2231

Title: Genetic relationships between native goat populations from Spain, Portugal and America

Presenting Author: Amparo Martínez Martinez. Departamento de Genética, Universidad de Córdoba, Spain.

Other authors (name only):

Abstract:
Twenty seven microsatellite markers were analysed in 1450 goats belonging to 38 native goat populations from Spain, Portugal and related Creole populations from America in order to establish their genetic relationships. The genetic differentiation between populations was moderate with a mean FST value of 0.13. The neighbour-joining tree constructed with the D2 genetic distance values demonstrate that Creole populations from Argentina, Bolivia and the Spanish goat from USA grouped in the same cluster that the Iberian populations while the Cuban Creole goat grouped with the populations from Canary and Cabo Verde Islands. Clustering of populations and estimates of the proportions of the individual genomes that were derived from the respective inferred clusters were obtained using the model-based clustering program Structure (Pritchard et al. 2000). Results for k=2 showed that populations from the Iberian Peninsula (Spain and Portugal) and most Creole goats grouped in the same cluster, while the Brazilian, Canarian, Cuban and Arapawan populations formed another cluster. Setting k=3 generated a cluster with the Canarian, Cuban and Arapawan populations. The results obtained suggested a clear influence of the Iberian populations in the Creole goats, while the Cuban Creole goat is more influenced by the Canarian breeds than by Iberian goats.

Poster 2232

Title: Annotation, polymorphism and expression analysis of the bovine INS-IGF2 locus

Krzysztof Flisikowski1, Pawel Lisowski2, Hermann Schwarzenbacher1, Ruedi Fries1

1 Lehrstuhl für Tierzucht, Technische Universität
Abstract:
Transcription of insulin (INS) and insulin like growth factor (IGF2) genes results in a bicistronic mRNA. Expression of the two genes may therefore be subject to common regulation. Based on the bovine draft sequence Btau_3.1 and EST databases of different species, we determined the intron-exon structure and identified regulatory regions including five promoters at the INS-IGF2 locus. Eight novel polymorphisms were identified in the regulatory regions, among them three in the INS promoter that potentially alter the binding capacity of AP-2 and NF-1 transcription factors. Additionally, a detailed quantitative analysis of tissue- and age-dependent expression of the IGF2 derived sequences was performed. The mRNA level differed significantly between tissues. Highest expression was observed in the liver and kidney, whereas it was only spurious in muscle and spleen. Expression was fourfold higher in two-month male fetuses than in twelve-month old bulls. Examination of the DNA methylation patterns revealed full methylation (93%) of the INS-IGF2 promoter independent of the age. Two SNPs localized in this region create new methylated CpG sites. These polymorphisms could affect the cis-activation of the bovine INS-IGF2 transcript and may be associated with variation of developmental traits.

Poster 2233

Title: Genetic characterization of Italian heavy draft horse (IHDH) breed using microsatellite markers

Presenting Author: Fabio Maretto. Department of Animal Science, University of Padova, AGRIPOLIS, 35020 Legnaro (PD), Italy.

Other authors (name only):
1. Cinzia Ribeca
2. Roberto Mantovani

Abstract:
The IHDH breed was established in 1926 by the Italian government and originated from crosses of Norfolk-Breton stallions with local derived Hackney, Percheron and Bretons mares. The breed was developed for agricultural and draft uses as well for artillery transport by the Italian army; nowadays it is mainly used for meat production and heavy draft works. This study compares genetic diversity of IHDH and other two unrelated breeds (Italian Haflinger, IH and Quarter Horse, QH breeds) by means of 23 microsatellite markers. A total of 95 animals (IHDH n = 55; IH n = 19 and QH n = 21) were genotyped. A total of 165 alleles were detected in the whole sample and the number of alleles (N_a) varied between 3 (locus Htg7 in the IH breed) and 13 (locus Tky343 in the IHDH breed) with a mean of 6.06 ± 1.72. The average observed heterozygosity (H_o) was 0.66, 0.67 and 0.76 for IH, IHDH and QH breed, respectively. The IHDH breed showed a good genetic variability comparable to the other analysed breeds. This study provides the first baseline of data for the IHDH breed and contributes to the conservation and implementation of selection programme of the breed.

Poster 2234

Title: High throughput SNP discovery and validation in the Pig: towards the development of a high density swine SNP chip

Presenting Author: Crooijmans Richard P.M.A. Animal breeding and Genomics Centre, Wageningen University, Marijkeweg 40, Wageningen, The Netherlands

Other authors (name only):
1. Ramos, A.M.
2. Amaral, A.J.
3. Kerstens, H.
4. Bendixen, C.
5. Hedegaard, J.
6. Rohrer, G.
7. Smith, T
8. Tassell van, C.
9. Taylor, J.F.
10 Rothschild, M.
11. Nonneman, D.
12. Beever, J
13. Archibald, A.L.
14. Schook, L.
15. Groenen, M.A.M.

Abstract:
Recent developments in sequencing technology have allowed the rapid and inexpensive generation of millions of short read sequences, enabling the cost effective identification of the hundreds of thousands of SNPs. Currently, a high density swine SNP chip is being developed as part of an integrated effort of several European and U.S. institutions involved in swine genomics research. The future pig SNP chip will include previously validated SNPs and SNPs identified de novo using second generation sequencing on the Illumina Genome analyzer (Solexa) and Roche 454 FLX sequencer. Towards this end, 20 DNA libraries were prepared using pooled DNA samples from individuals from five breeds (Duroc, Landrace, Large White, Pietrain, Wild Boar) digested with three restriction enzymes (AluI, HaeIII, MspI). The Solexa...
short sequence and longer 454 reads will be filtered using several quality criteria to produce the dataset used for SNP discovery. Additional criteria for the selection of the 60K SNPs to be incorporated into an Illumina iSelect assay include the Illumina design score, Infinium I vs II, estimated minor allele frequency and genomic location of the SNPs. The 60K SNP chip will be an extremely valuable tool for a variety of applications including QTL and LD mapping, association studies and genomic selection.

Poster 2235

Title: Insulin induced gene 2 (INSIG2) is associated with fatty acid composition in the pig

Presenting Author: Anna Pertek, Lehrstuhl fuer Tierzucht, Technische Universitaet Muenchen, Hochfeldweg 1, 85354 Freising, Germany

Other authors (name only):
1. Hermann Schwarzenbacher
2. Krzysztof Flisikowski
3. Ruedi Fries

Abstract:
Insulin induced gene 2 (INSIG2) is involved in sterol regulatory element binding protein (SREBP) activation of fatty acids and thus a candidate gene for fatty acid composition traits. We investigated porcine INSIG2 in a Mangalitsa x Piétrain cross. The genomic sequence of INSIG2 was obtained by sequencing a porcine BAC clone. The coding sequence, flanking splice junctions and a putative promoter region were screened for polymorphisms in the parental animals of the cross. A total of 13 variants were identified, including a synonymous substitution (c.237A>G) and a microsatellite polymorphism (*121_122insGT) in the 3’ untranslated region (3’-UTR). Four haplotypes were derived based on 12 variable positions, but haplotype 4 was not transmitted to the F2 generation. An association study was conducted including 324 F2 animals. Haplotype 1 significantly decreases saturated fatty acids percentage in the intramuscular fat. The content of polyunsaturated fatty acids (PUFA) is significantly associated with haplotypes 1 and 3. While haplotype 1 increases PUFA content, haplotype 3 reduces it. Haplotype 3 significantly increases monounsaturated fatty acids percentage. An obviously functional polymorphism could not be identified. However, microsatellite variation in the 3’-UTR may affect the expression of INSIG2.

Poster 2236

Title: Expression profiling and eQTL analysis reveal biological pathways and candidate genes affecting water holding capacity of muscle

Presenting Author: Siriluck Ponsuksili, Research Institute for the Biology of Farm Animals (FBN), Research Group ‘Functional Genome Analysis’, 18196 Dummerstorf, Germany

Other authors (name only):
1. E. Jonas
2. E. Murani
3. C. Phatsara
4. T. Srikanthai
5. C. Walz
6. M. Schwerin
7. K. Schellander
8. K. Wimmers

Abstract:
There is considerable variation in the water holding capacity (WHC) of meat, which is an important economic factor in pork production. WHC depends on numerous genetic and environmental factors that affect muscle fibre properties and metabolic processes ante and post slaughter. We aimed to identify candidate genes for WHC by QTL mapping, expression profiling and expression QTL (eQTL) analysis. Expression microarray analysis of M. longissimus dorsi RNAs of 74 F2 animals of a resource population showed 1,279 transcripts with trait correlated expression. Negatively correlated transcripts were enriched in functional categories and pathways of extracellular matrix, receptor interaction, and Ca-signalling; positively correlated transcripts dominantly represented oxidative phosphorylation, mitochondrial pathways, and transporter activity. A eQTL analysis of trait correlated transcripts revealed 897 eQTL, with 104 eQTL matching previously identified QTL for WHC; 96 transcripts were trans regulated and 8 had cis acting regulation. Based on these findings, a priority list of genes was established out of the orchestra of genes of biological networks relevant to the liability to elevated drip loss. The complex relationships between biological processes in live skeletal muscle and meat quality are mainly influenced by the energy reserves and their utilisation in the muscle and by the muscle structure itself.

Poster 2237

Title: Identification of a SNP in TRPM1 and its association with Leopard Complex Spotting (LP) and Congenital Stationary Night Blindness (CSNB) in horses.

Presenting Author (name & address): Rebecca Bellone University of Tampa 401 W. Kennedy Blvd. Tampa, FL 33606 USA
Abstract:
The Leopard Complex Spotting patterns in horses is caused by an incompletely dominant gene (*LP*) mapped to a 6cM region on ECA1 containing the candidate gene *TRPM1* (Transient Receptor Potential Cation Channel, Subfamily M, Member 1). Homozygosity for *LP* is associated with congenital stationary night blindness (CSNB) in Appaloosa horses. One SNP, found within an intron, had a distribution identical to *LP* on a panel of 10 horses. To determine the strength of the association, 397 unrelated horses from different breeds were tested. There was a complete association of this SNP with *LP* (*P < 0.0005*) and CSNB (*N*=30, *P < 0.0005*) among Appaloosa and Knabstrupper horses. Among Noriker horses, a strong association was observed (*P < 0.0005*) however this association was not complete; 34 horses identified as non-patterned (*lp/lp*) possessed at least one copy of this SNP. These results suggest two possible conclusions: the SNP is not responsible for *LP* spotting but associated with the phenotype in Appaloosa and Knabstrupper horses while less so among Noriker horses; or the phenotypic assignments of *LP* were not accurate. Additional work is underway to identify other SNPs in the region and to assess their potential effects on the *LP* phenotype.
Poster 3001

Title: Expression of four muscle protein genes in the Günther’s walking catfish Clarias macrocephalus

Presenting Author: Supawadee Poompuang, Department of Aquaculture, Faculty of Fisheries, Kasetsart University, Bangkok 10900 Thailand

Abstract:
Skeletal muscle proteins represent more than half of the fish body mass. Expression of muscle genes affects development and growth of the animal. Because growth is an important trait in fish farming, it is useful to identify genes associated with muscle growth. In this study, we identified the complete cDNA sequences of muscle genes and analyzed their expression profiles from larval to adult stages of walking catfish. The complete cDNA sequences of parvalbumin (PV) (670 bp), troponin C (TnC) (1,065 bp), troponin I (TnI) (843 bp), and myosin light chain III (MLCIII) (953 bp) were obtained from the skeletal muscle cDNA library. The PV cDNA encoded a protein of 109 amino acids. The deduced amino acid sequences of TnC, TnI, and MLCIII were 160, 176, and 150 residues, respectively. The transcript levels of each muscle gene were determined by semi-quantitative RT-PCR. Significant variations of gene transcripts were observed during larval development, with the TnC mRNA showed its highest levels. Expression of PV, TnC, MLCIII, and TnI were highest in skeletal muscle of juvenile and adult fish with lower expression in skin and gills. The levels of PV, TnC, and MLCIII transcripts were similar in muscle of juvenile fish while higher expression of PV was observed in adult fish.

Poster 3002

Title: Characterization of seven microsatellites in the Black Sea stellate sturgeon (Acipenser stellatus)

Presenting Author: Georgescu Sergiu Emil, Splaiul Independentei 91-95, 050095, Romania

Abstract:
Sturgeon populations from Romania have declined because of anthropogenic influences. In this context, the necessity to develop conservation programs for these species requires knowledge of their genetic diversity.

The aim of this study was to optimize a protocol for microsatellites multiplexing in order to estimate genetic variability in Acipenser stellatus. Genomic DNA extracted from fin clips was amplified in two PCR reactions using seven primer pairs labelled with four different fluorescent dyes to detect seven different microsatellites (LS-19, LS-34, LS-54, LS-57, LS-68, Aox23 and Aox45). The amplicons analyzed with an ABI-Prism-310 Genetic Analyzer were examined with specialized software.

The number of allele peaks depends on the level of ploidy of the studied species and on whether the fish is heterozygote or homozygote. The size of the alleles at every locus varied between 95-200 bp. A high level of polymorphism was observed for the population studied. Seven to thirteen alleles where detected with a mean of eight alleles per locus.

The co-amplification of the microsatellites is a very good method allowing the evaluation of intraspecific genetic diversity, used for the first time in Romania. It will allow us to characterize the genetic variations in Black Sea sturgeon populations.

Poster 3003

Title: Genetic characterization of the hybrid between two commercially important Brazilian catfishes

Presenting Author: Denise Aparecida Andrade de Oliveria, Escola de Veterinária da UFMG Av.Antônio Carlos 6627, 31270-010 Belo Horizonte-MG BRASIL, (31)3409-2206 fax:(31)3491-3963

Abstract:
The Pimelodidea catfish species Pseudoplatystoma corrucans (surubim or pintado) and P. fasciatum (cachara) are large, migratory fishes that represent an important fishery resource in their respective areas of occurrence in Brazil. The hybrid form of the two species is of great importance to the local aquaculture industry. In this study, it was investigated whether the two species could be distinguished genetically by sequencing the mitochondrial 16S rRNA of pure strains and hybrids. Samples of P. corrucans were collected in the São Francisco River over a period of 3 years. These were compared with captive P. fasciatum and with hybrids of the two species. Five 16S rRNA
haplotypes of *P. corruscans* were recovered. A fourth haplotype was found in *P. fasciatum* and in hybrids only, indicating that the two species can be genetically distinguished and that the hybrids sequenced in this study are descendants of the females of *P. fasciatum*. (This work was supported by FAPEMIG-CBB 243/04)

**Poster 3004**

**Title:** Culturing diversity? Contrasting genetic diversity between wild and cultured populations of Pacific lion-paw scallop

Presenting Author: Jessica L. Petersen, University of California Davis, Dept of Animal Science, Davis CA 95616, USA

Other authors (name only):
1. Ana Maria Ibarra
2. Bernie May

**Abstract:**
Because cultured lion-paw scallops achieve sexual maturity before harvest, introgression with adjacent wild populations is possible. Eleven microsatellite loci were used to evaluate if cultured populations in the Lagoon Guerrero Negro, Mexico, encompass the genetic variation present in the wild by examining the genetic diversity of: 1) wild lion-paw scallops around the Baja California Peninsula, Mexico, 2) three artificially spawned cohorts of scallops grown for aquaculture, and 3) a sample of wild spat collected by producers for the purpose of future artificial spawns. Genetic data coupled with observed shell abnormalities in one aquaculture cohort infer high levels of inbreeding. Pairwise *F*<sub>ST</sub> values show all three aquaculture cohorts are significantly different than the wild aggregations, even from the supposed source populations. In contrast, the wild spat collected for future spawns was not different than the wild source population, suggesting that spawning methodology and/or unrecognized selection pressures in aquaculture produce populations that do not capture the diversity found in the wild. These results warn that unless aquaculture propagation methods are amended to conserve natural diversity, introgression of aquaculture populations into the wild could alter the natural genetic make-up of the species at the possible demise of fitness and potential for adaptation.

**Poster 3006**

**Title:** Molecular investigation of resistance to amoebic gill disease (AGD) in Atlantic salmon.

Presenting Author: Rebl, Alexander, Forschungsinstitut für die Biologie landwirtschaftlicher Nutztiere, (Research Institute for the Biology of Farm Animals), Molecular Biology Research Unit, Wilhelm-Stahl-Allee 2, 18196 Dummerstorf, Germany

Other authors (name only):
1. Goldammer, Tom
2. Wimmers, Klaus

**Abstract:**
The Gram-negative bacterium *Aeromonas salmonicida* causing furunculosis, a systemic disease in salmonid fish, is responsible for considerable losses to the aquaculture industry. Selection of Steelhead rainbow trouts has been started in 1975 at the fishery institute in Born in order to breed fish with an elevated fitness in brackish water. Current investigations prove that the strain BORN exhibits a higher resistance towards *A. salmonicida* than the non-selected STEELHEAD trouts. We aim to examine the physiological mechanisms underlying this improved resistance. In the present pilot study, we compared gene expression profiles in liver tissue of healthy rainbow trout from the strains STEELHEAD and BORN via microarray technology. Among the 151 genes that were identified as being significantly differentially expressed we chose two genes to validate expression in a variety of tissues by real-time RT-PCR. Alfa<sub>1</sub>-antitrypsin gene is significantly up-regulated in kidney and liver of BORN trouts. This proteinase inhibitor is an acute-phase protein contributing to early immune response in mammals. However, dehydrodolichyl diphosphate synthase, a key enzyme of dolichol biosynthesis, was significantly down-regulated in muscle, kidney, spleen, gills, liver, and heart of BORN trouts. Further microarray experiments are planned to identify different molecular features in both strains after *A. salmonicida* infection.
of teleost hosts. While a moderate heritability for AGD resistance in Atlantic salmon has been reported previously, the mechanisms by which individuals resist the proliferative effects remain poorly understood. To gain more knowledge of this commercially important trait, we compared the gill transcriptome of Atlantic salmon putatively resistant and putatively susceptible to AGD using the 17k TRAITS/SGP Atlantic salmon cDNA microarray. We identified a large number of transcripts that were differentially expressed between the AGD resistant and susceptible animals, many of which were involved in the immune and cell cycle responses. Individuals resistant to AGD displayed significantly higher expression of genes involved in adaptive immunity and negative regulation of the cell cycle. In contrast, AGD susceptible individuals showed higher expression of acute phase proteins and positive regulators of the cell cycle. Expression of a subsample of transcripts was further examined with real-time quantitative RT-PCR (qPCR) in the AGD resistant and susceptible animals, as well as non-infected naïve fish. The identification of these responses has provided new insight into the molecular mechanisms controlling AGD resistance.

Poster 3008

Title: Molecular characterization of Alpine and Northern European populations of Arctic charr Salvelinus alpinus (Linnaeus, 1758) by means of nuclear and mitochondrial markers

P. Ajmone Marsan, Ist. Di Zootecnica, Università Cattolica del Sacro Cuore- Via E. PArense 84, 29100 Piacenza-Italy

Other authors (name only):
1. Licia Colli
2. Andrea Gandolfi
3. Fatima Chegdani
4. Lorenzo Bomba
5. Rosanna Marino
6. Marco Pellecchia
7. Francesco Nonnis-Marzano
8. Stella Grando
9. Davide Perini
10. Riccardo Negrini

Abstract:

The Arctic charr, Salvelinus alpinus, is an Holarctic Salmonid species adapted to cold-water habitats and whose populations, either anadromous or landlocked, are characterized by an extremely variable morphology. The presence of S. alpinus in Southern Europe is deemed as a relict of the Pleistocenic glaciations which strongly influenced both the distribution and the genetic variability of the species due to repeated isolation and bottleneck events. Previous genetic studies revealed a very low mitochondrial diversity in contrast with the high genetic variability, both within and among populations, highlighted by microsatellites. In this study, 537 specimens from 35 Alpine and 5 Scandinavian populations were characterized by AFLP polymorphisms and mtDNA control region sequences.

The results of AFLP analyses were evaluated by Factorial Correspondence Analysis (individual level) and by Multi-Dimensional Scaling (population level), while hidden genetic structures were investigated by a Bayesian clustering approach. The control region data were aligned with GenBank sequences to build a Median-joining network. Multivariate statistical analyses showed that individuals and populations
mostly clustered according to the geographic origin. Furthermore, some populations displayed low levels of genetic polymorphism which is probably due either to past demographic fluctuations or to a strong impact of anthropic activities in the recent decades.

**Poster 3009**

**Title:** MOLECULAR MARKER-ASSISTED COHORT SELECTION OF YELLOW PERCH

Presenting Author: Han-Ping Wang, Ohio Aquaculture Research and Development Integration Program (OARDIP), Ohio State University, 1864 Shyville Road, Piketon, Ohio 45661 USA

Other authors (name only):
1. Geoff K. Wallat
2. Hong Yao
3. Paul O’Bryant
4. Jacob D. Rapp
5. Laura G. Tiu
6. Bonnie Brown

**Abstract:**
Approximately 800 broodfish from eight stocks representing major populations were obtained from eight states. Those fish serve as a base population for a genetic improvement program. Genetic variance analysis suggested that all strains had genetic value to the program. Two improved lines were created in 2006 from selected broodfish of 2004 year-class, and three improved lines in 2007 from selected broodfish of 2005 year-class by performing cross-breeding of the base population. For each generation, approximately 500 fish were selected and genotyped from the base population and base generation using microsatellite to construct a pedigree. Among the 500 fish, 100 pairs of the least related, with highest breeding value were selected and divided into five cohorts based on their pedigree. The selection lines were created by pair-mating 20 pairs within each cohort to found the next generation of improved lines. On-station and on-farm tests showed that the two improved lines of 2006 grew 54.0% and 28.0% faster than unimproved fish, respectively, and the improved lines of 2007 grew 25% - 42% faster than controls. Parentage analysis techniques using six molecular markers in yellow perch have been developed. Two mapping families have been established and reared to maturation for QTL mapping.

**Poster 3010**

**Title:** Genetic structure from the anchovy of the Bay of Biscay using microsatellites and SNP discovery in the European anchovy

Presenting Author: Iratxe Zarraonaindia Martínez, Genetics, Physical Anthropology and Animal Physiology Department. Faculty of Science and Technology. University of the Basque Country.

Other authors (name only):
1. Miguel Angel Pardo
2. Mikel Iriondo
3. Carmen Manzano
4. Andone Estonba

**Abstract:**
Our interest is focused on unravelling the genetic structure of anchovy populations in the Bay of Biscay using DNA molecular markers, microsatellites and single nucleotide polymorphisms (SNPs). Our aim is to provide useful data for an effective conservation programme in this species, recently suffering a severe biomass reduction. In all, 10 microsatellite markers were analysed in 96 samples from each of two ICES divisions of the Bay of Biscay, and 96 samples from the Gulf of Cadiz and the Gulf of Lyon. Genetic structuring was detected in the Bay of Biscay’s anchovies, suggesting that they do not behave as a panmictic population. On the other hand, SNPs are rapidly emerging as the preferred genetic markers for molecular genetic analysis, but there is a lack of information about SNPs in the anchovy genome. That is why we carried out a SNP discovery development using two approaches: comparative sequencing of random DNA fragments and the exon primed intron-crossing (EPIC) method, which have proven to be successful in SNP identification.

**Poster 3011**

**Title:** Use of SNP-chips and selective DNA pooling for identifying disease resistance markers in Atlantic salmon


Other authors (name only):
1. Håvard Bakke
2. Ben Hayes
3. Sigbjørn Lien

**Abstract:**
Diseases such as infectious salmon anemia (ISA) continue to present a large problem for Atlantic salmon aquaculture in Norway. Current approaches to improve resistance by selective breeding involve challenge testing of a small number of individuals from a large number of families to obtain family rankings and breeding values. This strategy doesn’t exploit within family variation, meaning there is great
potential for the application of marker assisted selection. Simulation studies have shown that combined linkage / linkage disequilibrium analysis has high power to detect QTL using this population structure with a high density SNP marker resource. This project aims to identify QTL for ISA resistance using challenge test samples stored in a biobank from the SalmoBreed breeding population, and a SNP chip containing up to 60,000 markers. In addition, the selective DNA pooling methodology, where DNA pools of extreme individuals are typed rather than individual samples, will be evaluated as a means of dramatically reducing genotyping requirements. Families showing high variation for ISA resistance have been identified and preparation of DNA from these animals is currently underway. Markers identified in LD with disease resistance QTL will provide a powerful tool for improved selection for disease resistance across the whole breeding population.

Poster 3012

Title: GENETIC CONSEQUENCES OF DIFFERENTIAL SURVIVAL RATES IN CULTURED SILVER-LIPPED PEARL OYSTERS, Pinctada maxima.

Presenting Author: Curtis E. Lind, Aquaculture Genetics Group, School of Marine & Tropical Biology, James Cook University, Townsville QLD, 4810

Other authors (name only):
1. Brad S. Evans
2. Joseph J. U. Taylor
3. Dean R. Jerry

Abstract:
The pearlimg industry can benefit substantially from selective breeding programs; however, sustained genetic response to selection is highly dependent on the maintenance of genetic diversity within cultured populations. It has been recently shown that genetic diversity of Pinctada maxima is significantly reduced due to the culture process. What are still unknown are the underlying factors contributing to this loss. A possible strategy to overcome this and maintain genetic diversity is to equalise family sizes, which can be performed in the hatchery prior to communal rearing. However, this effort may be inconsequential if differential family survival through the larval rearing period ultimately results in highly variable family sizes. Through DNA pedigree analyses, this presentation scrutinises how genetic diversity is affected by differential survival of communally reared families, and whether equalising family sizes prior to communal rearing will influence effective genetic sizes. Results show that variance in family survival rates are present and can have detrimental effects on effective population sizes and genetic diversity. In one instance, complete dropout of families was observed. Consequently, the practice of equalising family sizes to maximise diversity may be ineffective and alternative approaches should therefore be investigated to avoid diversity loss and maximise response to selective breeding.

Poster 3013

Title: The Atlantic Cod Genomics and Broodstock Development Project (CGP)

Presenting Author: Rise M, Ocean Sciences Centre, Memorial University of Newfoundland, St. John’s, Newfoundland and Labrador, Canada

Other authors (name only):
1. Trippel EA
2. Gamperl AK
3. Johnson SC
4. Rise ML
5. Robinson A
6. Culver K
7. Garber A
8. Borza T
9. Hubert S
10. Tosh J
11. Lush L
12. Neil S
13. Afonso L
14. Hori T
15. Were K
16. Nardi G
17. Moir J
18. Powell F
19. Walsh A
20. Symonds J
21. Bowman S

Abstract:
The objective of the CGP is to develop breeding programs and fundamental genomics tools which will be used to supply the developing Atlantic cod aquaculture industry in Canada with improved broodstock. Three major spawning seasons have been completed in Newfoundland and New Brunswick / New Hampshire. Evaluations of progeny include assessment of family performance related to growth, survival and overall health. In addition, investigations related to cod physiology and immunology are conducted. Considerable variations among families in growth, tolerance to elevated temperatures and stress have been observed. Heritability estimates for growth are high, suggesting that the breeding programs can select fish for improved performance. The CGP has dramatically improved availability of genomic resources for cod. Approximately 158,000 sequences have been submitted to GenBank. Genes of interest
have been selected and development of a 20,000 element oligonucleotide microarray is underway. Development of gene-linked markers and a high density genetic map is ongoing. Marker identification has yielded >4,500 “predicted informative” SNPs and >140 microsatellite markers. A substantial investment has been made by Genome Canada and other partners to fund the CGP (www.codgene.ca). Resources developed by the CGP will enable marker assisted selection, and provide valuable tools for researchers interested in Atlantic cod.

Poster 3014

Title: Recent advances in the development of genomic resources in Crassostrea gigas: towards the sequencing of the Pacific oyster genome.

Presenting Author: Pierre Boudry, UMR M100 Physiologie et Ecophysiologie des Mollusques Marias, Ifremer - Technopole de Brest-Iroise, BP 70, 29280 Plouzane, France

Other authors (name only):
1. Patrick Gaffney
2. Elodie Fleury
3. Arnaud Huvet
4. Christophe Lelong
5. Pascal Favrel
6. Dennis Hedgecock

Abstract:
An international community, the Oyster Genome Consortium, uniting >70 biologists from 12 countries, presents the Pacific oyster Crassostrea gigas as a genome-sequencing candidate. In the USA and Europe, several EST sequencing projects have been successfully initiated by the Genoscope (Evry, France), the Joint Genome Institute (Walnut Creek, California) and the Max Plank Institute (Berlin, Germany), considerably increasing the number of ESTs. In Europe, the collaborative work developed within Marine Genomics Europe (www.marine-genomics-europe.org) and Aquafirst (aquafirst.vitamib.com) notably led to the development of (1) a cDNA microarray to study the functional genomics of survival during the reproductive period and (2) a dedicated EST database (www.sigenae.org/aquafirst). In the USA, inbred lines, F2 mapping families, SSR linkage maps, and transcriptomic analyses are aimed at understanding causes of growth heterosis. EST and BAC sequencing are currently being performed by the Joint Genome Institute. A BAC library made with DNA from a fourth generation inbred male was fingerprinted at the Genome Sciences Center (Vancouver, Canada) and the construction of BAC contigs-based physical map is well under way. It is now anticipated that the Institute of Oceanology (Chinese Academy of Sciences) and the Beijing Genomics Institute will sequence the oyster genome using the new generation sequencing technology.

Poster 3015

Genomic tools for breeding, fisheries and evolutionary biology of European Sea Bass

Filip AM Volckaert¹, Luca Bargelloni², Costas Batargias³, Adelino Canario⁴, Dimitris Chatziplis⁵, Alexander Chistiakov⁶, Richard Guyon⁷, Chris S Haley⁸, Bart Hellemans⁹, Georgios Kotoulas⁵, Heiner Kuhl¹⁰, Jacques Lagnel¹¹, Angelo Libertini¹⁰, Bruno Louro¹², Cécile Massault⁶, Richard Reinhardt¹⁰, Fabrice Senger¹, Erika Souche¹, Costas S Tsigenopoulos¹³, Guo Yuameti¹⁴, Francis Galibert⁷ (The Sea Bass Genomics Consortium)

¹ Laboratory of Animal Diversity and Systematics, Katholieke Universiteit Leuven, Charles Deberiotstraat 32, B-3000 Leuven, Belgium
² Department of Public Health, Comparative Pathology and Veterinary Hygiene, University of Padova, Viale dell’Università 16, Agropoli, I-35020 Legnaro, Italy
³ Nireus Chios Aquaculture S.A., Kardamila, GR-82300, Chios, Greece; present address: Faculty of Agricultural Technology, Department of Aquaculture and Fisheries, Technological Education Institute of Messolonghi, Nea Ktiria, GR-30200 Messolonghi, Greece
⁴ Centre of Marine Sciences, University of Algarve, Campus de Gambelas, P-8005-139 Faro, Portugal
⁵ Animal Breeding and Genetics, Department of Animal Production, School of Agricultural Technology, Alexander Technological Educational Institute of Thessaloniki, P.O.Box 141, GR-57 400 Sindos, Thessaloniki, Greece
⁶ Department of Molecular Diagnostics, National Research Center GosNIIgenetika, 1st Dorozhny Proezd 1, 113545 Moscow, Russia
⁷ UMR 6061 CNRS, Génétique et Développement, Faculté de Médecine, F-35043 Rennes Cedex, France
⁸ Roslin Institute (Edinburgh), Roslin, Midlothian EH25 9PS, Scotland, U.K
⁹ Hel lenic Centre for Marine Research (HCMR) Crete, Institute of Marine Biology and Genetics (IMBG), Thalassocosmos, P.O.Box 2214, Gournes Pediados, 715 00 Heraklion, Crete, Greece
¹⁰ Max-Planck-Institut für molekulare Genetik, Ihnestr. 63-73, D-14195 Berlin, Germany
¹¹ CNR – ISMAR, Istituto di Scienze Marine - Biologia del Mare, Riva 7 Martiri, 1364/A, I-30122 Venezia, Italy
Abstract
European sea bass (Dicentrarchus labrax L.) sustains a regional fishery and is commonly farmed in the Mediterranean basin; it has been subjected to genetic improvement for a short time. The Sea Bass Genomics Consortium developed a suite of new molecular tools over the past eight years. Genetic markers amount to 221 (176 mapped) AFLPs, 323 (196 mapped) microsatellites, more than 20 allozymes and an increasing number of SNPs. 30,000 EST traces from 14 cDNA tissue libraries have been generated. They have been used to prepare an updated linkage map of 374 markers and a radiation hybrid map of more than 1000 markers. The radiation hybrid panel of 92 cell lines was constructed from spleen tissue. A previously constructed BAC library has been assembled to a physical map of BAC end sequences. All three maps have been linked. The consortium has compared the genomes of European sea bass and several model teleosts, first as a modest undertaking based on 186 microsatellite flanking regions, but more recently by high-throughput BAC end sequencing and comparative mapping. A high percentage (>80%) of evolutionary conserved regions were spotted with the genome of the three-spined stickleback. Conservation of synteny dropped with phylogenetic distance as analysed in spotted pufferfish, fugu, medaka and zebrafish. Combining synteny data of stickleback and medaka the comparative BAC map of D. labrax has improved to a physical BAC map. The European sea bass genome is also in the process of being sequenced by whole genome shotgun (to date ~1.1 Gb / ~2 fold coverage). Based on the physical BAC map, the homologous group to stickleback chr 21 (~11.7 Mb) has been sequenced with high coverage through a combination of Sanger sequencing and pyrosequencing; fragments are being assembled. Based on a growth and stress experiments under commercial conditions, QTLs for growth have been detected and several BACs related to the QTLs have been sequenced. Additional BACs for immunological traits have been sequenced and the sequence assembled. Our findings contribute to a better understanding of fish evolution, provide excellent prospects for evaluating variation in the genome and enhanced selection in sea bass aquaculture, provide tools for tracing of fisheries products and open new perspectives towards understanding the application and effects of natural selection on locally adapted natural populations.

Poster 3016
Title Towards selective breeding of the Silver-lip pearl oyster (Pinctada maxima) understanding genotype by environment (GxE) interactions
Presenting Author: Dean R. Jerry, Aquaculture Genetics Research Program, School of Marine and Tropical Biology, James Cook University, Townsville, Qld 4810, Australia
Other authors (name only):
1. Renate Kvingedal
2. Brad. S. Evans

Abstract:
Before selective breeding involving a new aquaculture species it is important to identify populations/strains to be used as a founder genetic base and that possess “superior” characteristics for commercial production. Also of interest is an understanding of how phenotypic expression of important traits is influenced by local environmental conditions (genotype by environment interactions). Currently, there is interest in the selective breeding of the silver-lipped pearl oyster P. maxima in Indonesia. Before a breeding programme commences for this species we need to identify populations on which to base the founder stock and because improved oysters will be grown at multiple sites, determine the influence local environmental conditions have on the expression of oyster growth and pearl quality. In this experiment we produced spat from 27 full-sib families derived from three wild Indonesian P. maxima populations (Bali, Raja Empat, Aru) and communally stocked them at two different grow-out sites. Microsatellite-based DNA parentage analyses were used to assign oysters to their family and population of origin and growth and morphometric data obtained at 6,12 and 18 months. Results from this experiment will be presented and the implications of possible genotype by environment interactions on a future selective breeding programme for P. maxima discussed.

Poster 3017
Title: Status of Channel Catfish Genomics
Presenting Author: Sylvie Quiniou, ARS-USDA / CGRU, 141 Experiment Station Rd Stoneville, MS 38776

Abstract:
Channel catfish production is now the leading aquaculture species in the U.S., with 600 millions pounds processed annually. To support selective breeding programs, molecular tools are being developed to help characterizing the catfish genome and identify genomic regions that control important production traits. Briefly, framework genetic linkage maps have been produced based on microsatellite loci and Amplified Fragment Length Polymorphism loci. 400,000 Expressed Sequence Tags have been identified from several tissues and cell lines, and will be clustered and annotated in the Gene Index Project. One cDNA microarray and one high density
oligonucleotide array have been developed for global gene expression studies. Three large-insert catfish Bacterial Artificial Chromosome (BAC) libraries have been produced. More than 57,000 BAC ends have been sequenced. A physical map has been generated to integrate the genetic map with catfish chromosomes, allowing for fine mapping of phenotypic trait alleles such as Quantitative Trait Loci, effective positional cloning of genes controlling economically important traits to improve germplasm and for comparative genomic analyses. The physical map will also provide a minimal tiling path for a whole genome sequencing project.

Poster 3018

Title: Developing genomic resources for new and emerging aquaculture species in New Zealand

Presenting Author: P. J. Fisher, Invermay Agricultural Centre, Private Bag 50034, Mosgiel 9053, New Zealand

Other authors (name only):
J. E. Symonds
S.P. Walker
R.C. Rickman
R. McLaren
R. Anderson
K.M. McEwan

Abstract:
Commercial aquaculture, with notable exceptions, has been slower than terrestrial livestock industries to invest in selective breeding. Recognizing the value of aquaculture genetics, NIWA has initiated a program to build broodstock resources for three high value species: yellowtail kingfish (Seriola lalandi), groper (Polyprion oxygeneios) and paua (abalone, Haliotis iris). The goal is to identify and select high performing broodstock individuals by evaluating their offspring through multiple crosses. Key performance traits of commercial value such as growth, product quality and yield, will be assessed in these families. Species specific methodologies are being developed to manage genetic diversity within the broodstock and identify the parentage of the evaluated offspring. For paua, individual controlled 1:1 family crosses have been established and one significant issue is whether these families should be reared separately or pooled. For large marine finfish such as kingfish and groper, communal spawning is standard, but this provides little control over individual matings. Therefore, determining parentage (with DNA markers) is essential for all three species. We describe the identification of highly variable microsatellites in each species (using long read, single pass 454 sequencing technology) and the downstream procedures for developing them into parentage testing panels.

Poster 3019

Title: Preliminary linkage map of Haliotis midae based on AFLP and microsatellite markers

Presenting Author: Rouvay Roodt-Wilding, Department of Genetics, University of Stellenbosch, Stellenbosch, South Africa

Other authors (name only):
1. Daleen Badenhorst
2. Peizheng Wang

Abstract:
This study presents first-generation preliminary linkage maps for Haliotis midae, the only commercially important abalone species found in South Africa. Linkage maps were constructed using AFLP and microsatellite markers segregating in a F1 family following a pseudo-testcross mapping strategy. Twelve AFLP primer combinations, which produced 573 segregating bands and 10 microsatellite markers, were genotyped in the parents and 108 progeny of the mapping family. Of the 573 segregating AFLP peaks genotyped, 241 segregated in a 1:1 ratio and 332 in a 3:1 ratio. Of these AFLP markers, 90 segregated according to the expected 1:1 Mendelian ratio and 164 segregated according to the expected 3:1 Mendelian ratio at the P = 0.05 level and were used for linkage analysis. Of the 10 microsatellite markers genotyped, nine were informative for linkage mapping analysis. A total of 12 and 10 linkage groups were detected for the female (601.6cM; 41 markers) and male map (431.9cM; 33 markers). This linkage map forms a starting point for more detailed study of the H. midae genome and will act as a scaffold for basic and applied studies in abalone and initiate the mapping of quantitative trait loci (QTL) of commercially important traits and future marker assisted selection (MAS) selective breeding programmes.

Poster 3020

Title: QTL for resistance against Infectious Pancreatic Necrosis in Atlantic salmon.

Presenting Author: Thomas Moen, Nofima, PO Box 5010, N-1432 Aas, Norway

Other authors (name only):
1. Matthew Baranski
2. Anna Sonesson
3. Sissel Kjoeglum

Abstract:

A genome scan was used to identify QTL for resistance to Infectious Pancreatic Necrosis (IPN) in Atlantic salmon, a major cause of mortalities and losses in the salmon farming industry and a potential target for marker-assisted selection. The disease affects fry as well as post-smolt. The genome scan, performed on 10 large full-sib families of challenge-tested post-smolt, revealed a single major QTL explaining 80% of the genetic variation, in addition to one other experiment-wide significant QTL (1% of genetic variation). The major QTL was highly significant in 10 of 20 mapping parents and was mapped to a 3 cM interval. Challenge-tested fry, siblings of the investigated post-smolt, were used to show that the QTL had the same effect on fry as on post-smolt. Subsequently, ~100 full-sib groups off fry were genotyped for a larger set of markers, in order to fine-map the QTL, possibly detecting population-level associations, and to determine linkage phases in QTL heterozygous broodstock animals. The results from fine-mapping will be presented at the conference. The QTL is identical to the one detected by Houston et al. (Genetics 2008), and has been implemented in the breeding programme of Aqua Gen Ltd as of 2007.
**Poster 4001**

**Title:** Search for the imprinting mark required for parent-of-origin dependent \(H19\)ICR methylation using two different mouse models

Presenting Author: Claudia Gebert, National Institutes of Health (NIH), Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD), Program on Genomics of Development (PGD), Bldg. 6B, Room 2B206, 9000 Rockville Pike, Bethesda, MD 20892 (USA)

Other authors (name only):
1. Karl Pfeifer

Abstract:
A 2.4 kb DNA region, the \(H19\) imprinting control region (\(H19\)ICR), lies between the Insulin-like growth factor 2 (Igf2) and \(H19\) genes on mouse chromosome 7, and is important for normal imprinting of the two genes.

Interestingly, the \(H19\)ICR becomes only methylated in sperm. However, the mark required for paternal specific methylation is still not known. To learn more about the nature of this imprinting mark, we first wanted to know whether it is linked to the \(H19\)ICR, and whether this 2.4 kb region alone is sufficient for imprinting. We addressed these questions by generating two mouse models that carry the \(H19\)ICR upstream of the Alpha fetoprotein (Afp) on chromosome 5 and between the CD3 gamma and delta genes on chromosome 9, respectively. This approach allows us to study the unique function of the \(H19\)ICR outside an imprinting environment. Methylation analyses revealed that while the \(H19\)ICR becomes normally methylated on only the paternal allele at both exogenous loci in somatic tissues, it stays unmethylated in sperm. Thus, the imprinting mark is linked to the \(H19\)ICR, and the 2.4 kb region is sufficient to imprint somatic tissues. However, DNA methylation is not the imprinting mark that distinguishes the two parental alleles.

**Poster 4003**

**Title:** Establishment of an in vivo model for canine prostate cancer and development of an AAV mediated gene therapeutic approach

Presenting Author: H Murua Escobar, Small Animal Clinic, University of Veterinary Medicine, Bischofsholer Damm 15, 30173 Hannover, Germany

Other authors (name only):
1. JT Sollek
2. M Fork
3. S Willenbrock
4. KA Sterenczak

Abstract:
The dog (Canis lupus familiaris) is the only mammalian species besides the man that spontaneously develops cancer of the prostate and thus is estimated as a unique model for the human neoplasia. The aim of this study was to develop an in vivo animal model that mimics the behaviour of an original tumour as basis for the development of therapeutic approaches. We used HMGA2 gene expression as marker due to the fact several human and canine tumours -including canine prostate carcinomas- were reported to show a correlation between their dignity and the expression of HMGA2 gene. We derived the cell line CT1258 from a highly malignant canine adenocarcinoma of the prostate showing high expression of HMGA2 and inoculated 1x10\(^E6\) cells in 9 NOD-SCID mice s.c. as well as 1x10\(^E5\) and 5x10\(^E5\) cells respectively i.p.. The induced tumours emulated the original tumour regarding biological behaviour, HMGA2 expression, and the presence of cytogenetic markers. Finally, we developed a gene therapeutic approach using recombinant HMGA antisense adeno-associated viruses (rAAV) resulting in significant inhibition of cell proliferation and apoptosis of CT1258 cell in vitro.

**Poster 4004**

**Title:** Bovine \(PRNP\) polymorphisms in Asian native populations and Japanese breeds

Presenting Author: George Msalya, Takeshi Shimogiri*, Shin Okamoto
Kotaro Kawabe, Mitsuru Mizezawa, Takao Namikawa, Yoshizane Maeda

Other authors (name only):
1. J. Bullerdeg
2. N. Reimann-Berg
3. M. Dorsch
4. H. Hedrich

Abstract:
The dog (Canis lupus familiaris) is the only mammalian species besides the man that spontaneously develops cancer of the prostate and thus is estimated as a unique model for the human neoplasia. The aim of this study was to develop an in vivo animal model that mimics the behaviour of an original tumour as basis for the development of therapeutic approaches. We used HMGA2 gene expression as marker due to the fact several human and canine tumours -including canine prostate carcinomas- were reported to show a correlation between their dignity and the expression of HMGA2 gene. We derived the cell line CT1258 from a highly malignant canine adenocarcinoma of the prostate showing high expression of HMGA2 and inoculated 1x10\(^E6\) cells in 9 NOD-SCID mice s.c. as well as 1x10\(^E5\) and 5x10\(^E5\) cells respectively i.p.. The induced tumours emulated the original tumour regarding biological behaviour, HMGA2 expression, and the presence of cytogenetic markers. Finally, we developed a gene therapeutic approach using recombinant HMGA antisense adeno-associated viruses (rAAV) resulting in significant inhibition of cell proliferation and apoptosis of CT1258 cell in vitro.
We genotyped six polymorphic sites of a bovine prion gene (PRNP) and estimated haplotypes in Asian native populations and Japanese breeds. Six sites were 23-bp indel within promoter, 12-bp indel within intron1, nonsynonymous SNPs (K3T and S154N), octapeptide repeats within ORF and 14-bp indel within 3'UTR. At the 23-bp indel, the frequency of - (deletion) allele (0.55-1.00) was higher than that of + (insertion) allele in all groups except Bos frontalis. 12-bp indel frequency ranged from 0.00 to 0.93 for + allele and from 0.07 to 1.00 for - allele. + allele frequency was higher than - allele in all Asian populations except the Mongolian. In most Japanese breeds, - allele frequency was higher than + allele. For the 14-bp indel, + allele frequency (0.58-1.00) was higher than - allele. 6-repeats allele was common allele at the octapeptide repeats site. 5/5 genotype was detected in Japanese black and the Mongolian population. 4-repeats allele was detected only in Bos frontalis. K3T site was polymorphic in all Asian populations except the Mongolian. S154N was polymorphic in all Asian populations. We estimated 30 different haplotypes from these genotypic data. A ‘23-12-K6S14+’ haplotype was found in all groups with frequencies from 0.05 to 1.00.

Poster 4006

Title: Development of DNA markers for discrimination between Japanese and Australian beef cattle

Presenting Author: Yutaka SUEKAWA, Laboratory of Animal Breeding and Genetics, Graduate School of Agricultural Science, Kobe University, Kobe, Japan

Other authors (name only):
1. HAZUKI MUTO
2. KANAKO YOSIZAWA
3. SINJI SASAZAKI
4. Fumio MUKAI
5. Hideyuki MANNEN

Abstract:
In the meat industry, correct breed information in food labelling is required to assure meat quality. As an element of traceability system, genetic markers are useful to identify breeds. The aim of this study was to develop the DNA markers discriminating between Japanese and Australian beef. SRY and ND5 were used as gene markers specific to Bos indicus and MC1R gene as to red coat color. In addition, amplified fragment length polymorphism (AFLP) method was employed to develop additional markers. A total of 1696 AFLP primer combinations provided four candidate polymorphisms that were converted into single nucleotide polymorphisms (SNPs) markers for high-throughput genotyping. The allele frequencies in cattle from both countries were investigated for estimating the discrimination ability. The allele frequencies in Australian beef were 0.194 (SRY), 0.471 (ND5) and 0.410 (MC1R). The allele frequencies of AFLP-SNPs ranged from 0.000 to 0.001 in Japanese cattle and 0.032 to 0.094 in Australian beef. Combining these seven markers, the probability of identifying Australian beef was 0.909 and probability of misjudgement was 0.008. The genetic system using these DNA markers could be a
powerful tool for breed identification and would contribute to prevent falsified display beef in Japan.

**Poster 4007**

**Title:** Discovery of porcine miRNAs through comparative genomics approach and profiling from skeletal muscle tissues during development

Ting-Hua Huang, Meng-Jin Zhu, Xin-Yun Li, Shu-Hong Zhao

*Key Lab of Agricultural Animal Genetics, Breeding, and Reproduction of Ministry of Education, Huazhong Agricultural University, Wuhan, 430070, PR China.

**Abstract:**

MiRNA (microRNA) plays critical roles in many important biological processes such as growth & development in mammals. In this study, we used in silico prediction followed by microarray hybridization to identify porcine miRNAs. Alignment of porcine genomic sequence against human, mouse and rat miRNA stem-loop sequences identified 775 predicted porcine miRNAs. An initial screen by microarray hybridization with RNA samples prepared from the 33 days whole embryo and extra embryo membrane validated the expression of 296 of these candidates. To investigate the decisive miRNAs involved in muscle growth & development processes, we subsequently examined the miRNA expression profiles in longissimus muscle tissues collected from 33 days, 60 days of gestation fetuses and adult pigs. A total of 140 miRNAs were differentially expressed among the ages investigated. The differentially expressed miRNAs were classified into four categories of distinctive expression patterns, suggesting possible involvement of special biological processes respectively. Five of the differentially expressed miRNAs detected by the arrays were validated by real-time PCR method. Further bioinformatics analysis of the miRNA-mRNA interaction sites suggested that the potential mRNA targets of the differentially expressed miRNAs may play important roles in muscle growth & development. Our study have expanded the miRNA catalog of the pig, and for the first time, provided miRNA transcriptome landscape of the prenatal and adult porcine skeletal muscle. The novel data and the bioinformatics analysis presented in this study may facilitate the hypothesis-based functional studies of miRNAs in muscle.

**Poster 4008**

**Title:** Muscle Fiber Typing of the Blind Subterranean Mole Rat Reveals Adaptive Features to Sustained Activity Under Hypoxia

Presenting Author: Mark Band, University of Illinois, 1201 W. Gregory Dr., Urbana, IL, 61801

Other authors (name only):
1. Alma Joel
2. Pessia Shenzer
3. Raymond Coleman
4. Aaron Avivi

**Abstract:**

Mammalian muscles are composed of three primary types of fibers, type I oxidative, type IIa oxidative and type IIb glycolytic. Most mammalian species present a distribution of the various fiber types allowing plasticity for adaptation to changing physiological conditions. Mole rats of the Spalax ehrenbergi superspecies are blind subterranean rodents that require intense activity under hypoxic conditions for survival. Fiber typing of neck (trapezius) and leg (gastrocnemius, quadriceps) muscles using standard histochemical techniques (succinic dehydrogenase, myosin ATPase) showed that the muscle fibers of mole rats in natural settings, as well as after extended captivity, were predominantly type IIa. The same muscles in laboratory rats showed the full range of fiber types. In contrast, the hearts of the mole rats and the laboratory rats were very similar. Previous investigations have shown a correlation of VEGF levels to a shift from type IIb to type IIa in mice under intense endurance activity. VEGF levels in mole rat skeletal muscle are naturally elevated as compared to other terrestrial mammals. Our results indicate that skeletal muscle in the mole rat appears to have evolved in response to specific environmental demands to permit intensive endurance burrowing activities under conditions of severe or chronic hypoxia.

**Poster 4009**

**Title:** The association between Split Upper Eyelid and multiple horns observed in sheep is maintained in multihorned goats.

Presenting Author: M.Teresa Tejedor Hernández, Department of Anatomy, Embriology and Genetics, Faculty of Veterinary Sciences of Zaragoza (Spain), C/Miguel Servet 177. 50013-ZARAGOZA (Spain)

Other authors (name only):
1. Luis V. Monteagudo
2. Isidro Sierra
3. Mariano Herrera

**Abstract:**

We have recently detected the existence of autochthonous populations of multihorned goats in the
region of Extremadura (Spain). The flocks contain around 7% multihorned females, since multihorned males are not selected for reproduction. Our first observations clearly indicate that this character is associated to Split Upper Eyelid (SUED). This was a well known association in several multihorned sheep breeds, such as Hebridean, Churro Navajo and Jacobs, but scientific reports on caprine populations carrying it were not published. As in sheep, caprine SUED shows a wide range of morphological alterations: Slight changes in the position of the eyelashes, small notches and finally, in a small proportion of individuals, completely split eyelids, exposing most of the eye. Multiple horns and SUED are completely associated, as the result of the observation of the modified eyelids at birth is completely concordant with the future eruption of multiple horns. At present, even if the exact inheritance model for this character remains undisclosed in both sheep and goat, its complete conservation in both species seems evident, suggesting that it appeared prior to their evolutionary segregation.

Poster 4010

Title: Phylogeny relationship of GDF5 sequence between a number of mammalian species.

Presenting Author: Tomasz Ząbek, National Research Institute of Animal Production, Immuno- and Cytogenetics Department, Krakowska 1, 32-083 Balice, Poland

Other authors (name only): 1. Ewa Słota

Abstract:
Growth differentiation factor 5 (GDF5) is a member of the bone morphogenetic protein (BMP) family and the TGF-beta super family. The members of this family are regulators of cell growth and differentiation in both embryonic and adult tissues. 441 base pairs exonic sequence of GDF5 gene, including GCA repeats was compared between 13 species belonged to Bovidae, Cervidae, Canidae, Suidae, and Equidae family and human as reference for primer design. Sixty nine single nucleotide differences were found in GDF5 sequence when compared between all species. Except to the region spanning tandem repeat of dog, (GCA)nGTA(GCA)n structure were common for GDF5 repeat region of all species, with the largest (GCA)n motive in domestic horse, Przewalski horse, and onager. Based on phylogeny distance data GDF5 sequence is the most variable inside Bovidae family in relation to other investigated taxa.

Poster 4011

Title: Genomic organisation and mapping of the canine High Mobility Group A1 (HMGA1) gene

J.T. Soller1,2, C. Beuing1, M. Muth3, S. Wagner2, G. Dolf3, C. Schelling4, A. Richter2, S. Willenbrock1,2, N. Reimann-Berg1,2, S. Winkler2, I. Nolte1, J. Bullerdiekel, H. Murua Escobar1,2

1Small Animal Clinic, University of Veterinary Medicine Hannover, Germany. 2Centre for Human Genetics, University of Bremen, Germany. 3Institute of Animal Genetics, Nutrition and Housing, University of Berne, Berne, Switzerland. 4Department of Animal Sciences, Swiss Federal Institute of Technology Zurich and Vetsuisse Faculty Zurich, University of Zurich, Switzerland.

Abstract:
The high mobility group A1 proteins (HMGA1a/HMGA1b) are highly conserved between mammalian species and widely described to participate in various cellular processes. In humans, chromosomal aberrations affecting the HMGA1 gene locus on HSA 6p21 were described for various benign mesenchymal tumours, while high titres of HMGA1 proteins were shown to be associated with the neoplastic potential of various types of cancer. Due to the various similarities in biology and presentation of human and canine cancer, knowledge about the structure of cancer related genes and proteins is of significant value. Herein we report the characterisation of the genomic structure of the canine HMGA1 gene consisting of 7 exons and 6 introns spanning in total 9524bp, the in vivo localisation of the HMGA1 protein to the nucleus, and chromosomal assignment of the gene by FISH to CFA12q11. Additionally, we evaluated a described canine HMGA1 exon 6 SNP in 55 Dachshunds. The performed characterisations will allow doing comparative analyses of aberrations affecting the human and canine gene and protein as basis for revealing mechanisms involved in HMGA1 related pathogenesis in both species.

Poster 4012

Title: New loci and allele characterization of pig MHC (SLA) classical class I gene

Presenting Author: Maiko Tanaka-Matsuda, Institute of Society for Techno-innovation of Agriculture, Forestry and Fisheries, 446-1 Kamiyokoba, Ippaizuka, Tsukuba, Ibaraki 305-0854, Japan

Abstract:
Swine MHC (SLA) class I genes are classified into two groups, classical and non-classical ones. In pigs, the role for presentation of intracellular peptidic antigens to T cell receptors is mainly played by three types of class I antigens, SLA-1, SLA-2 and SLA-3. To date, 44 alleles have been reported for SLA-1 locus, and the number of SLA-1 alleles is considered to increase by further analysis. On the other hand, there are several observations for duplication of SLA-1 locus in single haplotypes, and some haplotypes seem to lack expressed SLA-1. We classified the SLA-1 alleles detected in newly sequenced SLA haplotypes in addition to the alleles known to date. One of the haplotypes we newly sequenced had two loci sharing characteristics with the known SLA-1 alleles. Interestingly, we found loci that possessed some features distinguished from SLA-1, although they shared several characteristics with SLA-1, which have not been observed in SLA-3 or other SLA loci. Sequences belonging to this type were also observed in dbEST for pigs. We have further to investigate locus organization of SLA classical class I genes on the genomic sequence in various haplotypes to build the appropriate classification of SLA loci and alleles.

Poster 4014
Title: Evidence for Adaptive and Negatively Selected Chromosome Rearrangements in Amniote Evolution
Presenting Author: Denis M. Larkin, Department of Animal Sciences, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA.
Other authors (name only):
1. Greg Pape
2. Ravikiran Donthu
3. Loretta Auvil
4. Michael Welge
5. Harris A. Lewin

Abstract:
The genome assemblies of 10 amniotes were compared at the level of chromosome organization; human, chimpanzee, macaque, cattle, pig, dog, rat, mouse, opossum, and chicken. Blocks of homologous synteny (HSBs) and evolutionary breakpoint regions (BR) were identified from pairwise comparisons of all genomes. The rates of rearrangements were compared. After the divergence from marsupials, an elevated rate of chromosomal rearrangements was observed in placental mammals (~1.3 breakpoint per My), followed by a decrease (~0.8 breakpoint per My) until the Cretaceous–Tertiary (K-T) boundary. After the K-T boundary the rate of chromosomal rearrangements in placental mammals increased to ~1.4 breakpoint per My. The rearrangement rate in marsupials is ~0.8 breakpoints per My, in birds it is ~0.7 breakpoints per My. We found that large HSBs conserved in all species contain significantly more genes controlling essential processes of organismal development than the genome average. One of these regions, a 23.6 Mbp HSB on HSA2, contains the HOXD cluster. In contrast, BRs contain a high frequency of genes associated with adaptive responses to the environment and sequence features that are characteristic of evolutionary hotspots. We propose that natural selection plays an important role in determining structural and functional features of amniote chromosomes.
Poster 4015

Title: Three groups of Endogenous Retroviruses found in Crocodilians

Weerachai Jaratlerdsiri1, Chandramaya Siska Damayanti1, Lee Miles1, Sally Isberg2, Lorna Melville3, Chris Moran1, and Jaime Gongora1.*

1Centre for Advanced Technologies in Animal Genetics and Reproduction, Faculty of Veterinary Science, RMC Gunn Building B19, University of Sydney, NSW 2006, Australia.
2Porosus Pty Ltd, PO Box 86, Palmerston NT 0831, Australia.
3OIC Berrimah Veterinary Laboratories, Department of Primary Industry, Fisheries and Mines, GPO Box 3000 Darwin 0801.
*Corresponding author: j.gongora@usyd.edu.au

Abstract
Two families, Crocodylidae, and Alligatoridae are unambiguously recognised, with gharials either lumped with Crocodylidae or assigned to a separate family Gavialidae. Endogenous retroviruses (ERVs) are copies or remnants of exogenous retroviruses that were integrated into the host genome at some stage in the past. ERVs are vertically transmitted from the host to its progeny. Although most ERVs are defective due to inactivating mutations, functional ERVs are potential agents of disease. Previous studies have identified the presence of a distinct clade of ERV from six species of crocodilians. Here we analyse the functionality, distribution and phylogenetic relationships of ERVs in thirteen species of Crocodylidae and seven species of Alligatoridae from across the world. The ERV reverse transcriptase (pol) gene fragment (0.8-1 kb) was amplified, cloned and sequenced. Preliminary analyses of sixty-six crocodilian ERV pol DNA sequences show that these retroelements possess stop codons and deleterious mutations. Thus, crocodilian ERVs are generally, if not universally, defective as has been observed in many other hosts. Phylogenetic analyses show that crocodilian ERVs are generally, if not universally, defective as has been observed in many other hosts. Phylogenetic analyses show that crocodilian ERVs cluster in three major clades, one Crocodylidae-specific, one Alligatoridae-specific and one found in all Crocodilians including gharials. Further analyses of the evolutionary relationships and evidence for ERV functionality within crocodilians are underway.

Poster 4016

Title: A yeast-based assay to detect TP53 functional mutations: study on spontaneous feline tumors.

Presenting Author: Paola Modesto Istituto Zooprofilattico Sperimentale del Piemonte, Liguria e Valle d’Aosta- P.zza B.go Pila 39/24 16129 Genova, Italy

Other authors (name only):
1. Alberto Inga.
2. Virginia Andreotti.
4. Umberto Cardellino.
5. Angelo Ferrari.
6. Claudio Petterino.
7. Alessandra Ratto.

Abstract:
In feline neoplasms little is known about occurrence and nature of p53 mutations, although the few available data suggest a lower frequency compared to humans.

In order to understand the involvement of p53 mutations in feline carcinogenesis, we previously developed an yeast-based approach revealing that feline p53 (fe_p53) can be studied in this model system and is equally susceptible to inactivating mutations as human p53.

In this report we examined 30 bioptic samples from spontaneous feline tumors using the yeast-based assay to score p53 mutations based on the frequency of phenotypically mutant fe_p53 cDNA alleles individually examined upon in-vivo cloning. In this assay non-transactivating p53 mutations result in the appearance of red colonies while white colonies are instead indicative of active p53.

Four samples resulted in a frequency of red colonies higher than 30% indicative of the presence of clonal p53 mutations in the biopsy. Sequencing of these cDNAs identified a splice donor site mutation (G>A) resulting in an 11 amino-acid insertion (exon 6) in an hepatocarcinoma and an in-frame deletion of 12 nucleotides in a schwannoma (exon 5 starting at codon 167). Direct DNA sequencing of genomic DNA is in progress and definitive results will be presented in poster.

Poster 4017

Title: Comparative genomics of Toll-like receptor signalling reveals candidate genes for disease resistance traits

Presenting Author: Oliver Jann, The Roslin Institute and R(D)SVS, University of Edinburgh, Roslin, Midlothian, EH25 9PS, Scotland

Other authors (name only):
1. Susan I. Anderson
2. Kirsty Jensen
3. Tahar Ait -ali
4. Chunhua Wu
5. Noelle E. Cockett
6. Elizabeth J. Glass
Abstract:
Studies in human and mouse indicate that polymorphism in genes involved in Toll-like receptor (TLR) signalling can affect immune related traits and thus might explain part of the observed variation in disease resistance of livestock. Here we use an interspecies comparative approach to investigate the importance of 20 TLR related genes for resistance traits by associating their genomic localisation with previously published immune-related QTL regions and by analysing their transcript levels in breeds of divergent immune-related phenotypes. We report the localisation of these genes in cattle, sheep and pig and compare their positions to the corresponding genes in human and mouse. Several of the loci are located within regions affecting variation of related health traits in different species. Polymorphism in some of the genes have been directly associated with effects on relevant traits in humans and mice. Many of the genes display divergent transcript levels in animal breeds of different resistance status. Thus, this intergeneric approach has identified several strong candidate genes for a number of important immune traits like clinical mastitis, tolerance to Trypanosoma infection and general disease resistance in pig, sheep and cattle. Further studies are required to investigate the potential role of polymorphism within these genes and their direct effects on resistance traits.

Poster 4018

Title: SPECIES IDENTIFICATION OF MEAT AND ESTIMATING THE PREVALENCE OF BUSH MEAT UTILIZATION IN RAW MEAT OUTLETS USING CYTOCHROME B SEQUENCES.

Presenting Author: … KARISA BRIAN KAHINDI, ILRI P.O BOX 30709, NAIROBI
b.karisa@cgiar.org

Other authors (name only):
1. CHARLES, N. KIMWELE
2. ROB SKILTON
3. JOSEPH JUNGA
4. OLIVIER HANOTTE

Abstract:
In Africa, identification of poached animal species relies greatly on morphological features, (Malisa et al. 2005). In cases when the morphologically variable features are absent in confiscated carcasses, the accuracy in the identification reduces. DNA and protein based methods have been used in the identification of species with varying success rates. DNA techniques using either nuclear or mitochondrial DNA markers show greater potential for use in conservation and wildlife management (Cronin et al., 1991). Of the two, mitochondrial DNA analysis usually serves as a more powerful tool for detecting population genetic structure due primarily to its maternal mode of inheritance, lack of recombination, high mutation rate and the process of lineage extinction within populations (Brown et al., 1982).

In this study, the species of 565 randomly sourced samples of raw meat were determined by PCR amplification of genomic DNA using universal Cytochrome b primers, (Verma and Singh, 2003). The PCR products were sequenced and the nucleotide sequences were run on a BLAST search of sequence database in NCBI, (www.ncbi.nlm.nih.gov).

The 550 samples analysed in this study were all identified as domestic animal species. Significant substitution of animal species was observed especially in the substitution of goat meat with sheep meat. The substitution between species is a clear indication that there is fraud with possible economic and health implications.

Poster 4019

Title: Identification and characterization of a perfect mono-nucleotide microsatellite within 16S rRNA gene in chicken mtDNA sequence

SHENGGUO ZHAO1, 2
1CAAS-ILRI Joint Laboratory on Livestock and Forage Genetic Resources, Institute of Animal Science, Chinese Academy of Agricultural Sciences (CAAS), No. 2, Yuan Ming Yuan Xi Lu, Haidian District, Beijing 100193, China
2College of Animal Science and Technology, Gansu Agricultural University, Lanzhou 730070, Gansu, China

Other authors (name only):
1. OLIVIER HANOTTE
2. JIANLIN HAN

Abstract:
An alignment of 13 chicken and red junglefowl complete mtDNA sequences reveals a perfect mono-nucleotide microsatellite repeat unit consisted of between 8 and 13 Cs within the 16S rRNA gene in seven samples, however, the remaining six sequences have an interrupted non-polymorphic fragment ((C)nT(C)m). To further characterize this microsatellite, 350 samples of indigenous chickens, commercial breeds and wild red junglefowls were
genotyped. Thirty-five samples were also sequenced for the same region. Both genotyping and sequencing results confirmed the presence of these two types of structures in chicken mtDNA genome, with the wild red junglefowl samples showing the interrupted fragment. Among the domestic chickens analysed, the two structures show a clear geographic distribution pattern. The perfect repeats are detected in indigenous chickens from West Asia including Bangladesh, Myanmar and Pakistan and in the majority of commercial fowls. The interrupted fragment occurs among indigenous chickens from South and South-East Asia including China, Korea and Papua New Guinea. This perfect mono-nucleotide microsatellite, the first of its kind being validated in the avian mitochondrial genome, could be useful in providing insights into the evolutionary mechanisms of polymorphism and/or instability of mtDNA microsatellites and a better estimation of mutation rate of such microsatellites in birds.

Poster 4020

Title: Sequences of MCW0371 reveal complex structures having alleles of either perfect mono-nucleotide repeats or compound microsatellite repeats

Presenting Author: JIANLIN HAN1,3

1 CAAS-ILRI Joint Laboratory on Livestock and Forage Genetic Resources, Institute of Animal Science, Chinese Academy of Agricultural Sciences (CAAS), No. 2, Yuan Ming Yuan Xi Lu, Haidian District, Beijing 100193, China

3 International Livestock Research Institute (ILRI), P.O. Box 30709, Nairobi 00100, Kenya

Other authors (name only):
1. LINA JIN1,2

2 CAAS-ILRI Joint Laboratory on Livestock and Forage Genetic Resources, Institute of Animal Science, Chinese Academy of Agricultural Sciences (CAAS), No. 2, Yuan Ming Yuan Xi Lu, Haidian District, Beijing 100193, China

2 College of Animal Science and Technology, Gansu Agricultural University, Lanzhou 730070, Gansu, China

2 SHENSCHENG ZENG1

1 CAAS-ILRI Joint Laboratory on Livestock and Forage Genetic Resources, Institute of Animal Science, Chinese Academy of Agricultural Sciences (CAAS), No. 2, Yuan Ming Yuan Xi Lu, Haidian District, Beijing 100193, China

2 OLIVIER HANOTTE3

3 International Livestock Research Institute (ILRI), P.O. Box 30709, Nairobi 00100, Kenya

Abstract: A total of 378 samples from nine indigenous chicken populations and three commercial breeds were genotyped using MCW0371 locus, a microsatellite mapped within MHC-B region on GGA16. Twelve alleles between 199 and 210 bp were detected. Some genotypes exhibited peaks that had a pattern similar to those of perfect mono-nucleotide repeats; however, irregular patterns predominated in the samples and it was impossible to determine their allele sizes accurately. To verify these results, PCR products from 24 samples with likely homozygous genotypes were selected for direct sequencing. Most of them turned out to be heterozygous. To further validate the sequence structure at this locus, 45 samples were chosen for sequencing after cloning. A total of 49 haplotypes were identified with two structures in their repeat units: the first being perfect mono-nucleotide repeats having 14 to 20 ‘As’; and the second, a compound microsatellite having 1 to 6 ‘AAC’ repeats followed by 9 to 17 ‘As’. In addition, some SNPs and in/dels were present in flanking and repeat regions. These structures tend to complicate the accurate determination of allele sizes at this locus when using conventional capillary electrophoresis. This calls for thorough validation of such markers before wide application in genetic studies.

Poster 4021

Title: Sequencing of full-length 5’ end of canine Ataxia Telangiectasia Mutated cDNA and characterization of the promoter region

Presenting Author: Maria Elena Turba; Genefast srl, Via della Pace 33/a 41050 Castelnuovo Rangone Modena, Italy

Other authors (name only):
1. Fabio Gentilini
2. Monica Forni
3. Stefano Cinotti

Abstract: The predicted canine ATM locus is annotated in a canine chromosome 5 region (Dog Genome assembly 2.0, Ensembl release 49) syntenic with human chromosome 11. Using 5’ RACE PCR, cloning and sequencing of mRNA purified from canine blood, the transcription start site of canine ATM was found on the reverse strand at CFA5: 27307661. To predict the putative promoter region, 700 bp genomic sequences upstream of the 5’ cap site were analyzed using Proscan software v1.7. A putative TATA-less bi-directional promoter region was found in the 371bp region upstream of the cap site. The core promoter harbours different conserved regulatory motifs: CREB, CCAAT boxes, SP1, GCF, XRE, ETS, Cre and c-Myb. When compared to the human and pig ATM/NPAT promoter, the canine cis-regulatory elements are highly conserved.
Two exons, not annotated in the public database, were found in the 5’UTR. The cloned cDNA fragment extended from the transcription start site to exon 6. The putative translation start codon (XM_845871.1) corresponding to the N-terminus of human isoform 1, was found in exon 3. Two additional in-frame Met codons were located downstream in exons 5 and 6. The cloning of 5’ UTR could allow the investigation of the alternative splicing of canine ATM N-terminus.

**Poster 4022**

**Title:** Analysis of non-orthologous genes between cow and the genomes of other species.

Presenting Author: John L. Williams, IDRA, Parco Tecnologico Padano, Via Einstein, Lodi 26900, ITALY

Francesco Strozzi¹, Raffaele Mazza², Sem Genini³, Dimos Kapetis³, Francesca Panzitta³, Roberto Malinverni³, Paolo Ajmone-Marsan³ and John L. Williams¹

¹IDRA Parco Tecnologico Padano, Via Einstein, Polo Universitario, Lodi 26900, ITALY  
²Istituto di Zootecnica, Università Cattolica del S. Cuore, via E. Parmense, 84, 29100 Piacenza  
³Parco Tecnologico Padano, Via Einstein, Polo Universitario, Lodi 26900, ITALY

**Abstract:**
Automated annotation of the bovine genome sequence identified more than 25K genes. In the present study the Ensembl Biomart tool was used to identify genes that are orthologous between cattle and the human, mouse and dog genomes. Comparing the genes annotated in cattle with other species 865 were absent, whereas among the genes annotated in cattle 6,247 had no orthologs in the human, mouse and dog genome.

There are several reasons why genes annotated in cattle are missing in the other genomes; these include missing or incorrectly assembled bovine sequence and structural variations in the genes between species. However, there large number of genes in cattle for which orthologs are missing in the other species was not expected. An in silico pipeline was developed to make pair-wise alignments of the “missing” genes between genomes and the evidence from sequence alignments was used to cluster the genes into different classes. The possible systematic failure of the automated protocols to annotate particular classes of genes was investigated using annotation information available in Ensembl, ontology descriptions and manual curation for each subset of genes. This work will help the annotation of the bovine genome and will assist in the development of improved gene models for automated annotation protocols.

**Poster 4023**

**Title:** High-resolution comparative mapping between X chromosomes in sheep, cattle, and human

Presenting Author: Tom Goldammer, Forschungsinstitut für die Biologie landwirtschaftlicher Nutztiere (FBN), Wilhelm-Stahl-Allee 2, 18196 Dummerstorf, Germany

Other authors (name only):
1. Ronald M. Brunner
2. Alexander Rebl
3. Chun H. Wu
4. Koh Nomura
5. Tracy Hadfield
6. Clare Gill
7. Brian Dalrymple
8. James E. Womack
9. Noelle Cockett

**Abstract:**
X chromosome morphology and banding structure differ very strong between sheep and cattle, contrary to most other chromosomes in both species. We focused on the extensive analysis of the X chromosome in sheep (Ovis aries, OARX) to study the differences of chromosome organization in sheep and cattle and constructed a high-resolution physical map including 169 loci based on a comparative mapping approach. DNA sequences for mapping were selected by using the Virtual Sheep Genome Browser v1.1. Performed genotyping of DNA sequences in the USUoRH5000 ovine whole-genome radiation hybrid panel followed by RH map construction using the Carthagene1.1R software package revealed RH loci order on OARX. The RH map consists of four linkage groups with an average loci density of 1.3 megabases. Assignment of numerous microsatellite markers to the RH map of OARX anchors available linkage maps physically to the chromosome. The cytogenetic map based on FISH of BAC clones of the CHORI-243 library. FISH loci represent cytogenetic anchors and support loci order within the constructed RH map. Except for little rearrangements, the map is concordant with the virtual OARX map. A developed comparative map for sheep, cattle, and human X chromosomes identifies several new evolutionary break points and chromosome rearrangements.

**Poster 4024**

**Title:** Mapping and sequencing of major histocompatibility complex class I loci in rabbit
Order of the authors: Hassina OULD-HAMOUDA, Angélique TEILLAUD, Mark STAM, Claire ROGEL-GAILLARD

Presenting Author: Claire ROGEL-GAILLARD

1 INRA UMR314, Laboratoire de Radiobiologie et Etude du Génome, Jouy-en-Josas, 78350, France.
2 CEA/DSV/iRCM/SREIT/LREG, Jouy-en-Josas, 78350, France

Other authors (name only):
1. Hassina OULD-HAMOUDA
2. Angélique TEILLAUD
3. Mark STAM

Abstract:
The Major Histocompatibility Complex (MHC) is a conserved locus in jawed vertebrates and thus constitutes an interesting region for phylogenomic analyses and, more generally, for addressing the question of vertebrate evolution. Rabbit MHC referred to as RLA (Rabbit Leukocyte Antigen) complex, localized on OCU12q11 is still poorly characterized. Our aim was to produce data on the RLA complex in order to include rabbit in MHC comparative analyses among Mammals. We have built a physical map of the RLA complex and produced two contigs for class I and III adjacent regions. Contig1 spans 700 kilobases from MOG to VARSL and contig2 spans 1500 kilobases from CDSN to NOTCH4. We have identified two clusters of class I genes. A first cluster is intermingled between TRIM26 and TRIM39 in contig1 and a second cluster is situated between POU5F1 and BAT1 anchoring genes in contig2. Unlike the human MHC, no third class I gene cluster was detected close to ZNRD1. Six BACs covering the class I gene clusters were sequenced. Our results demonstrate the presence of six class I loci in each cluster and identification of putative functional loci is in progress. Phylogenetic studies of class I loci will be discussed.

Poster 4025

Title: Goat mammary gland expressed genes representative of immunity function during pregnancy before the acquisition of its secretory phenotype

Presenting Author: FAUCON Félicie Unité GPL Bâtiment 221 INRA Jouy en Josas Domaine de Vilvert 78352 Jouy en Josas

Other authors (name only):
1. REBOURS E.
2. HELBLING J.C.
3. BEVILACQUA C.

4. AUBERT J.
5. MARTIN-MAGNIETTE M.L.
6. ROBIN S.
7. MARTIN P.

Abstract:
Mammary tissue differentiation into a secretory epithelium that synthesizes milk occurs during pregnancy. Impaired differentiation during pregnancy may provoke lactating default. Molecular mechanisms associated to mammary tissue terminal differentiation remained incompletely understood especially in ruminants. Morphological differences between mice and ruminant mammary tissues before pregnancy let’s think that there are differences between these two mammals in the differentiation mechanisms. A transcriptional analysis on 5 physiological stages (4 during pregnancy and 1 during lactation) appeared as the best approach to get overview of the differentiation process. An appropriate experimental design was drawn to follow gene expression profiles during differentiation of mammary tissue. Using 3 nulliparous or primiparous goats per stage (a small ruminant model economically acceptable), the comparison was done on 9 dye-swaps, using 22k bovine oligoarrays. Statistical analysis revealed that 2 330 genes varied significantly at least once. These genes were divided into 19 clusters. Identification of biological functions revealed that mechanisms described for mice can be applied to goats. As for example the fall down in lactation of genes implicated in cell growth and proliferation and the increase in lactation of expression of genes implicated in lipid metabolism. A special observation, not described for mice, was the expression of immunity genes at mid-pregnancy.

Poster 4026

Title: In silico characterization of the genome of bovine endogenous retrovirus γ4 based on the bovine genome sequencing draft.

Authors:
Presenting Author
Name: Chankyu Park
Address: Laboratory of Mammalian Genomics, Department of Animal Biotechnology, Konkuk University, Seoul 143-701, South Korea.

Conference registration number of Presenting Author: 0192

Other authors (name only)
1. Rui Xiao
2. Kwangha Park
3. Younshin Oh
Abstract
The genome of the replication-competent bovine endogenous retrovirus (BERV) γ4 provirus, which is the most abundant ERV family in the bovine genome, was characterized in detail. The BERV γ4 genome revealed that BERV γ4 harbors 8576 nucleotides and has a typical 5′-long terminal repeat (LTR) gag provirus from pro pol env LTR. The retroviral organization with a long leader region positioned before the gag open reading frame. Multiple sequence analysis showed that the nucleotide difference between 5′ and 3′ LTRs was 4.2% (mean value 0.042) on average, suggesting that the provirus formed at most 13.3 million years ago. Gag was separated by a stop codon from pro pol in the same reading frame, while env resided in another reading frame lacking a functional surface domain. According to the current bovine genome sequence assembly, the full-length BERV γ4 provirus sequences are only found in chromosomes 1, 2, 6, 10, 15, 26, 28, X, and unassigned, although the partial sequences are almost evenly distributed throughout the entire bovine genome. This is the first detailed study describing the genome structure of BERV γ4. Combined with our recent reports on the characterization of bovine ERVs, this study will contribute to illuminating ERVs in cattle in which no information was previously available.

Poster 4028
Title: Molecular characterization of the putative regulatory regions of the caprine prion protein gene (PRNP)


Other authors (name only):
1. Fabio Zuccon.
2. Maria Grazia Maniaci.
3. Maria Caramelli
4. Pier Luigi Acutis

Abstract:
Transmissible spongiform encephalopathies (TSEs) are a group of inevitably fatal neurodegenerative diseases of humans and animals. Several nonsynonymous single-nucleotide polymorphisms (SNPs) within the PRNP coding sequence have been associated with resistance/susceptibility to TSEs in humans, sheep, goats, deer and elk. Recent studies indicate that PRNP regulatory polymorphisms may also influence TSE occurrence. However, no basic information currently exists regarding the sequence composition and/or the frequency of genetic polymorphisms within the putative regulatory regions of the goat PRNP gene. Therefore, we generated sequences for the caprine PRNP gene regions that are homologous to known mammalian PRNP regulatory regions. Promoter prediction analysis using CpGProD software yielded a single goat PRNP promoter that was homologous to regions of known promoter activity in sheep and cow. Sequence analysis and haplotype cloning revealed 25 nucleotide polymorphisms and 3 insertion/deletion (indel) polymorphisms in the putative promoter and intron I. No variation was detected within the predicted exon 1. The four conserved PRNP promoter motifs previously described for mammalian species were also conserved within goat sequence. The results of this study provide the initial genomic foundation for future investigations on the range of transcription factor binding motif profiles in the goat PRNP gene promoter.

Poster 4027
Title: Polymorphism and evolution of the major histocompatibility complex DRA and DQA genes in Equidae

Presenting Author: Dr Petr Horin, Institute of Animal Genetics, University of Veterinary and Pharmaceutical Sciences, Brno, Czech Republic

Other authors (name only):
1. Eva Janova
2. Jan Matisovic
3. Jiri Vahala
4. Roman Vodicka

Abstract:
As a result of host and pathogen interactions, the major histocompatibility complex (MHC) genes coding for antigen binding and presenting molecules are the most polymorphic genes in the vertebrate genome. The diversity of MHC class II genes DQA and DRA was studied in the family Equidae. In addition to 11 previously reported DRA and 26 DQA alleles, 6 new DRA alleles and 11 new DQA alleles were identified in the genus Equus, species Equus kiang, E. hemionus, E. zebra, E. grevyi and E. burchelli. Phylogenetic analysis of both DQA and DRA sequences of the family Equidae provided further data supporting the trans-species hypothesis. The phylogeny trees constructed did not match the standard Equidae taxonomy, expressing specific features of the MHC genes. Analysis of selection showed differences between the less variable DRA and more variable DQA genes. DRA alleles were more often shared by more species. The DQA sequences analyzed showed strong positive selection, the selected amino acid positions mostly corresponded to selected positions in rodent and human DQA genes.
**Poster 4029**

**Title:** Polymorphism of PrP gene in Polish pig breeds.

Presenting Author: Agata Piestrzyńska-Kajtoch, National Research Institute of Animal Production, Department of Immuno- and Cytogenetics, 32-083 Balice, Poland

Other authors (name only):
1. Barbara Rejduch

**Abstract:**
Host-encoded prion protein is believed to be a component of an infectious agent of transmissible spongiform encephalopathies (TSE). All analyzed mammalian genomes have a copy of PrP gene. Many polymorphisms within the coding region have an influence on the TSE susceptibility, pathogenesis and transmissibility. There are no reported TSE cases in pigs until today, but it was experimentally proved that pigs can become infected. The aim of our study was to investigate the polymorphism in the coding part of the PrP gene in Polish pig breeds.

DNA was isolated from different tissues of 152 pigs [Sus scrofa domestica] from 3 breeds (56 unrelated Polish Large White [WBP], 55 unrelated Polish Landrace [PBZ] and 41 Pulawska – some of which were half-siblings). The coding region of the gene was amplified, sequenced and analyzed in freeware genetic computer program.

Three silent polymorphisms were found (counted from the beginning of the coding region): T198C, C354T and C723G. 3.6% of WBP were homozygotes T198C, 30.3% of WBP and 9.1% of PBZ were heterozygotes T198C/T, 21.9% of Pulawska and 1 WBP pig were heterozygotes C354T/C. Only 1 WBP pig was heterozygote C723G/C.

These polymorphisms have not been published before.

---

**Poster 4030**

**Title:** Comparative annotation of Sus scrofa chromosome 1

Presenting Author: Zhihua Jiang, Department of Animal Sciences, Washington State University, Pullman, WA 99164-6351 USA

Other authors (name only):
1. Natasha M. Robinson
2. Jennifer J. Michal

**Abstract:**
The pig whole genome is currently being sequenced by The Wellcome Trust Sanger Institute through funding provided by Cooperative State Research, Education and Extension Service at the United States Department of Agriculture (CSREES-USDA). This project uses a clone-by-clone sequencing strategy, based on the MTP of BAC clones. To date, more than 1,200 clones have been selected and used to generate initial shotgun sequencing data for Sus scrofa chromosome 1 (SSC1). We have assigned these clones to their orthologous regions on human chromosomes 6, 9, 14, 15 and 18, which are consistent with the current comparative relationship between these two species. On the other hand, SSC1 might have some small regions orthologous to human chromosomes 1, 10, 11 and 20, respectively. Interestingly, the human orthologous regions harbor a total of 1,412 genes, including 917 known function genes, 170 genes similar to the other genes, 137 hypothetical genes, 114 ORF genes and 74 pseudogenes. However, one-to-one orthologous hits only confirmed 820 (89%) known function genes, 47 (28%) genes similar to other genes, 66 (48%) hypothetical genes, 82 (72%) ORF genes, and 27 (36%) pseudogenes in pigs. These results indicate that different categories of genes might have different evolutionary mechanisms.
**Poster 5001**

**Title:** Evaluation of feed efficiency in beef cattle under low and high energy feedlot diets

Durunna, Obioha N., Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, Canada, T6G 2P5.

Other authors (name only):
1. Mujibi, Fidalis D.N
2. Wang, Zhiquan
3. Moore, Stephen. S

**Abstract:**
Residual feed intake (RFI) measures the difference between an animal’s actual and expected feed intakes (EFI) based on the maintenance of body weight and its growth. Little is known about the effects of different diets on animals’ RFI measures. Our objective was to evaluate the effect of diet energy density on animal RFI ranking. One hundred and seventy six randomly chosen steers, 5-7.5 mo of age were used for the feeding trials at the Kinsella ranch, University of Alberta. The steers were crosses between Angus, Charolais or Alberta hybrid bulls and hybrid dams. The diets were offered sequentially (ad-libitum) with low energy diet in Period 1 (P1) then high energy diet in Period 2 (P2). Individual feed intakes were collected using the Growsafe® feeding system. Body weights were measured biweekly. Average daily gain (ADG) and EFI were estimated using a linear regression analysis for each period. RFI for each animal was calculated separately in each period. Means of RFI were 0.00 ± 0.03 and 0.00 ± 0.06 respectively for P1 and P2. Pearson and Spearman-rank correlations between P1 and P2 RFI indices were 43% and 41% respectively. One hundred and four steers (59.1%) changed in their RFI rankings. This re-ranking of RFI may be an indication of genotype by environment interaction.

**Poster 5002**

**Title:** Population Studies for Arabian Horses by 15 Microsatellite loci

Presenting Author: Dorota Monies, King Faisal Specialist Hospital and Research Centre, P.O.Box.3354, Riyadh 11211, MBC-03, Saudi Arabia

Other authors (name only):
1. Abu Al Saud N.
2. Sahar N.
3. Meyer BF.

**Abstract:**
Prior to the introduction of any DNA marker as a tool for human paternity test and personal identification, a population genetic database should be established. A similar database should also be made for establishing equine paternity. There are several commercial available kits for equine DNA genotyping but population genetic studies are limited. This paper describes allele frequencies together with some statistical parameters of parentage interest for 15 STRs (VHL20, HTG4, AHT4, HMS7, HTG6, AHT5, HMS6, ASB23, ASB2, HTG10, HTG7, HMS3, HMS2, ASB17, HMS1) included in the Equine StockMarks and the Equine Panel1.1 systems in a sample of 283 unrelated Arabian horses. Genetic parameters were calculated using Cervus (v.3.0) and PowerStats (v.1.2) software. No significant deviations from the Hardy – Weinberg Equilibrium (HWE) were detected, with the exception of the HMS3 locus. The polymorphic information content (PIC) varied from 0.35 to 0.75 and the power of discrimination (PD) from 59.5% to 91.8%. The combined power of exclusion (CPE) for 8 out of 9 loci (without HMS3) recommended by the International Society for Animal Genetics (ISAG) was 98.34% and the CPE for 14 markers was 99.84%. Analyzed parameters demonstrate that the abovementioned STR markers are suitable for paternity investigation in the Arabian horse population, but with some limits to estimating a paternity as certainty.

**Poster 5003**

**Title:** Developmental validation of a single-tube amplification of the 17 loci for parentage of horse with StockMarks for Horses

Presenting Author: Zaulet Mihaela, 91-95 Spl Independentei street, county 5, Bucharest.

Other authors (name only):
1. S. Georgescu
2. S. Kevorkian
3. A. Manea
4. R.A. Vatasescu-Balcan
5. M. Rebedea

**Abstract:**
The PCR technology provides a sensitive method for parentage verification and individual identification of horses. For the 17 microsatellites designated (VHL20, HTG4, AHT4, HMS7, HTG6, AHT5, HMS6, ASB23, ASB2, HTG10, HTG7, HMS3, HMS2, ABS17, LEX3, HMS1, CA425) , amplification of the STR loci was realized by multiplex PCR using StockMarks for Horse Paternity PCR Typing Kit (AppliedBiosystems). Characterized by defined critical control points, the assay was conducted within a quality assurance system compliant with ISO/IEC 17025:2005 guidelines.
Taking into account that we are employing an “in house”- developed method the necessary validation...
process was carried out with horse DNA samples extracted from blood. The key parameters of method validation include specificity, LOD (limit of detection), robustness, repeatability, reproducibility, selectivity and the reproducibility of DNA extraction. In conclusion microsatellites can give sufficient and reliable information for paternity testing and in horse breeding this technology has the potential to be of great use in monitoring levels of genetic variation.

Poster 5004

Title: Developmental Validation of a Sensitive and Discriminating Panel of Novel Canine STR Markers for Forensic Testing

Presenting Author: Elizabeth Wictum, Forensic Unit, Veterinary Genetics Laboratory, University of California, Davis, California, USA

Other authors (name only):
1. Teri J. Kun
2. Nguyen V. Truong
3. Christina D. Lindquist
4. Julia Collins
5. Aaron Wong

Abstract:
The molecular analysis of forensic evidence has revolutionized the criminal justice system. Its strength has been used to both incriminate the guilty and free the innocent. While analysis of human biological material has been extensively vetted in courtrooms worldwide, forensic analysis of animal DNA is still gaining acceptance. As the oldest domesticated species, dogs (*Canis lupus familiaris*) inhabit 39% of households in America and, after humans, are the species of greatest forensic interest. Numerous genetic markers have been employed to individualize canids for parentage verification, breed identification, phylogeny, and diversity assessment; however, forensic analysis requires the application of more stringent marker selection criteria. To address the lack of a standardized canid forensic panel that meets those criteria, tools were developed to mine the 7x dog genome sequence data. Fifteen unlinked highly discriminating tetranucleotide-repeat markers were identified and assembled with a sex-determination marker into a multiplex capable of generating a full profile with less than 0.1 ng of nuclear DNA. This panel has the potential to be not only a valuable tool for the emerging field of veterinary forensic science but also demonstrates utility for parentage verification in highly inbred dog populations and for phylogenetic analyses of unique canid populations.

Poster 5005

Title: Structure and breed assignment of river buffalo populations using microsatellite markers (preliminary results)

Presenting Author: DANIELA IAMARTINO, LGS – Laboratorio Genetica e Servizi, Via Bergamo 292, 26100 Cremona Italy.

Other authors (name only):
1. CHIARA LOFFI
2. ZEFFIRINA RONCHETTI
3. MARIELA CASTRILLO
4. ANGELO COLETTA
5. MICHELE BLASI

Abstract:
Given the increasing importance of food traceability for safety and quality and with aim to safeguard of the made in Italy products, we have investigated on possible genetic differences among river buffalo populations coming from different geographical areas: Italy, North-Africa, East Europe, China and Brazil. Fourteen microsatellite loci, amplified with a protocol of one multiplex-PCR, were used to assess genetic diversity within and across the populations. We analysed a total of 318 animals: 60 from Italy, 60 from North Africa, 58 from Romania, 67 from Bulgaria, 8 from China and two groups from Brazil, respectively of 32 and 33 animals. The mean number of alleles was 5.7 and the expected heterozigosity ranged from 0.526 (Jaffarabadi, Brazil) to 0.720 (Murrah, Bulgaria). Although there was abundant genetic variation, genetic differentiation between populations was low (Fst=0.075±0.01). Individuals were clustered applying a parametric genetic mixture analysis implemented in the Structure 2.2 software. The number of genetic clusters K was tested using the admixture model, a burning period of 100,000 followed by 100,000 MCMC repeats. Even if the demarcation within East Europe groups is lower than the others, the method resolved 7 clusters, being able to correctly identify buffalo populations.

Poster 5006

Title: Admixture analysis of European and Chinese pig breeds

Presenting Author: Christoph Knorr, Institute of Veterinary Medicine, University of Göttingen, Burckhardtweg 2, 37077 Göttingen, Germany

Other authors (name only):
1. Helge Täubert
2. Daniel G. Bradley
3. Kefei Chen
4. Lusheng Huang
5. Bertram Brenig

**Abstract:**
We present population genetic analysis of microsatellite variation based on 17 microsatellite markers in 15 different pig breeds from Europe (Angeln Saddleback, Bunte Bentheimer, German Edelschwein, German Landrace, Göttingen Minipig, Hampshire, Pietrain, European Wild boar and Swabian-Haellian swine) and Asia (Jiangquhai, Luchuan, Minzhu, Rongchang, Yujijiang, and Tibetan). The European group included the native European wild boar and the breed Göttingen minipig; in total 336 pigs were investigated. Chinese breeds showed a higher average mean number of alleles per locus (4.7) than the European population (3.9). Unique alleles were present in 6 breeds. To analyse admixture between these populations following approaches were used: the systematic selection of population associated alleles for European and Asian breeds and examination of their frequency; examination of genetic distances between breeds; and a model-based Bayesian admixture analysis. The results show an influence of Asian breeds in European populations, more in commercial dam lines than in sire lines. Within the European breeds, the Göttingen minipig, a crossbred of Asian and European breeds, showed strongest influence of Asian admixture. Tibetan pigs were the most admixed and allelic diverse Chinese sample. For two Asian breeds (Tibetan and Minzhu), geographical outliers in the sample, signature of European admixture was found.

**Poster 5007**

**Title:** Polymorphisms in DRB3 binding pockets affect immune response to a FMDV peptide in cattle

Presenting Author: Rebecca Baxter, The Roslin Institute and R(D)SVS, University of Edinburgh

Other authors (name only):
1. Zen Lu
2. Elizabeth J Glass

**Abstract:**
Immune responsiveness is a complex trait involving many genes, but Major Histocompatibility Complex (MHC) genes are consistently implicated as having a significant impact. However their relative importance in out-bred species such as cattle is less apparent. In this study we have investigated the effect of bovine MHC class II (BoLA) - DRB3 polymorphism on the response to a Foot-and-Mouth disease virus (FMDV) peptide in a genotypically diverse Charolais-Holstein F2 cross population. The majority of functionally relevant polymorphisms alter the conformation of ‘pockets’ within the peptide binding cleft. The pockets act as anchors for pathogen peptides and thus these polymorphisms determine the binding affinity of peptides to MHC molecules. We are evaluating how the polymorphisms in the pockets affect the binding of the FMDV peptide and the consequent response of both B and T cells. We found that the specific polymorphisms in pocket 4 significantly affect anti FMDV IgG1 and IgG2 responses. To enable us to fully comprehend the interactions between the differences in the DRB3 pockets within the DR molecule and the FMDV peptide we are currently using a modelling based approach.

**Poster 5008**

**Title:** Structural equation modeling to find the relationship between different genetic loci and different phenotypes influencing body composition

Presenting Author: Gudrun A. Brockmann, Humboldt-Universität zu Berlin, Institut für Nutztierrassenwissenschaften, Invalidenstraße 42, 10115 Berlin, Germany

Other authors (name only):
1. Shiu-Wern Tsaih
2. Christina Neuschl
3. Gary A Churchill
4. Renhua Li

**Abstract:**
We used the method of structural equation modeling to find the relationship between different genetic loci and different phenotypes influencing body composition in mice. We use an intercross population between the mouse lines NMRI8 (selected for high body weight) and DBA/2 (control) in which QTLs for body weight, adiposity, and muscle weights were mapped. The traits were measured at 6 weeks, when animals became fertile and have finished the fastest growth phase. Using the technique of structural model analysis, we considered body weight as lean mass and fat mass and were able to distinguish genetic loci that affect adiposity from those that affect lean mass. The results show that the NMRI8 alleles of the selection line on chromosomes 7 and 14 have pleiotropic positive effects on both muscle and fat tissue mass, while a locus on chromosome 13 contributed to the selection response only by increased fat deposition. The fat mass was also affected by a complex pattern of interaction between loci on chromosomes 6 and 14. The sex of individuals affected the fat mass either directly or indirectly via the muscle as a mediator. The analysis sheds new light on the action of genes controlling body weight as composite trait of fat and muscle tissues.
Poster 5009

Title: Plasmid-Mediated Muscle-Targeted Gene Nutrition: The Porcine GHRH myogenic expression plasmid for enhancing body weight in rat

Presenting Author: Qingyong Meng, College of Biological Science, China Agricultural University, No. 2 Yuanmingyuan west Road, Beijing 100094, China

Other authors (name only):
1. Yali Zhang
2. Xia Gong
3. Zhengquan Yu
4. Qifa Xie
5. Tonghui Lin
6. Jing Fei
7. Ning Li

Abstract:

Plasmid-mediated myogenic expression was used to enhancing animal growth as gene nutrition. It was a promising, simple, and inexpensive approach in this field. In this report, a myogenic expression plasmid, ppG-H6, containing the porcine GHRH cDNA under the control of a alpha-actin promoter was constructed. Based on the expression of this plasmid transfer in C2C12 cells, ppG-H6 showed the ability of driving GHRH expression in muscle cell. A single 300ug dose of this plasmid was injected intramuscularly into rat. Follow-up evaluation demonstrated increased body weight and GH level by the plasmid. After injection with ppG-H6, these rats showed a more than seven-fold increase in serum GH and the increased body weight as 234.99±5.73g vs 203.19±13.11g of control group ppG-CK. These results suggest plasmid-mediated GHRH ectopic and myogenic expression are methods for increasing the body weight of an animal.

Poster 5010

Title: EVALUATION OF MARKERS FOR SPECIES IDENTIFICATION IN VETERINARY FORENSIC GENETICS.

Presenting Author: Riina Maria Vittoria – CEA, Istituto Zooprofilattico Sperimentale del Piemonte, Liguria e Valle d’Aosta – Turin, Italy.

Other authors (name only):
1. Domenis Lorenzo
2. Colussi Silvia
3. Trisorio Silvia
4. Robetto Serena
5. Caramelli Maria
6. Orusa Riccardo

Abstract:

Identifying the species of origin of biological traces is important in veterinary forensic caseworks. A morphological approach is not always sufficient or feasible and the analysis of genetic markers can be useful.

We analyzed mitochondrial genetic markers to evaluate their suitability for species identification and discrimination. We also describe their practical application in a case report. Samples from domestic and wild animal species were analyzed sequencing cytochrome b (cytb) and cytochrome oxidase I (COI) genes. Obtained sequences were compared with those deposited in public databases.

For the case report we also carried out an alignment for a direct comparison between the sample and the sequences of the most suspected involved species. The genetic distance was calculated and a phylogenetic tree was constructed. The bootstrap test confidence was used to determine the statistical significance of the species assignment.

Obtained results showed that cytb is an effective marker for species identification and is a promising tool for forensic genetic purposes. COI showed a lower discriminatory power regarding philogenetically close species.

The developed analysis protocol appeared a good approach for veterinary forensic investigations, where, different than in human forensics, numerous animal species need to be discriminated and no standardized methods are available.

Poster 5011

Title: Patterns of genetic diversity near genes under selection in cattle

Presenting Author:   Pam Wiener, The Roslin Institute, University of Edinburgh, Roslin, EH25 9PS United Kingdom

Other authors (name only):
1. Beatriz Gutiérrez-Gil

Abstract:

Because of their well-documented histories, domestic species provide an important resource for understanding the patterns of variation left by selection. Such information is useful for interpreting patterns in other species as well as for identifying as yet undiscovered genes in domestic species. This study characterizes diversity patterns across genes under selection in cattle, with associated phenotypes related to both domestic use (muscle conformation) and physical appearance (coat color). The analysis includes
two marker types (microsatellites and SNPs) and applies various statistical approaches, from which comparisons of statistical power and inferences can be made. While results of these analyses demonstrate that selection can be detected using comparisons of breeds bred for different phenotypes, some limitations in the methodologies and markers are also highlighted. This work has important implications for assessing the methodologies available for detecting past selection and for understanding the process of domestication.

Poster 5012

Title: Microsatellite mutation rates in the eastern tiger salamander (Ambystoma tigrinum tigrinum)

Presenting Author: Zafer Bulut, Selcuk University Faculty of Veterinary Medicine Department of Biochemistry Campus/Konya/TURKEY


Abstract:
Microsatellites are commonly used for mapping and population genetics because of their high heterozygosities and allelic variability (i.e., polymorphism). Microsatellite markers are generally more polymorphic than other types of molecular markers (e.g., allozymes or SNPs) because the insertions/deletions that give rise to microsatellite variability are relatively common compared to nucleotide substitutions. Nevertheless, empirical evidence on microsatellite mutation rates (MMRs) is lacking in most vertebrate groups despite the importance of such estimates to key population genetic parameters such as the effective population size. Here, we present empirical data on MMRs in eastern tiger salamanders (Ambystoma tigrinum tigrinum). We conducted dozens of captive breeding trials, then genotyped over 1000 offspring (i.e., >2000 meioses) at a suite of microsatellite loci in an effort to determine a) genetic parentage and b) MMRs. Our results provide the first estimates of MMRs in urodèles (newts and salamanders), and they illustrate that MMRs can vary by more than an order of magnitude across loci within a given species.

Poster 5013

Title: Data validation and algorithmic design for chromosome grouping of SNPs from a 7K genome-wide porcine chip

Presenting Author: Vivi Raundahl Gregersen, Dept. of Genetics and Biotechnology, University of Aarhus, Blichers allé 20, P.O. box 50, DK-8830 Tjele, Denmark


Abstract:
High-throughput marker systems increases the data output drastically, hence the need for rapid validation methods. An Illumina Infinium® SNP BeadChip was produced for mapping and QTL purposes. Initially, a data set of about 1,000 animals divided in ten sire families was used together with 100 replicate samples to generate a configuration template for the clustering of the SNPs. The dataset was divided into unsuccessful, not segregating and segregating SNPs and deviation from Hardy-Weinberg Equilibrium was calculated. Secondly, additional 350 animal samples of different breeds were included in the sample set and processed. About 10,000 animals were genotyped by both the original and new configuration templates and discrepancies were further investigated.

The resulting set of 4,366 segregating SNPs was grouped into the 18 porcine autosome groups using two-point linkage calculated by the Crimap 4.2 package. Prior knowledge of location for 400 SNPs was used as a backbone. Ten SNP sets overlapping each other by 500 SNPs was used for the initial group placement (LODS>50). New linkage sets was produced of the remaining SNPs combined with each of the autosome groups. This procedure was performed until the LODS could be relaxed to seven. A total of 4,181 SNPs could be uniquely divided into the 18 groups. The remaining 185 SNPs were either singletons or linked together in small groups.

Poster 5014

Title: The Mitochondrial DNA (mtDNA) Haplogroup Compositions of Three Native Turkish Sheep Breeds and Their Implications on the Conservation Studies

Presenting Author: Eren Yüncü, Middle East Technical University Department of Biological Sciences Lab:147

Other authors (name only): 1. Evren Koban 2. İnci Togan

Abstract:
Sheep domestication is believed to have occurred at least by three separate events in different domestication centres. Archaeological studies suggest
that Turkey might be harbouring the earliest one. Since, there were no wild sheep in Europe before domestication, European domestic sheep might be mainly extends of this centre. However, native Turkish sheep might still be harbouring variability which is absent in extend breeds. Their high genetic variability can be a signature for maintained diversity since domestication. Therefore, they must be conserved. Furthermore, Turkey is close to other assumed domestication centres. Hence Turkish breeds might also be admixture of products of these domestication centres. The evolutionary history of breeds may help to resolve different causes of high diversity in Turkish breeds. The aim of the study is to determine composition of sheep mtDNA haplogroups indicating three domestication events, among native breeds, as a part of a national project TURKHAYGEN-I. Three native Turkish breeds (Karayaka, Akkaraman, Gökçeada) are studied for mtDNA control and ND4 region by RFLP and SSCP methods, respectively. Results are employed to enlarge existing data to reveal evolutionary history of the Turkish sheep breeds. Results will also be used to prioritize Turkish domestic sheep breeds in conservation studies.

**Poster 5015**

**Title: The Bos taurus Genome Database: an integrated database for Bovine Genome research**

Presenting Author: Dajeong Lim, National Institute of Animal Science, RDA, 564 Omockchun-dong Gwoneon-gu


**Abstract:** We constructed a database to study function genomics in the Bovine genome. The BGD integrated into UniGene, markers, QTLs, transcripts and SNP information and then provides a useful tool for studying genomics and biological mechanisms in the bovine. We experimentally determined 63,625 ESTs (Expressed Sequence Tags) from 3 full-length enriched cDNA libraries of Korean Native Cattle (KNC). QTL and marker were retrieved from the public databases. The EST data were clustered and assembled into unique sequences, contigs, and singletons. The BGD database provides functional annotation, identification of SNPs, genome mapping of coding sequences, and characterization of trait loci controlling. It also provides KNC specific information compare other breeds of cattle. Especially, we are developing network model that is coexpressed genes for QTL region. This is accessible online and can answer queries using several search options, including clone IDs, marker, QTL and genes. The BGD supported advanced search interface, disease browser and comparative gene map of pig. Graphical map view and genome browser show ESTs and contigs from National Institute of Animal Science (NIAS). BGD is accessible at http://bgd.nabc.go.kr.

**Poster 5016**

**Title: Genetic variability in Bolivian Llamas (Lama glama) using microsatellites: preliminary results**

Authors: Julia Barreta¹, Volga Iñiguez¹, Roberto Chiri², Fernando Romero¹, Tito Rodríguez², Beatriz Gutiérrez-Gil³ and Juan-Jose Arranz⁴

¹ Instituto de Biología Molecular y Biotecnología, UMSA, La Paz (Bolivia). ² Banco de Germoplasma de Camélidos UTO, Oruro, (Bolivia). ³ Instituto de Investigaciones Agropecuarias, UMSA, La Paz (Bolivia). ⁴ Dpto. Producción Animal Universidad de León, (Spain).

**Abstract:** Bolivia has the highest census of llamas in the world with approximately 2.400.000 animals. This species present an important degradation process; it is caused principally by the human press on the land. As a consequence, the study of this genetic resource is critical for the establishment of management and conservation plans that assures genetic conservation of llamas. In a preliminary study in coordination with the conservation plans of BANCAMEL, genetic variability at 10 microsatellite markers was examined in Bolivian Llamas. A total of 108 animals belonging to following ecotypes: Azanaque, Cruce Culta , Kara, Ketena, Morejon, Orinoca, Rivera, Sajama, Totora, Tiutiri and Yawaroko were analyzed. Microsatellites were amplified in multiplex and alleles were identified using an ABI3130 sequencer and GeneMapper software. The number of alleles per loci, heterozygosity, and departures from Hardy Weinberg equilibrium were calculated for all microsatellite loci in the whole populations. All the populations show, in general, a good fit to HWE. The observed number of alleles per loci in the 108 animals varied between 6 for YWLL44 marker and 24 in YWLL29. The observed average heterozygosity in the populations ranges from 0.656 in Morejon ecotype to 0.759 in Titutiri llamas.
Poster 5017

Title: QTL detection for milk protein composition of bovine milk

Presenting Author: G.C.B Schopen, Animal Breeding and Genomics Centre, Wageningen University, Wageningen, P.O. Box 338, 6700AH, The Netherlands

Other authors (name only): 1. M. H. P.W. Visker 2. J.A.M. van Arendonk 3. H. Bovenhuis

Abstract:
The objective of this study was to map QTLs for milk protein composition in Dutch Holstein-Friesians. From 1912 first parity cows, one morning milk sample was collected and analyzed for the major milk proteins using capillary zone electrophoresis. The 1912 phenotypes were corrected for systematic environmental effects. The heritabilities, i.e. the proportion of phenotypic variation due to genetics, were moderate to high for the major milk proteins. DNA was isolated from blood samples from 849 cows and semen samples from their 7 sires. A whole genome scan using 1536 single nucleotide polymorphisms was performed. The QTL analysis consists of a multimarker regression approach for interval mapping in half-sib families. In this study, we covered 2831 cM of the cattle genome. Several genome-wise significant (P<0.05) QTLs were found for the major milk proteins distributed over eight chromosomal regions. Fine mapping of QTL regions to reduce confidence intervals of the detected QTLs, to facilitate new candidate genes that affect milk protein composition, is in progress.

Poster 5018

Title: Phylogenetic and population structure of Thai indigenous pigs and Thai wild boars assessed by mitochondrial DNA sequence polymorphism

Presenting Author: Rangsun Charoensook, Department of Animal Science, Agriculture Faculty, Chiang Mai University, 50200, Chiang Mai, Thailand Institute of Veterinary Medicine, Georg-August-University, Burckhardtweg 2, 37077 Göttingen, Germany


Abstract:
The study of mitochondrial DNA (mtDNA) D-loop sequences were carried out using 72 Thai indigenous pigs and 11 Thai wild boars collected at twelve locations in five provinces in Northern Thailand. To infer phylogenetic relationships between Thai pigs, other Asian pigs, and European pigs, published D-loop sequences from GenBank were used for comparison with our data. Moreover, molecular diversity and population demographic statistics were calculated to determine the genetic structure of Thai pig populations. The genetic distances between groups further indicate that Thai pigs are closely related with other Asian pigs (mean distance ± SD; 0.0074 ± 0.0016), but are very different from European pigs (0.0205 ± 0.0039). In total, 24 different mtDNA haplotypes of Thai pigs have been described. Finally, 19 haplotypes specific for Thai indigenous pigs, and four haplotypes specific for Thai wild boars belonged to the Asian group and only one haplotype - although specific for Thai indigenous pigs - fits into the European group. This study proposes firstly that the indigenous Thai pigs are closely related with Thai wild boars and may eventually go back to the common Asian ancestor. Secondly, Thai and Asian pigs are distinctly different from European pigs.

Poster 5019

Title: Genetic diversity of Churra Tensina breed sheep, and relationship with other Churra group breeds using microsatellites.

Presenting Author: Jorge H. Calvo & Unidad de Tecnología en Producción Animal, CITA, Avda de Montañana 930, 50059 Zaragoza. Spain


Abstract:
Churra Tensina sheep is an endangered local course-wooled hardy breed belonging to Churra group which is raised for lamb production, although it has been milked for years. Thirty microsatellites proposed by FAO and ISAG for biodiversity studies have been used to characterize Churra Tensina, Churra, Churra Lebrijana and Latxa sheep breeds. All microsatellite loci analysed were found to be polymorphic, showing an observed heterozygosity ranged from 0.6599 to 0.7084 for Churra Lebrijana and Latxa sheep breeds. All microsatellite loci analysed were found to be polymorphic, showing an observed heterozygosity ranging from 0.6599 to 0.7084 for Churra Lebrijana and Latxa, respectively. All populations showed significant deviations from Hardy-Weinberg proportions. The fixation indices showed statistically significant differentiation between breeds (Fst=0.119), low inbreeding related to the whole population (Fit=0.146) and within population
Results from Geneclass assignment test revealed that 100% animals were assigned to the population they were collected from. Structure programme inferred 4 clusters for the best estimated of the log probability of the Churra Tensina population, indicating that Churra Tensina was structured in 4 sub-populations. Reynolds distance results showed the higher and lower distance between Churra Tensina and Churra Lebrijana, and Churra Tensina and Latxa pairs, respectively. In conclusion, Churra Tensina sheep breed showed good diversity values despite its low effective population size.

**Poster 5020**

**Title:** Conservation genetics of the endangered “Altamurana” sheep breed from Southern Italy

Presenting Author: Elena Ciani, General and Environmental Physiology Department, University of Bari, Via Amendola 165/a 70126 Bari, Italy

Other authors (name only): 1. Francesca Cecchi 2. Elisa Mazzanti 3. Elisabetta Castellana 4. Roberta Ciampolini

**Abstract:**
Altamurana is an originally triple-purpose sheep breed from Southern Italy that has experienced a drastic population size contraction. A total of 89 animals, sampled from two different farms in the province of Bari (Apulia), were genotyped at 19 microsatellite loci belonging to the ISAG-FAO panel. Samples were chosen according to genealogical records in order to minimize relatedness among animals. A high level of linkage disequilibrium was observed in the total sample, also among non syntenic locus pairs, suggesting the presence of population substructuring. A genetic partitioning at the farm level was supported by the FST value between the two farms (0.056; P < 0.0001) and by two different population assignment tests. In particular, using a Bayesian approach, different ancestralities within a single farm were highlighted, corresponding to different farm origins. Within-group linkage disequilibrium greatly decreased when clustering the animals based on the different ancestralities. These results confirm the power of STR loci to detect fine-scale genetic structure and highlight a noticeable differentiation among the ancestralities, probably arisen from reproductive isolation and/or different selection strategies across farms. This should be seriously taken into consideration, due to critical implications for the breed conservation.

**Poster 5021**

**Title:** Dynamic annotation of DNA sequences in agricultural species.

Presenting Author: Alessio Valentini Department of Animal Production, Università della Tuscia, Viterbo, Italy


**Abstract:**
Several millions of DNA sequences exist in public databases deriving from ESTs projects and various other contributions. Moreover, parallel sequencing nowadays permits to achieve a throughput of hundreds of thousands of cDNA sequences in a single experiment. Annotation of these sequences is required to perform several tasks, like microarray generation, species comparison for expressed genomes, metabolic pathway assessment in orthologous species etc. We developed a public domain portal where large databases of sequences can be automatically annotated starting from those of model species related to the one of interest. The portal has been developed for agricultural species and particularly for animals and plants. Users can request annotation of sequences available in specific NCBI databases or supply a set of his/her own set of data. The search is performed sequentially on a list of databases ranked according to their suitability to retrieve the best information for the species of interest. Here we present some examples concerning cattle and sheep.

**Poster 5022**

**Title:** GENETIC AND PHENOTYPIC RELATIONSHIPS BETWEEN BLOOD GAS PARAMETERS AND ASCITES-RELATED TRAITS IN BROILERS

Presenting Author: Ane Marie Closter, Wageningen University, Animal Breeding and Genomics Centre, Marijkweg 40, 6709 PG Wageningen, The Netherlands

Other authors (name only): 1. Pieter van As
Abstract:
Ascites is a metabolic disorder in broilers associated with rapid growth and high demand for O₂. The tendency of broilers to develop ascites is heritable. Successful selection against ascites susceptibility requires good and easy-to-measure indicator traits. In juvenile chicken, blood gas parameters have been suggested as indicator traits. The aim of the present study was to estimate heritability and genetic correlations between blood gas parameters (measured during week 4), body weight and heart ratio (post-mortem indicator for ascites).

For this purpose total blood gas traits were available on nearly 3000 broilers from generation 7 and 8 of an advanced intercross line from 2 dam lines. The birds were challenged under cold conditions. Heritability for heart ratio was 0.43, for pH 0.15, and 0.19 for HCO₃⁻. Heritabilities for \( p_{vCO_2} \) (0.02), \( p_{vO_2} \) (0.03) and \( sO_2 \) (0.07) were low. Genetic correlation between heart ratio and HCO₃⁻ was 0.31 (±0.15), -0.62 (±0.21) between heart ratio and \( p_{vO_2} \), and -0.04 (± 0.45) between heart ratio and \( p_{vCO_2} \). Heritability for blood gas parameters and the genetic correlations with heart ratio estimated in the current data do not support the suggestion that at that age blood gas parameters are useful traits in selection against ascites susceptibility.

Supported by Hendrix Genetics B.V. (The Netherlands) and by grant 401-138-8000 from STW (The Netherlands).

Poster 5023
Title: Centromeric/pericentromeric junction within the MHC locus on chromosome 7 in pig

Mark STAM
1,2

1 INRA, DGA, UMR314, Laboratoire de Radiobiologie et Etude du Génome, Jouy-en Josas, France.
2 CEA/DSV/iRCM/SREIT/LREG, Jouy-en-Josas, 78350, France

Other authors (name only):
1. Hélène HAYES
2. Maude BERTAUD
3. Angélique TEILLAUD
4. Gaetan LEMONNIER
5. Claire ROGEL-GAILLARD

Abstract:
In swine, the centromere of chromosome 7 (SSC7) splits the Major Histocompatibility Complex (MHC) into two regions, the class III (p arm) and class II (q arm) regions. Although swine MHC is completely annotated, its centromeric junction remains uncharacterized. Screening our pig BAC library with primers specific of the centromeric genetic marker SW472 identified three BACs containing 3.6Kb-long tandem repeats. Fluorescence in situ hybridization showed that the 3.6Kb repeat and BACs are specific of the SSC7 centromere. Screening with the 3.6Kb probe isolated additional BACs localized at the SSC7 pericentromeric/centromeric junction, eight of which were sequenced with a threefold coverage. Pig VEGA annotation and sequence comparison with BAC ends from the swine genome physical map revealed that one BAC mapped to a locus 21Kb from the BTNL4 gene on SSC7 q arm. Some BACs contained SINE elements characteristic of euchromatin and one BAC end was similar to the mc2 centromeric sequence, previously described as specific of metacentric pig chromosomes. Therefore, we have identified BACs spanning the pericentromeric/centromeric junction within MHC on the q arm of SSC7. These results represent the first step in the sequence analysis and further expression studies of the junction between euchromatin and heterochromatin regions in pig.
program PHASE v2.1.1 in absence of familiar data. The analyzed microsatellites are highly polymorphic in this breed compared with other breeds. In consequence, the haplotype variability is remarkably high (73 haplotypes), requiring specially designed statistical tests able to deal with this variability. Moreover, sample structuration by flock was included in the analysis. The statistical tests were performed by SASv9.0 and Rv2.7.0. Several haplotypes were found to be significantly associated with susceptibility/resistance to both OPA and MV, although the results are sensitive to the inclusion of flock-effect in the analysis. It can be concluded that the parameters used in this work can improve the haplotype-based association studies in domestic animal species.

Poster 5025

Title: In vitro Conservation and Preliminary Molecular Characterization of Some Turkish Native Domestic Animal Genetic Resources-I (TURKHAYGEN-I)

Kurar E1, Arat S2, Ertugrul O3, Pabuccuoglu S4, Sagirkaya H3, Tasdemir U6 and Togan I7

1Faculty of Veterinary Medicine, Selcuk University, Konya, Turkey
2TUBITAK Institute for Genetic Engineering and Biotechnology, Gebze, Turkey
3Faculty of Veterinary Medicine, Ankara University, Ankara, Turkey
4Faculty of Veterinary Medicine, Istanbul University, Istanbul, Turkey
5Faculty of Veterinary Medicine, Uludag University, Bursa, Turkey
6Lalahan Livestock Central Research Institute, Ministry of Agriculture and Rural Affairs Ankara, Turkey
7Department of Biological Sciences, Middle East Technical University, Ankara, Turkey

Abstract:
Dramatic chances of environmental conditions including global warming and environmental pollution have irreversibly affected the world’s flora and fauna. As in other countries, several endemic domestic animals have been totally lost or under danger in Turkey. This matter has been discussed on various platforms in Turkey and was declared as a priority and various action plans have been prepared. In order to solve this problem, the Turkish Ministry of Agriculture and Rural Affairs (TKB) has taken precautions by application of conservation programs for some animal genetic resources. By the contribution of the TKB, Turkish National Science and Research Foundation (TUBITAK) and 10 universities, a consortium was founded to take other efficient measures. The main objective of TURKHAYGEN-I is the application of in vitro conservation programs by foundation of two cryo-banks in which duplicated samples will be preserved. The project covers endemic sheep (13), cattle (6), goat (5), horse (5) breeds and Anatolian water buffalo. A total of 1500 animals have being collected from different parts of Turkey and the herds are placed in research farms of the participating institutions where somatic cells, embryos, DNA, and semen samples are collected. All preserved animals are also characterized at DNA level using microsatellite and mtDNA markers.

Poster 5026

Title: Animal QTLdb: A tool set to warehouse and compare cattle, pigs, chicken and sheep QTL within and between species.

Presenting Author: Zhi-Liang Hu, Department of Animal Science, Iowa State University, 2255 Kildee Hall, Ames, IA 50011

Other authors (name only):
1. James M. Reecy
2. Eric Ryan Fritz,
3. Svetlana Dracheva,
4. Behnam Abasht,
5. Stacey N. Meyers,
6. Denis M. Larkin,
7. John Bastiaansen,
8. Carol Scott,
9. Xiao-Ping Li,
10. Wonhee Jang,
11. Kwan-Suk Kim,
12. Donna Maglott,
13. Sean Humphray,
14. Jonathan E. Beever,
15. Susan J. Lamont,
16. Harris A. Lewin
17. Jane Rogers,
18. Jill Maddox,
19. Max F. Rothschild,

Abstract:
The Animal QTL database (QTLdb; URL: http://www.animalgenome.org/QTLdb/) was developed to aid with identifying causal mutations responsible for economically important traits by allowing comparative viewing of all published porcine QTL data. The capability of AnimalQTLdb has since been expanded by: (1) modifying the database so that QTL from multiple species can be warehoused and comparisons made within and across species; and (2) extending QTL map alignments to a greater range of structural genomic features, such as radiation hybrid markers, microsatellites, finger printed contig BAC clones, SNPs, and microarray elements. AnimalQTLdb now contains QTL data for cattle,
swine, chicken and sheep; serves as both a data warehouse and search tool; and a powerful QTL oriented, map based data mining tool through easy-to-use and user-friendly web interfaces. Animal Trait Ontology (ATO) was introduced to improve consistency of trait nomenclature and organise traits for database management. Current initiatives include extending the utility of AnimalQTLdb by integrating mouse, rat and human QTL information and by working with livestock, rat and mouse communities to extend the ATO to facilitate cross-species trait comparisons. As new types of data (e.g. copy number variation and metabolomics) become available these will be incorporated into AnimalQTLdb to further enhance data mining.

**Poster 5027**

**Title:** Molecular estimation of effective population size – a new perspective for the assessment of risk status for livestock breeds?

Presenting Author: M. Tapio, International Livestock Research Institute ILRI, PO Box 30709, Nairobi 00100, Kenya

Other authors (name only):
1. C. Flury
2. F. Ytournel
3. H. Simianer
4. O. Hanotte
5. S. Rieder

**Abstract** (maximum 200 words):
For livestock breeds of the developing world demographic information is often lacking and effective population size cannot be estimated accurately. With dense marker information the application of LD-based methods for the estimation of effective population sizes might offer new perspectives to overcome this problem. In an international project genome-wide SNP data from two African cattle breeds (Sheko and N’Dama) and one Swiss breed (Eringer) will be evaluated. So far, information for 17 Sheko individuals and 22 N’Dama individuals, respectively genotyped within the bovine HapMap-project were available. Thereof three chromosomes with frequent indications of selection and three “neutral” chromosomes were selected. Effective population size was estimated based on r2. Levels of LD and development of estimated effective population size over generations were compared between chromosomes. A decreasing population trend was found for both breeds, however with large standard deviations. The practical results were compared with simulated results. The goal of the overall study is the evaluation of marker based estimation of effective population size for the management of livestock populations in developing countries.

**Poster 5028**

**Title:** ANEXdb: An Intuitive Animal Microarray Expression and Functional Annotation Database

Presenting Author: Oliver Couture, Iowa State University, Ames, IA, USA

Other authors (name only):
1. Keith Callenberg
2. Neeraj Kaul
3. Remy Younes
4. Jack Dekkers
5. Vasant Honavar
6. Christopher Tuggle

**Abstract**:
The Animal Expression Database (www.ANEXdb.org) allows users to upload microarray experiments with Affymetrix® GeneChip®-based expression data using a user-friendly web interface. The database can accommodate any species or any chip based on the Affymetrix® platform. Data can be stored as private information within the database, and the database streamlines public submission to GEO using an automatic SOFT formatter. In addition to storing expression data, the database is connected to an annotation database, annotDB, which currently contains an in-house alignment of 1,691,062 porcine sequences assembled into 98,452 consensus sequences and 192,736 singletons. Through BLAST, these sequences will not only be connected to the Affymetrix® porcine GeneChip®, but to other species using NCBI’s RefSeq database. This BLAST analysis allows additional information from NCBI Gene, KEGG, Pfam, GO, and Ensemble to be tied to the sequences for further annotation. The design of annotDB is species-independent, and can accommodate any type of sequence. The annotation and expression databases are modular in design, and can be independently updated. EBI’s Bio-Mart will be used as an interface for querying both expression and annotation simultaneously. Currently data for over 400 Porcine GeneChip® hybridizations are being uploaded to ANEXdb.

**Poster 5029**

**Title:** Single nucleotide polymorphisms of the KITLG and KIT genes in pigs

Presenting Author: Naohiko OKUMURA, Second Research Division, STAFF-Institute, 446-1 Ippaiizuka, Kamiyokoba, Tsukuba, Ibaraki 305-0854, Japan.
Other authors (name only):
1. Toshimi MATSUMOTO
2. Noriyuki HAMASIMA
3. Takashi AWATA

Abstract:
Here we report the variability in genes encoding KIT tyrosine kinase receptor and its ligand KITLG by determining single nucleotide polymorphisms (SNPs) in 384 individuals constituted by 16 pig groups of 11 breeds, two synthetic-line cross pigs, two cross breeds, and Japanese wild boar. SNPs and indels within the coding sequence regions of KITLG and KIT and their 5′-flanking regions were detected by aligning sequences from 8 pigs; subsequently, the SNPs were genotyped using MALDI TOF-MS (SEQUENOM MassARRAY system). Principal component analysis using allele frequencies in the SNP loci showed a distant relationship between Asian and Euro-American pig groups, except for Berkshire and Tokyo X breeds. These breeds were located within the mid-portion of the distribution in the first principal component. The Hampshire breed was distant from the other pig groups on the axis of the second principal component. Haplotype estimation revealed each unique haplotype of KIT gene being in the white and Hampshire breeds because of the causative gene for their coloration. On the other hand, the haplotypes of KITLG and KIT detected in the Berkshire breed were prevalent in Asian pig groups; this tendency is different from that observed in other Euro-American pig breeds.

Poster 5030

Title: Genetic analyses of Ghanaian chicken ecotypes: diversity and relationships with other breeds

Presenting Author: Boniface Baboreka KAYANG, Wildlife Research Center of Kyoto University, c/o JASSO, Tanaka-Sekiden-cho, Sakyo, Kyoto 606-8203, JAPAN

Other authors (name only):
1. E. Inoue
2. R. Osei-Amponsah
3. A. Naazie
4. K. Kinoshita
5. M. Mizutani
6. K. Nirasawa
7. A. Nakamura
8. K. Nagao
9. H. Hayakawa
10. K. Fujiwara
11. S. Ito
12. M. Inoue-Murayama

Abstract:
This study investigated the genetic diversity of local chicken populations in three different agro-ecological zones of Ghana and examined their extent of genetic differentiation and phylogenetic relationships with other breeds. A total of 494 birds randomly sampled from one Red Jungle Fowl and nine domestic fowl populations, comprising the Ghanaian ecotypes (Interior Savannah, Forest and Coastal Savannah), Japanese native breeds (Nagoya, Mikawa and Gifujidori), White Leghorn, Rhode Island Red and Fayoumi, were genotyped across 21 microsatellite loci from the recommended FAO panel. Out of a total of 178 alleles observed, 27 were unique to the Ghanaian populations. The average number (± SD) of alleles/locus and the observed heterozygosity were, respectively, higher in the Ghanaian populations (7.76 ± 3.39 and 0.591 ± 0.110) compared to the rest (6.24 ± 2.79 and 0.374 ± 0.100). Genetic differentiation (F_{ST}), heterozygote deficiency (F_{IS}), Nei’s genetic distance and cluster analysis (STRUCTURE) indicated that the three Ghanaian ecotypes were distinct from the other populations but were closely related to one another and showed high diversity but lacked sub-structuring. The results suggest that the Ghanaian populations are derived from a mixture of egg- and meat-type breeds and comparison of phenotype and genotype will provide information for selection.

Poster 5031

Title: Probabilistic Expert Systems (PES) as a tool for parentage testing in horses with lack of parental genotypes.

Presenting Author: Paolo De Iuliis, UNIRELAB s.r.l., Laboratorio di Genetica Forense Veterinaria, Via Campobello 7/c, 00040 - Pomezia (RM) Italy

Other authors:
1. Margherita Bonuglia
2. Chiara Bocci
3. Cinzia Grasso
4. Marina Dobosz

Abstract:
Accurate determination of relatedness is of great importance in horse breeding industry. Currently the majority of horse breed registries adopted DNA technology in order to verify pedigree records and to solve queries of parentage. When genetic data of one of the individuals in comparison are missing, performance of a parentage test could be questionable. We present a new approach to analyze complex cases of identification inference. In a case of disputed maternity involving a thoroughbred foal, whose mare was dead, the relationship was initially excluded by genetic incompatibility between the foal and a bone that was supposed belonging to the mare. This result led us to bring forward two different hypothesis:
1. the bone belongs to the mare and relationship is excluded;
2. the bone doesn’t belong to the mare of the foal.

Due to the impossibility to verify the identity of the bone, the relationship was then verified by statistical analysis of other genotypes in the family of the putative mare through Probabilistic Expert Systems. One of these systems (FINEX) was recently developed for forensic purposes and is particularly valuable for disputed paternity/maternity cases in humans. We will illustrate how this case has been successfully worked out by FINEX.

Poster 5032

Title: Genetic diversity of the mtDNA in five Italian horse populations.

Presenting Author: Chiara Bocci, UNIRELAB s.r.l., Laboratorio di Genetica Forense Veterinaria, Via Campobello 7/c, 00040 - Pomezia (RM) Italy

Other authors:
1. Marina Dobosz
2. Cristina Martinez-Labarga
3. Margherita Bonuglia
4. Cinzia Grasso
5. Paolo De Iuliis

Abstract:
Genetic variability of a 510bp fragment in the D-loop region of the mtDNA in five horse breeds bred in Italy was analysed. Two breeds are autochthonous: Maremmano (N=40) and Murgese (N=23). Other Italian populations were also included: Arabian (N=40), Thoroughbred (N=40) and Sicilian Oriental Thoroughbred (N=7). Murgese samples were selected at random, while selection for other breeds has been performed using pedigree information in order to maximise maternal diversity. Genomic DNA was extracted from blood on FTA® papers. PCR primers were designed according to Xu and Arnason (1994). Sequences obtained were aligned and compared with reference sequence from GenBank X79547. For a total of 150 individuals, 44 polymorphic sites representing 51 haplotypes were observed. Genetic distances were estimated by Slatkin’s linearized Fst’s (Schneider, Roessli, and Excoffier 2000) method using Arlequin package software 2.1 (Schneider et al. 2000). Our data were compared with 22 mtDNA sequences present in literature representing Italian horse breeds. An Equus asinus sequence was used as the outgroup. The aim of this study is to evaluate matrilineal relationships of these breeds in order to enlarge genetic information about horse populations bred in Italy and to confirm their multiple origin reported by other investigators.

Poster 5033

Title: Genome-wide analysis of pig alternative splicing with full-length enriched cDNA libraries.

Yong-Min Cho, Dajeong Lim, Kyung-Tai Lee, Mi-Jeong Byun, Yeonkyung Kang, Heebal Kim, Seok-ki Im

NIAS (National Institute of Animal Science, RDA), Seoul National Univ.

Abstract:
We produced 75,110 expressed sequence tags (ESTs) from a full-length enriched 6 cDNA libraries: porcine abdominal fat, porcine fat cell, porcine loin muscle, backfat, liver and pituitary gland. Alternative splicing is an important role of the functional complexity in mammalian genes, developmental stage and pathological state. Particularly, EST/mRNA alignment is important method for in silico detection of AS (Alternative Splicing) because ESTs and mRNAs are the most abundant resources and are regarded as a direct evidence of the expressed genes. Defining alternative splicing isoforms with EST fragment data faces the multiassembly problem. The multiassembly problem is the problem for reconstructing the most likely set of full-length isoform sequences from a mixture of EST fragment data. Our algorithm identifies novel alternative splicing events based on sequence similarity searches between each EST dataset and known gene set without multiassembly problem. We determined alternative splicing events using BLAST and validated AS events with SIM4 program. As a result, we found 717 sequences of case1 type (EST insertion) and 255 sequences of case2 type (EST deletion).

Poster 5034

Title: Evidence for cattle major histocompatibility complex (BoLA) class II DQA1 gene heterozygote advantage against clinical mastitis caused by Streptococcus and Escherichia species..

Presenting Author: Shin-nosuke Takeshima

Other authors (name only):
1. Yuki Matsumoto
2. Shou Chen
3. Tatsuyuki Yoshida
4. Harutaka Mukoyama
5. Yoko Aida

Abstract:
Mastitis, which is a mammary gland inflammatory response to bacterial infection, is a major problem in the dairy industry. We genotyped bovine major histocompatibility complex (BoLA)-DRB3 and BoLA-DQA1 in 120 cattle with clinical mastitis and
85 randomly selected Holstein cattle in Japan by polymerase chain reaction-sequence based typing (PCR-SBT). The mastitis cattle were divided into four groups according to the bacterial species causing the mastitis (Staphylococcus aureus, Streptococcus, Escherichia and coagulase-negative staphylococci). The BoLA-DRB3 and BoLA-DQA1 heterozygosity of each group was compared to that of the control cattle. The expected heterozygosities based on Hardy-Weinberg proportions and the observed heterozygosities for each locus were compared for each group. The Escherichia and Streptococcus mastitis groups showed significant differences between their expected and observed heterozygosities with regard to their BoLA-DQA1 genes. No differences were observed for any group with regard to the BoLA-DRB3 genes. We then found that two BoLA-DQA1 alleles promoted susceptibility to mastitis caused by Streptococcus and Escherichia, respectively. This is the first report in Japan showing that heterozygosity of the BoLA-DQA1 gene is associated with resistance to mastitis progression.

Poster 5035

Title: The “Osservatorio Italiano HCM Felina”: a Feline Inherited Diseases Network in Italy

Presenting Author: M. Longeri, Università degli Studi di Milano, Dipartimento di Scienze Animali, Sez. Zootecnica Veterinaria, Via Celoria, 10 - 20133 Milano (Italy)

Other authors (name only):
1. G. Pertica
2. P. Ferrari
3. P.G. Brambilla
4. C. Cozzi
5. M. Strillacci
6. B. Gandolfi
7. P. Valiati
8. C. Scilini
9. M. Polli
10. L. Guidobono Cavalchini

Abstract:
In Italy cat breeders are scattered in many associations and breed fan clubs. The availability of new genetic tools for inherited disease identification and the possibility of shipping buccal swabs directly from the cattery to diagnostic Labs, prevent a real control of veterinaries, geneticists and breed associations on diagnosis and selection of the diseases. In 2008 to overcome the absence of a network the Italian Observatory on Feline Hypertrophic Cardiomyopathy (HCM) has been constituted. So far it associates the Italian Main Coon and other breed clubs, the Milan University (Spin-off Vetogene) and selected clinicians. Main targets: 1) scientific monitoring on the cat inherited disease in Italy, mainly HCM, so far; 2) supporting to breeder in the selective choices; 3) constitution of a biological and data bank for further scientific studies; 4) improving link among cat breeders, selected veterinarians and scientific community, for the first time in Italy. A cat joining the Observatory is submitted to microchip implant, periodical clinical controls, DNA tests and blood storage. Data are anonymously available to Observatory associated.

Breed statistics, correlation between clinic and genetic results and some discussion on possible spin-off on feline genetic research are also proposed.

Poster 5036

Title: The trend of the inbreeding in the course of time in “Bracco Italiano” dog breed.

FRANCESCA CECCHI1, ROBERTA CIAMPOLINI1, BRUNO DE SANCTIS2, FABIO CASETTI3, SILVANO PRESCIUTTINI1

1Centro Interuniversitario di Ricerca e di Consulenza sulla Genetica del Cane, Università di Pisa, Viale delle Piagge 2, 56124 Pisa, Italy; 2USL Viterbo; 3Società Amatori Bracco Italiano (SABI), Via Dosso 7, 26856 Mirabello di Senna Lodigiana, Italy.

Abstract:
We refer on a genealogical study on the “Bracco Italiano” dog breed, with the aim to study the investigate the trend of the inbreeding during a period of 38 years. The complete electronic record of the breed, including 20,499 animals born between 1970 and 2007, was downloaded from the ENCI database. Animals were divided in eight periods according to their year of birth. The following genetic parameters were calculated, using the program CFC: a) the level of inbreeding (F) of the total population; b) the individual inbreeding; c) the number and the percentage of inbreds and the average inbreeding coefficient in the inbreds animals for five-year periods.

The inbreeding coefficient was <0.05 in 8,103 dogs (39.53%) whereas it was >0.20 in 487 dogs (2.51%). The average inbreeding coefficient over all animals resulted 3.6% (from 0.4% in 1970-1975 to 5.5% in the last two years). The percentage of inbreds per year increased from 2.44% in 1976-1980 to 100% in the past two years. It is concluded that a regular monitoring of genetic variability of the population must be adopted, in order to avoid the danger of an
excessive increase of inbreeding in the future, which would result in significant inbreeding depression.

Poster 5037

Title: The prion protein gene (Prnp) polymorphisms in Slovenian sheep breeds

Presenting Author: Jelka Zabavnik Piano, Veterinary Faculty, University of Ljubljana, Gerbičeva 60, Ljubljana, Slovenia

Other authors (name only):
1. Marko Cotman
2. Ivan Ambrožič
3. Drago Kompan
4. Polona Juntes
5. Milan Pogačnik

Abstract:
Scrapie is the transmissible spongiform encephalopathy (TSE) that occurs naturally in sheep. Three scrapie-linked polymorphisms in the prion protein gene (Prnp) located at the codons 136, 154 and 171 modulate the susceptibility to classical type of scrapie. In order to evaluate the Prnp polymorphisms in Slovenian sheep breeds and to evaluate the genetic susceptibility through the years sheep of four breeds (Bela Krajina Pramenka, Bovska sheep, Istrian Pramenka, Jezersko-Solcava) Prnp polymorphisms were analyzed from the year 2005. Polymorphisms at codons 136, 154 and 171 were determined by nucleotide sequencing of the Prnp and by allelic discrimination assay. Each year different sheep were randomly sampled for Prnp genotyping. The most frequent genotype in all Slovenian sheep breeds is ARQ/ARQ. Animals carrying this genotype are moderately susceptible to scrapie. The allelic variant VRQ, known to carry very high risk of scrapie is only poorly represented in the population of the examined Slovenian sheep breeds and is decreasing through the years. Slightly more abundant is the allelic variant ARR that is typical for sheep resistant to scrapie. This allelic variant is increasing through the years.

Poster 5038

Title: Viral-vector Mediated Transgenesis in the Göttingen Minipig Brain

Presenting Author: Claus Hedegaard, Dept. of Genetics and Biotechnology, Faculty of Agricultural Sciences, University of Aarhus, Blichers Allé, 8830 Tjele, Denmark.

Other authors (name only):
1. Andreas Nørgaard Glud
2. Mette Slot Nielsen
3. Carsten Reidies Bjarkam
4. Jens Christian Sørensen
5. Poul Henning Jensen
6. Christian Bendixen
7. Knud Larsen

Abstract:
Transgenic large mammals for modelling human diseases are receiving increasing interest as potential elaboration of rodent models. Pigs offer splendid anatomic, physiologic, and genetic recapitulation of humans, avoiding ethical concerns with primates used for experiments. Particularly the architecture of the pig CNS provides an attractive basis for modelling neurological diseases. The considerable longevity expected for transgenic pigs to develop age-related neurodegenerative disorders has made us attempt to establish a proof of concept with surgery based, virus mediated gene transfer in Göttingen minipigs. Three anaesthetised minipigs were MRI-scanned for determination of coordinates for substantia nigra (SN). The animals then received six stereotaxic, unilateral deep brain injections with recombinant lentivirus harbouring the EGFP gene. After four weeks, the animals were euthanised, brains dissected, and relevant tissues recovered. Histology showed injection points targeting SN as visualised by Nissl staining while subsequent immunohistochemical staining with anti-EGFP demonstrated transgene expression and transport to striatum. PCR analysis confirmed presence of the EGFP gene in SN but not in striatum, reflecting the presence of the neuronal cell nuclei in SN, and axons extending to striatum. The obtained results indicate the present technique to be an efficient means of transducing discrete, selective tissues of the Göttingen minipig brain.

Poster 5039

Title: Profile of the Circulating DNA in Healthy Cattle

Presenting Author: Julia Beck, Chronix Biomedical GmbH, Goetheallee 8, 37073 Goettingen, Germany

Other authors (name only):
1. Julia Beck
2. Howard B. Urnovitz
3. Wilhelm Wemheuer
4. Bertram Brenig
5. Ekkehard Schütz

Abstract:
Circulating nucleic acids (CNAs) have been shown to have diagnostic utility in human diseases. The aim of this study was to sequence and organize DNA obtained from serum of apparently healthy cattle providing an extensive catalog of the circulating DNA profile.
Serum DNA from 79 cattle was extracted, amplified, sequenced using high-throughput pyrosequencing (454/Roche) and categorized by: 1) origin (bovine versus exogenous), 2) functionality (repeats, coding or non-coding sequences), and 3) chromosomal localization. 7.81E+05 sequences were obtained consisting of 1.27E+08 nucleotides, of which 78% was attributed to known database sequences. Of the known sequences 95% were genomic and 5% were of bacterial or viral origin. Of the genomic sequences, 48% originated from repeats and 34% from annotated genes, which is within the expected range. Interestingly, coding sequences (CDSs) were significantly underrepresented in the circulating DNA pool. Whereas, sequences matching to repetitive elements were found in amounts as expected from the genome. The under-representation of CDSs argues strongly against a solely apoptotic/necrotic origin of CNAs. The normal bovine circulating DNA profile described here and the technique of high-throughput sequencing together with a detailed analysis of the origin of the circulating DNA may open a diagnostic horizon.

Poster 5040

Title: Genetic diversity among goat populations in Turkey

Genetic Diversity Among Goat Populations in Turkey

Bulut Z¹, Kurar E², Ozsensoy Y², Altunok V¹, Nizamlioglu M¹
¹Department of Biochemistry, Faculty of Veterinary Medicine, Selcuk University, Kampus, Konya, TURKEY
²Department of Genetics, Faculty of Veterinary Medicine, Selcuk University, Kampus, Konya, TURKEY

Abstract

Microsatellites are one of the favored markers used in population genetics studies because of their high heterozygosity and versatility. In this study, 11 microsatellites loci were used as molecular markers to determine the intra and inter genetic diversities of eight different goat breeds including Kil, Angora, Kilis, Yayladag, Shami, Honamli, Saanen, and Alpine. The genetic differentiation between breeds was considerable as a result of the statistically significant (P<0.001) pair wise FST values of each pair of breeds. Exceptionally, FST values calculated for Honamli and Kil breeds were statistically non-significant (P>0.05). Heterozygosity values were ranged between 0.62 and 0.73. According to the Structure and assignment test, individuals of Angora and Yayladag goat were assigned to the breed they belong to, while other breeds were assigned to two more different groups. Because, this study for the first time presented genetic data for the Yayladag goat, structure analysis and assigned test results were interesting for the Yayladag goat. Additional and different molecular markers are needed for future analyses.

Poster 5041

Title: Metanalysis of microsatellite data allows a molecular classification of European cattle

Presenting Author: J.A. Lenstra, Faculty of Veterinary Medicine, Utrecht University, Yalelaan 2, 3584 CM, Utrecht, Netherlands

Other authors (name only):
1. European Cattle Genetic Diversity Consortium

Abstract:

Several European research groups have studied the molecular diversity of cattle by microsatellite genotyping. A combined dataset now contains genotypes of >10,000 animals from 186 European, African and Asian taurindicine breeds, 98 of which have been analyzed with all 30 FAO-recommended microsatellites. Combination of data from different sources and different marker panels was validated by metaanalysis of 39 breeds sampled and genotyped by different laboratories. Model-based clustering and distance analysis resulted in a molecular classification of European cattle that correlates better with geographical origin than with breeding objectives:

(1) Northern-European breed clusters: Lowland Pied, Baltic-Highland Red, North-West Intermediate, British, Nordic, Nordic-Ayrshire and Russian cattle;
(2) Central-European clusters: Southern French, Central Brown and Western-Central breeds, the last cluster containing Central Spotted, Western-Alpine and German Yellow subclusters;
(3) Mediterranean clusters: Balkan, Podolian and Iberian breeds, the last cluster containing Cantabrian, Andalusian, Morena and Lidia as distinct subgroups.

SNP analysis of 1079 Y-chromosomes revealed two haplotypes Y1 and Y2. Y1 is predominant in most Northern-European breeds but is also carried by several Northern-Spanish bulls. The geographical distribution of breed clusters and paternal lineages is proposed to reflect successive migrations as well as relatively recent expansions of successful breeds.

Poster 5042

Title: Presence of Southeast Asian and Indian subcontinent maternal lineages in a genetically unstructured Zimbabwean chicken population

Presenting Author: Weigend, Steffen; Institute of Farm Animal Genetics, Friedrich-Loeffler-Institute, 31535 Neustadt, Germany
Other authors (name only):
1. Muchadeyi, Farai
2. Eding, Herwin
3. Wollny, Clemens
4. Simianer, Henner
5. Groeneveld, Eildert

Abstract:
Genetic diversity and population structure of village chickens of five agro-ecological zones in Zimbabwe was investigated using twenty-nine autosomal microsatellites and 455bp of the mitochondrial DNA D-Loop region. A comparison to a set of reference populations from Malawi, Sudan, Northwest Europe and six purebred lines was made. The maternal origin of the Zimbabwean chickens was investigated by comparing mtDNA haplotypes to those of the presumed centers of domestication. Both microsatellites and mtDNA analyses did not support that Zimbabwean chicken eco-types are substructured according to agro-ecological zones. Based on microsatellites, the five chicken eco-types of Zimbabwe clustered together. Marker estimated kinship values indicated that the level of genetic variation was high and very similar within and between eco-types. Zimbabwean chickens were distinguishable from the reference populations, particularly purebred lines. Analysis of mtDNA indicated that two distinct maternal lineages exist in the five Zimbabwean eco-types. For one of these lineages, chickens from Zimbabwe and Malawi shared major haplotypes with populations that have a Southeast Asian background. The second maternal lineage, probably from the Indian subcontinent, was common to the five Zimbabwean eco-types, Sudanese and European chickens as well as purebred broiler and layer lines.

Poster 5043
Title: DNA testing of urine identifies horses that have undergone post-race drug testing.

Presenting Author: Dr C.K. Harper – Veterinary Genetics Laboratory Faculty of Veterinary Science University of Pretoria Onderstepoort 0110

Other authors (name only):
1. Mrs. A. Ludwig
2. Dr. S. de Kok

Abstract:
This investigation followed a question regarding the identity of a specific horse that tested positive to an illegal substance on a post-race urine sample. The urine sample from the horse was submitted to the Veterinary Genetics Laboratory at Onderstepoort (VGL) for DNA testing by the National Horseracing Authority (NHA) of South Africa Drug Testing Laboratory. The resulting genotype was compared to the genotypes of South African Thoroughbred horses. It was matched, without exclusion, to 12 STR loci and Amelogenin. This included the 9 loci recommended by ISAG. The NHA requested that a standard sampling and transport methodology be set up as guideline to facilitate the DNA testing of urine samples in future in cases in which the test results are challenged. The NHA provided a total of 13 samples from various racecourses in the country. The samples were analyzed using the Gentra Puregene Cell and Tissue Kit (Qiagen) with the addition of Hydrochloric acid as an initial step to remove calcium carbonate present in horse urine. PCR was performed using 12 STR loci in twoplexes and Amelogenin. Results were analyzed and recommendations prepared for future sample collection, handling and analysis.

Poster 5044
Title: Molecular characterization of epistatic interactions in a M16i X Myostatin mouse cross


Department of Animal Science; Center for Integrated Animal Genomics, Iowa State University, Ames, IA 50011-3160 USA

Abstract:
Animals that lack functional Myostatin are known for their extreme muscle growth and lack of adipose accumulation. To identify QTL that have differential effects in the presence and absence of functional Myostatin, M16i mice, which was derived from a line which was selected for weight gain before six weeks of age, were reciprocally crossed with Myostatin-null mice to generate F1s, which were mated to generate F2s. A total of 1072 F2 generation progeny that were homozygous Myostatin wild-type or Myostatin-null were genotyped for 186 SNP distributed across the genome. Initial statistical analysis of body weight indicated that the main effects of Myostatin genotype and reciprocal cross were significant across all ages. The main effect of sex was significant at 1, 4, 5, 6 weeks of age. Single nucleotides that were statistically associated with body weight (P<10^-10) were clustered on several chromosomes. Significant (P<0.05) two-way interactions were detected between Myostatin and cross, sex and Myostatin, Myostatin and SNP. Significant Myostatin by SNP interactions may indicate epistatic interactions. Segregation distortion was observed on chromosome 1 and was independent of Myostatin genotype and reciprocal cross were significant across all ages. The main effect of sex was significant at 1, 4, 5, 6 weeks of age. Single nucleotides that were statistically associated with body weight (P<10^-10) were clustered on several chromosomes. Significant (P<0.05) two-way interactions were detected between Myostatin and cross, sex and Myostatin, Myostatin and SNP. Significant Myostatin by SNP interactions may indicate epistatic interactions. Segregation distortion was observed on chromosome 1 and was independent of Myostatin genotype. In contrast, the segregation distortion observed on chromosome X was only present in homozygous Myostatin wild-type mice. These preliminary results indicate that this population will be a valuable resource to better understand
epistatic interactions. Funded by a grant from USDA-NRI 2006-35205-16696.

**Poster 5045**

**Title:** Bayesian analysis of 15,110 single nucleotide polymorphisms in the bovine genome

Presenting Author: Jennifer J. Michal, Department of Animal Sciences, Washington State University, Pullman, WA 99164-6351 USA

Other authors (name only):
1. Zhihua Jiang
2. Xiao-Lin Wu
3. Ming Zhang
4. Raymond W. Wright, Jr.

**Abstract:**
In order to further understand molecular mechanisms of mutation and genome evolution, single nucleotide polymorphisms (SNPs) neighborhood patterns in cattle were determined using Bayesian analysis on 15,110 SNPs, each with flanking sequences of 500 bp. Approximately 70% of SNPs were transitions. Transversion SNPs occurred most frequently (38.6%) when the A+T content equaled two at their immediate adjacent sites. In addition, C\rightarrow T and A\rightarrow G transitions and A\rightarrow C and G\rightarrow T transversions had reverse complementary neighborhood patterns. This study revealed several previously unreported SNP neighborhood patterns. First, cattle and humans share an overall SNP pattern, which is indicative of a common mutation system. Second, true neighborhood patterns for A\rightarrow T and C\rightarrow G are unclear since the sense and antisense sequences flanking these mutations are not recognizable. Third, the neighborhood ratio between A+T and G+C was very different, being lowest for C\rightarrow G, followed by C\rightarrow T/A\rightarrow G, A\rightarrow C/G\rightarrow T, and highest for A\rightarrow T. Fourth, transitions were significantly increased when two immediate adjacent sites provided structures for CpG. Lastly, methylation induced deamination reactions were responsible for approximately 20% of all transitions due to the unequal occurrence between A\rightarrow G and C\rightarrow T in five paired neighboring structures. The conversion occurred at both CpG and non-CpG sites.

**Poster 5046**

**Title:** Mangalarga Marchador horse: genetic diversity and population structure

Presenting Author: Maria Raquel Santos Carvalho, Departamento de Biologia Geral, Instituto de Ciências Biológicas, Universidade Federal de Minas Gerais, Belo Horizonte - MG - Brazil

Other authors (name only):
1. DeAssis, J. B.
2. DeLaat, D. M
3. Peixoto, M. G. C. D
4. Bergmann, J. A. G.
5. Fonseca, C. G

**Abstract:**
In order to ascertain genetic diversity and structure for Mangalarga Marchador breed, 115 unrelated animals, born from October/2004 and May/2005, were sampled from three regions of Minas Gerais State, Brazil (South, Southeast and Northeast) and tested for 10 microsatellite loci. Under Hardy-Weinberg assumptions, seven markers were at equilibrium (LEX014, LEX017, LEX019, SGCV23, TKY321, VHL20, and VIASH39) while two (ABS3 and LEX031) presented significant homozygotes excess. Total number of alleles identified was 75, with a mean number of alleles of 8.22. Mean heterozygosity was 0.637 and mean PIC (Polymorphism Information Content) was 0.662. Most informative markers (PIC>0.7) were ABS3, LEX019, SGCV2, TKY321, and VHL20. FST values (0.06) indicates a mild geographical structure. A Bayesian based cluster analysis under a three cluster model was used to estimate population structure: 93.9% of the 115 animals were correctly assigned to the subpopulations from where they were sampled out. Comparisons of pairwise inbreeding coefficients resulted in 11% of the pairs sharing more than 20% of the alleles, thus suggesting that these animals are closely related. These results suggest that the population is genetically structured. Gene flow occurs between South and Southeast and South and Northeast regions, but less frequently between Southeast and Northeast regions.