The periodic coloration in birds forms through a prepattern of somite origin
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Understanding APP processing in Alzheimer disease using a mouse model of Down syndrome

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Trisomy of human chromosome 21, which causes Down syndrome (DS), is the most common genetic cause of dementia. By the age of 40, virtually all people with DS have amyloid plaques and neurofibrillary tangles, the main pathological hallmarks of Alzheimer disease (AD). The main component of amyloid plaques, amyloid beta, is produced through sequential cleavage of Amyloid Precursor Protein (APP). The APP gene is present on chromosome 21, but other genes are likely to also have a role in AD in DS (AD-DS). Our aim is to study the effect of triplication of chromosome 21 genes other than APP on APP processing. We use Mouse Embryonic Fibroblasts (MEFs) derived from a DS mouse model (Dp1Tyb) that carries a segmental duplication of 23Mb of mouse chromosome 16, homologous to chromosome 21. We will focus on the expression and half-life of APP-derived fragments and on alteration in the structure of endosomes, a main site of APP processing. The ultimate aim of our research is to identify the genes responsible for altered APP processing in AD-DS. To do this we plan to use mouse models that have segmental duplications of progressively smaller regions of the mouse chromosomes homologous of human chromosome 21.

Cohesin is continuously required to sustain neuronal gene expression

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Cornelia de Lange Syndrome (CdLS) is a developmental disorder caused by mutations that compromise the function of cohesin, a major regulator of 3-D genome organisation. A universal yet unexplained feature of CdLS is cognitive impairment. In particular, it is not known whether neuronal dysfunction arises primarily from developmental defects in neuronal progenitors, or from an ongoing requirement for cohesin in post-mitotic neurons. To address these questions, we investigated the impact on neuronal gene expression of depleting cohesin at distinct points in development - in the germline, in immature post-mitotic neurons, or in mature neurons. Unexpectedly, neuronal genes with key roles in synaptic transmission, connectivity, neuronal development and signaling were selectively downregulated, regardless of whether cohesin function was compromised throughout development, during the differentiation of immature post-mitotic neurons, or in mature neurons for just 24 hours. Our findings establish that cohesin is required to sustain the expression of functionally important neuronal genes, and provide a tractable approach to neuronal dysfunction in CdLS.
Is apical constriction synchronised with cell cycle progression in mammalian neurulation?

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Embryonic epithelia generate mechanical forces which sculpt the embryo. During neural tube closure, actomyosin-dependant neuroepithelial apical constriction and interkinetic nuclear migration (IKNM) facilitate apposition of the posterior neuropore (PNP) neural folds. Both apical constriction and IKNM are force generating mechanisms which co-regulate epithelial apical dimensions, but how they co-exist in the neuroepithelium is unknown.

To investigate this, E9 CD1 mouse embryos were cultured in the Rho-associated kinase (ROCK) inhibitor Y-27632. Progression past S-phase was blocked with hydroxyurea. Whole-mount confocal immunofluorescopy was used to visualise cell cycle markers (G2→M: pH3, M→G1: pRB) and cell outlines (Scrib).

ROCK inhibition widened the PNP and increased neuroepithelial apical dimensions without substantially changing mitotic index. In vehicle-treated embryos, pRB+ cells predominantly had small apical areas. ROCK inhibition (2hrs) significantly increased their apical dimensions. pH3+ cells’ apical dimensions ranged from the smallest to the largest observed; surprisingly, this was unchanged by ROCK inhibition (2hrs). This suggests neuroepithelial cells adopt small apical dimensions independently of ROCK during M-phase, but require ROCK to maintain apical constriction when entering G1. Trapping neuroepithelial cells in S-phase for 8hrs narrowed the PNP and prevented its widening following ROCK inhibition. Thus, ROCK-dependent apical constriction is temporally coordinated with IKNM during PNP closure.

The role of cellular senescence and its associated secretome in paediatric Adamantinomatous Craniopharyngioma (ACP)

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ACP is the most common non-neuroepithelial developmental brain-tumour. Mouse models generated in our group have demonstrated that mutations in CTNNB1 during development can drive this tumour-formation. SOX2 + ve pituitary stem cells targeted with oncogenic beta- catenin form senescent cell clusters that can induce tumourigenesis non-cell-autonomously, through a senescent-associated-secretory-phenotype (SASP), comprised of cytokines, chemokines, growth factors, etc. To elucidate the role of senescence in ACP I am generating a mouse model (p21FDR/+) allowing detection and ablation of senescent cells in-vivo. Selective ablation of cluster cells at time-points during tumour formation will determine whether the cells are required for tumour initiation, or solely in tumour maintenance and growth. Additionally a Rosa26St31+/mRFP+/g mouse will attenuate activities of SASP genetically, to determine the crucial secretory drivers of tumourigenesis, as well as to determine the developmental roles of cytokines and chemokines in the pituitary. This will provide insight into senescence in ACP pathogenesis and pituitary development.

Epigenetic revertant mosaicism in congenital melanocytic naevi – proof of concept for allele-specific silencing

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Revertant mosaicism is the process of in vivo correction of a mutation which results in phenotypic rescue, previously only described in autosomal recessive skin diseases with secondary somatic DNA mutations. We describe here epigenetic revertant mosaicism in three patients with congenital melanocytic naevi (CMN), a condition caused by in utero somatic heterozygous mutations in critical signaling pathway gene NRAS. Patients presented with new islands of normal skin within naevi, and biopsies were obtained from both areas. Unexpectedly, the known DNA mutation was present in all areas, as were equal numbers of naevas cells, however the expression within the phenotypically normal skin was strikingly monoallelic, and restricted to the normal allele. This was confirmed using an optimized and highly specific gene expression assay for the mutant alleles. Investigation of the underlying mechanism has excluded secondary promoter mutations in NRAS, and allele-specific differences in CpG methylation. This phenomenon of natural genetic therapy in vivo is proof of concept that therapeutic allele-specific targeting of gene expression would be sufficient to correct the clinical phenotype.

Refining mouse models for preclinical studies

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MRC Harwell is involved in many national and international projects using mouse models to study the relationship between genes and disease. We have many resources available to the biomedical community which we are keen to share.

Archiving - We have a frozen embryo and sperm archive where we can cryopreserve your mouse strains for free. These can then be sent anywhere in the world.

Genome Editing Mice for Medicine - Our Genome Editing Mice for Medicine programme is an MRC programme aimed at generating scientifically important, novel mouse lines taking advantage of advances in genome editing. UK scientists are invited to nominate genetically altered mouse lines to be made to advance their own research and be of widespread use in biomedical science.

Training - We have a range of different training courses for laboratory and animal science. These courses not only include technical courses in transgenesis, pathology and cryopreservation but also taught courses in mouse genetics and genome editing.
**IMPC** - We are a member of the International Mouse Phenotyping Consortium (IMPC) a global programme looking to find the function of every protein-coding gene in the mouse genome, with the data freely accessible online. If you are interested in a particular gene knockout, you can nominate it through us.

**Epithelial dynamics during development of the mammalian ear canal**

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Defects in ear canal development can cause severe hearing loss, however very little is known about how the canal initiates, extends and opens. Here we have studied mammalian ear canal development and show that the canal undergoes a complex system of closure and reopening as it forms. The more superficial part of the canal formed from an open primary canal, initiating at the junction between the first and second arch, which later collapsed and then reopened. In contrast, the deeper part of the canal formed from a solid meatal plate that extended from the primary canal into the first arch and later opened. As the ear canal developed, the different parts displayed distinct patterns of proliferation and keratin expression, with collapse of the primary canal linked to loss of periderm. Final opening of the canal was triggered by terminal differentiation of the epithelium. Interestingly, the meatal plate opened asymmetrically, associated with differential proliferation, to create the thin outer surface of the ear-drum. Understanding these complex processes involved in canal development can shed light on the underlying causes of canal atresia.

**Congenital Macular Dystrophy is caused by non-coding duplications downstream of IRX1 locus**

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Autosomal dominant North Carolina Macular Dystrophy (NCMD) is believed to represent a failure of macular development with consequent central vision loss. Prior genetic linkage pinpointed the disease locus to chromosome 6q16 and 5p15. The aim was to identify causative variants and further the understanding of the disease mechanism.

CRISPR technology was used to reproduce a 50 kb non-coding tandem duplication between IRX1 and ADAMTS16 (chr5 in hg19 and chr13 in mm9) recurrently identified in NCMD families by whole genome sequencing. Gene expression studies were performed on developing fetal eye and limb-bud. Chromosome conformation capture (c-Hi-C) and chromatin immunoprecipitation performed on the same tissues, mapped chromatin architectural folding and defined putative regulatory elements.

Expression of IRX1 and IRX2 was increased in the limb-bud tissue at E12.5 in the mouse model versus control. Subsequently these genes are downregulated at E14.5, suggesting temporal dysregulation of the IRXA cluster. c-Hi-C at the 5p locus showed unique chromatin interactions in developing wild-type eye versus limb-bud, suggesting distinct tissue-specific regulation.

Chromatin organization findings highlight the relevance of tissue specific DNA contacts in development and disease. Further characterization of mouse model and exploration of the nine unsolved cases may uncover additional molecular targets for NCMD.

**Tctex1d2, an integral transporter for (Intraflagellar Transport) IFT components during sperm development**

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Juvenile syndrome is a rare autosomal recessive ciliopathy affecting about 1 in 100,000 – characterized by shortening of long bones, constriction of the thorax and polydactyly. Here we characterize a Tctex1d2 gene trap mouse model creating successful knock-down of gene expression particularly in the testes and bone. Tctex1d2 is a light chain of the ciliary intra-flagellar transport (IFT) dynein-2 motor complex, essential for motor stability and trafficking of protein cargos required for ciliogenesis. Although no gross skeletal abnormalities were evident, micro-CT scans revealed mild elevation in long bone (tibia) mass confirming a skeletal phenotype. Furthermore, a significant reduction in sperm number and motility was observed in mutant epididymis with several spermato genesis defects including amorphous heads, short/bent flagella and swollen tail tips. During spermatid differentiation, protein transport is based on IFT and intra-manchette transport. In the mutants, Tctex1d2 protein was seen to be absent in the flagellum of developing spermatids and mature sperms. Hence, to determine the stage of spermato genesis at which Tctex1d2 is required for cellular cargo trafficking selective IFT components were studied. We found markedly increased staining along the flagella of IFT components in the mutants, suggesting that transport of IFT proteins is disrupted. We therefore conclude that Tctex1d2 is an integral component of the sperm retrograde IFT machinery.

**Characterisation of a SOX2-positive population in the postnatal adrenal medulla.**

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The adrenal gland is an endocrine organ responsible for the stress response and is involved in the regulation of the immune system and metabolism. The adrenal gland is composed of an outer cortex and an inner medulla, which have distinct functions. It is a dynamic organ, capable of responding to the variable physiological demand requiring production of steroids and catecholamines. While stem and progenitor cell populations in the adrenal cortex have been identified and characterised, investigation of the adrenal medulla cell hierarchy is still ongoing.

The adrenal medulla derives from the neural crest, from which a common progenitor generates both sympathetic neurons and neuroendocrine chromaffin cells. SOX2 is a marker of multiple stem cell/progenitor lineages and is found to be upregulated in many tumours including pheochromocytomas and parangliomas, tumours of the adrenal medulla.

Here we identify and characterise SOX2-positive cells in the murine adrenal medulla during homeostasis. SOX2-positive cells do not colocalise with other known progenitor markers in the medulla. Using genetic lineage tracing in the juvenile postnatal animal, we identify SOX2-positive cells to be an expanding population which gives rise to chromaffin cells. Taken together our data point towards a new candidate progenitor cell population of the adrenal medulla.

**The periodic coloration in birds forms through a prepattern of somite origin**

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Spots, stripes… These periodic patterns often seen in animals have been largely viewed as self-organizing. Do they also depend on preexisting positional information? In juvenile galliform birds we show that signaling from the somite sets the position of stripes on preexisting positional information? In juvenile galliform birds, we show that signaling from the somite sets the position of stripes on preexisting positional information. In juvenile galliform birds, we show that signaling from the somite sets the position of stripes on preexisting positional information?

**Surface ectoderm biomechanics depend on Grainyhead-like protein 2 (Grhl2) to regulate neural tube closure**

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Grainyhead-like proteins (Grhls) are evolutionary conserved transcription factors essential for neural tube closure. Grhl2 knockout models display both cranial and spinal neural tube defects (NTDs), whereas the Axial defects (Axd) model, a Grhl2 overexpressing model, has spinal defects. Grhl2 is expressed in the non-neural ectoderm or surface ectoderm, a monolayer of cells covering and wrapping around the neural folds that ultimately will give rise to the epidermis. We investigated the molecular and cellular mechanisms by which either excess or insufficient Grhl2 expression can cause NTDs. We performed RNA-seq experiments on both knockout and overexpressing models (Axd) and identified multiple pathways regulated inversely in both models. Genes encoding proteins of the junctional apical complex were highly regulated in both datasets. We further validated a number of adherens junction molecules (such as E-cadherin) and tight junction molecules (a number of claudins) to be regulated by Grhl2 in the caudal neural tube. Consistent with the fact that these molecules are pivotal for cell-cell adhesion and connected to the cytoskeleton, we also identified abnormalities in F-actin and p-MLCII organization. In addition, absence of Grhl2 altered the nature of the surface ectoderm cells which acquired a more neuroepithelial phenotype. Finally, functional experiments showed that the surface ectoderm cells in the presence of elevated or diminished Grhl2 expression, were under abnormal tension. In conclusion, Grhl2 defines the epithelial nature of the surface ectoderm monolayer, making it a determining factor for neural tube closure.

**Increased repair in neural crest frontal bones correlates with recruitment of sutural stem cells**

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The craniofacial skeleton is formed from the neural crest and mesodermal lineages, both of which contribute mesenchymal precursors during formation of frontal and parietal bones respectively. Increased healing capacity in frontal bones has been previously described. In culture, frontal bone cells are more osteogenic than parietals, as seen by formation of more mineralized nodules. Although differences in molecular signalling and osteogenic potential have been well studied, the recruitment of suture mesenchymal cells and specific stem cell subsets has not yet been investigated. Here we combine lineage tracing techniques in a calvarial wound model to assess the infiltration of previously reported stem cell populations, as well as the availability and recruitment of neural crest derivatives. 2 mm defects were performed on frontal and parietal bones of 6 weeks old mice. Gli1-CreERT; Rosa26R;R26R and Axin2-CreERT; Rosa26R mice were induced with tamoxifen 2 days prior to osteoctomy, whereas Wnt11-Cre; Rosa26R was constitutively active. We found that, within the first week post-surgery, not only a larger number of labelled stem cells reside on the inter-frontal suture when compared to the sagittal, but a higher proportion of them infiltrate the frontal wound in comparison to the parietal. Moreover we provide insights on the neural crest domain at the sagittal suture, which suggest that the position in relation to the surrounding sutures might be a determinant on the efficiency of parietal wound healing.
Postnatal skull development of c-Fos mice and the importance of osteoclasts across ontogeny

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The dynamic nature of bone remodelling enables skeletal elements to adapt to their environment, thus playing a crucial role in normal development. Here, we investigated the interaction between abnormal bone remodelling (c-Fos knockout mice) and postnatal skull growth by utilising 3D morphometrics to characterise phenotypic change and link such changes to cellular involvement. MicroCT scans indicated a delayed development of c-Fos mutants compared with WT littermates, illustrated by their domed skull phenotype which lacked tooth eruption and displayed altered morphology of several bones: parietal, nasal and mandible. The two principle components (PCs) contributing to overall shape variation were skull height (37%) and width (31%) with the c-Fos mutants occupying the heightened and wider region of morphospace. Interestingly, this phenotype of the mutants increasingly diverged from WT littermates across postnatal ontogeny. Unexpectedly, microCT reconstructions indicated premature suture fusion (craniosynostosis) in c-Fos mutants. In WT mice at early stages of postnatal development (P4), TRAP-positive osteoclasts lined the bone margins along the patent coronal and sagittal sutures. At later stages of development (P20), osteoclast activity had decreased and was associated with increased bone deposition at the suture locale. Our results suggest an unappreciated role for osteoclasts in maintain suture patency across ontogeny.