The most common human ADAR1p150 Zα domain mutation P193A is well tolerated in mice.

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Summary
ADAR1 mediated A-to-I RNA editing is a self/non-self discrimination mechanism for cellular double stranded RNAs. ADAR mutations are one cause of Aicardi-Goutières Syndrome, an inherited paediatric encephalopathy, broadly classed as a “Type I interferonopathy”. The most common ADAR1 mutation is a proline 193 alanine (p.P193A) mutation, mapping to the ADAR1p150 isoform specific Zα domain. We report the development of an independent murine P195A knock-in mouse, homologous to the human P193A mutation. The Adar1P195A/P195A mice are largely normal and the mutation is well tolerated. Contrasting with previous reports when the P195A mutation was compounded with an ADAR1 null allele, the majority of mice have only a modest reduction in weaning weight and survived long-term. Severe runting and shortened survival of Adar1P195A−/− animals are dependent on the parental genotype. The P195A mutation is well tolerated in vivo and the loss of MDA5 is sufficient to completely rescue the Adar1P195A−/− mice.

Keywords: ADAR1, A-to-I RNA editing, P193A mutation, Zα domain, MDA5, dsRNA, innate immunity
Introduction

One of the most common RNA modifications in mammals is the deamination of adenosine to inosine, termed A-to-I editing, in double stranded regions of RNA (dsRNA) by Adenosine Deaminase Acting on RNA 1 (ADAR1) and ADAR2 (Eisenberg and Levanon, 2018). A-to-I editing results in the non-reversible conversion from adenosine to inosine for the targeted nucleotide within the RNA. Inosine is usually interpreted as a guanosine during translation. Depending on where the editing site lies in the transcript A-to-I editing can have a variety of consequences (Eisenberg and Levanon, 2018; Solomon et al., 2017; Solomon et al., 2013; Walkley and Li, 2017). Editing within protein-coding sequences can change the amino acid codon, and therefore the protein product, from that genomically encoded (Licht et al., 2019). A-to-I editing can also impact RNA splicing, stability, translation, localization (Kapoor et al., 2020; Lev-Maor et al., 2007; Shoshan et al., 2015; Stellos et al., 2016) as well as the biogenesis of non-coding RNAs (microRNAs (Kawahara et al., 2007) and circRNAs (Ivanov et al., 2015)). Editing alters the base-pairing properties within RNA, both stabilising and destabilising the RNA secondary structure depending on context (Liddicoat et al., 2015; Solomon et al., 2017). A-to-I editing can be readily detected using sequencing methods, as A-to-I editing can be identified by A-to-G mismatches between the cDNA and genomic DNA, allowing genome-wide mapping (Levanon et al., 2004; Li et al., 2009; Ramaswami et al., 2013).

There are millions of A-to-I editing sites across all tissues within the human transcriptome, with the majority located in repetitive elements such as Alu repeats (Bazak et al., 2014; Gabay et al., 2022; Picardi et al., 2015; Tan et al., 2017). In mice there are between 50,000-150,000 editing events, also concentrated in evolutionarily related repetitive elements (SINE/LINEs) (Costa Cruz et al., 2020; Licht et al., 2019; Pfaller et al., 2018; Pinto et al., 2014). The key physiological function of ADAR2 is to recode the Gria2 transcript by editing a coding region of the transcript, resulting in an amino acid substitution and a correctly functioning GLUR2 protein (Higuchi et al., 2000; Higuchi et al., 1993). In contrast to the recoding of a single relevant mRNA target for ADAR2, it has been demonstrated by multiple groups that the physiologically most important function of ADAR1 editing is to prevent the cells own RNA being mistaken as foreign RNA by the innate immune system (Liddicoat et al., 2015; Mannion et al., 2014; Pestal et al., 2015). ADAR1’s primary physiological function is to counteract cytoplasmic innate immune sensing by melanoma differentiation-associated protein 5 (MDA5) of endogenous RNAs (Chalk et al., 2019; Hartner et al., 2009; Heraud-Farlow et al., 2017; Liddicoat et al., 2015), a species conserved function (Chung et al., 2018; Pestal et al., 2015). ADAR1 is expressed as two isoforms, a constitutive and primarily nuclear p110 protein and an inducible and mostly cytoplasmic p150 isoform. Recent studies have focussed attention on the role of...
the cytoplasmic p150 isoform and how this intersects with innate immune sensing (Kim et al., 2021; Pestal et al., 2015; Ward et al., 2011).

**ADAR (ADAR1)** loss of function mutations have been identified as one of the genetic causes of Aicardi-Goutières Syndrome (AGS) (Rice et al., 2012; Rice et al., 2017). AGS is an inherited paediatric encephalopathy, classed as a “Type I interferonopathy”, characterised by the upregulation of interferon (IFN) and interferon stimulated gene (ISG) expression (Crow and Manel, 2015). AGS patients with **ADAR** mutations most often have compound heterozygous mutations, with one mutation impacting the p150 isoform together with a second that affects both p110 and p150 isoforms (Rice et al., 2012; Rice et al., 2017). Mutations in **ADAR** have also been identified in bilateral striatal necrosis (BSN), where patients have a dystonic or rigid movement disorder associated with symmetrical abnormalities of the brain (Livingston et al., 2014). Similar to **ADAR** mutant AGS, BSN patients with **ADAR** mutation have a characteristic “interferon signature”. Recent knock-in mouse models have begun to address how distinct **ADAR1** mutations, including those reported in AGS and BSN, impact the function of **ADAR1** *in vivo* (de Reuver et al., 2021; Guo et al., 2021; Inoue et al., 2021; Maurano et al., 2021; Nakahama et al., 2021; Tang et al., 2021).

One of the most frequently reported **ADAR1** mutations is proline 193 to alanine (p.Pro193Ala; P193A) (Rice et al., 2017), which maps to the unique Zα domain of the p150 isoform (Herbert, 2021; Herbert et al., 1995; Herbert and Rich, 2001; Herbert et al., 1998; Nakahama and Kawahara, 2021). The Zα domain is found in only one other protein in the human genome, Z-DNA binding protein 1 (ZBP1), that activates cell death pathways during viral infection and in animals deficient in Receptor interaction protein kinase 1 (*Ripk1*) and caspase 8 (*Casp8*) (Jiao et al., 2020; Newton et al., 2016; Zhang et al., 2020). Proline 193 contacts the left-handed helix structure characteristic of Z-form DNA and RNA and is important in the interaction between **ADAR1**p150 and Z-form nucleic acid (Schwartz et al., 1999). Interestingly, unlike other **ADAR** mutations, the P193A mutation is present at an allele frequency of 0.002 of the human population globally (Herbert, 2020; Karczewski et al., 2020), however the effects of this mutation are not definitively understood. In human populations the P193A mutation is nearly always heterozygous (Karczewski et al., 2020). Maurano et al., recently reported development and characterisation of a murine p.Pro195Ala (P195A) mutant mouse model, homologous to the human P193A mutation (Maurano et al., 2021). They reported that the P195A allele was well tolerated in isolation and could be homozygous viable with a normal lifespan. When combined with a second mutation, either a p110/p150 null allele or p150 null allele, they reported a significantly shortened lifespan and reduced weaning weights, with
evidence of activation of an innate immune/interferon and integrated stress response gene expression program in the \textit{Adar1}^{P195A/p150}. They report rescue of the phenotypes by loss of MDA5 consistent with prior genetic rescue of ADAR1 loss of function alleles (de Reuver \textit{et al}., 2021; Liddicoat \textit{et al}., 2015; Nakahama \textit{et al}., 2021; Pestal \textit{et al}., 2015). However, unlike ADAR1 null/editing dead models (Liddicoat \textit{et al}., 2016b; Mannion \textit{et al}., 2014; Wang \textit{et al}., 2004), they also reported normalisation of both survival and weaning weight by concurrent loss of LGP2, IFNAR or PKR.

Here, we report generation and analysis of an independent \textit{Adar1}^{P195A} mutant mouse model. Consistent with the published model (Maurano \textit{et al}., 2021), we find that the P195A allele is well tolerated when heterozygous or homozygous. When the P195A mutation was combined with either a p110/p150 null allele or a point mutation that renders both p110 and p150 editing deficient (E861A), we observed long term survival of the majority of mice across all genotypes. Adult \textit{Adar1}^{P195A/-} and \textit{Adar1}^{P195A/E861A} animals had evidence of a mild ISG signature but not of activation of a PKR dependent integrated stress response. The loss of \textit{Ifih1} (MDA5), the primary sensor of unedited cellular dsRNA, prevented activation of the ISG response and rescued long term survival of all genotypes tested. These data do not support the conclusion that the P195A allele leads to fully penetrant PKR-dependent pathology and shortened lifespan in mice.
Materials and Methods

Ethics Statement
All animal experiments conducted for this study were approved by the Animal Ethics Committee of St. Vincent’s Hospital, Melbourne, Australia (Protocol number #016/20). Animals were euthanized by cervical dislocation or CO₂ asphyxiation.

Animals

Adar\textsuperscript{1P195A} mice were generated using CRISPR/Cas9 targeting in C57BL/6 zygotes by the Monash Genome Modification Platform (Monash University, Clayton, Australia). A repair oligo containing a CCT>GCT point mutation resulting in a p.P195A (Proline>Alanine) mutation. The repair oligo also had a silent point mutation (CCT>CCC; Proline>Proline, p.P194P) immediately upstream and a second silent mutation (TTG>CTG; Leucine>Leucine, p.L196L) immediately downstream of the P195A mutation, respectively, that created a \textit{BsrB1} restriction enzyme site and prevented Cas9 from cutting the repaired locus and to be used for genotyping using restriction digest of the genotyping PCR product. Introduction of the mutation was confirmed by Sanger sequencing of the region in both the founders and subsequent generations.

The \textit{Adar}\textsuperscript{E861A/+} (Adar\textsuperscript{1E861A/+}; MGI allele: Adar\textsuperscript{tm1.1Xen}; MGI:5805648) (Liddicoat \textit{et al.}, 2015), Adar\textsuperscript{-/-} (Adar\textsuperscript{1/-}; exon 2-13 deleted; MGI allele: Adar\textsuperscript{tm2Phs}; MGI:3029862) (Hartner \textit{et al.}, 2004), Adar\textsuperscript{fl/fl} (Adar\textsuperscript{1fl/fl}; exon 7-9 floxed; MGI allele: Adar\textsuperscript{tm1.1Phs}; MGI:3828307) (Hartner \textit{et al.}, 2004; Hartner \textit{et al.}, 2009), Ifih1\textsuperscript{1/-} (Ifih1\textsuperscript{1tm1.1Cln}) (Gitlin \textit{et al.}, 2006), and Rosa26-CreERT\textsuperscript{2} (Gt(ROSA)26Sor\textsuperscript{tm1(cre/ERT2)Tyj}) (Ventura \textit{et al.}, 2007) mice have all been previously described and were on a backcrossed C57BL/6 background. All animals were housed at the BioResources Centre (BRC) at St. Vincent’s Hospital. Mice were maintained and bred under specific pathogen-free conditions with food and water provided \textit{ad libitum}. For acute somatic deletion model (\textit{R26-CreERT\textsuperscript{2}}), all animals were ≥8 weeks of age at tamoxifen initiation; Tamoxifen containing food was prepared at 400mg/kg tamoxifen citrate (Selleckchem) in standard mouse chow (Specialty Feeds, Western Australia).

Genotyping

Genotyping of the P195A mutants was determined by PCR. The repair oligo carrying the P195A mutation also introduced a silent \textit{BsrB1} immediately upstream of the P195A mutation. The digestion of the genomic DNA PCR product was used to determine presence of the P195A allele and can discriminate heterozygous and homozygous mutants. The following primers: Primer P1 (5’-ACCATGGAGAGGTGCTGACG-3’) and P2 (5’-ACATCTCGGGCCTTTGGTGAG-3’) were used to obtain a 489bp product from the wild-type allele, which would yield two fragments (265bp and 224bp product) when digested with \textit{BsrB1}. 
(NEB) for the P195A mutant. The P1 primer was used for Sanger sequencing (Australian Genome Research Facility (AGRF), Melbourne) of the purified PCR product as required. Genotyping of all other lines used was performed as previously described (Heraud-Farlow et al., 2017; Liddicoat et al., 2015).

**Histology**

Three wild-type (Adar1 \(^{+/+}\), 2 female (F)/1 male (M)), three Adar1 \(^{P195A/+}\) (2F/1M) and four Adar1 \(^{P195A/E661A}\) (3F/1M) animals at 6-7 months of age were used for histopathology examination as previously described (Chalk et al., 2019; Heraud-Farlow et al., 2017). The wild-type animals were littermate controls of the mutant bearing animals. Tissue collection and histology was performed by the Phenomics Australia Histopathology and Slide Scanning Service, University of Melbourne. The wild-type animals were identified to the pathologists as “controls”, the remaining samples were genotype blinded to the staff and pathologist assessors. The following organs were assessed: adrenal glands, bladder, bone marrow, brain, cecum, cervix, colon, duodenum, epididymes, eyes, gall bladder, harderian glands, head, heart, hind leg (long bone, bone marrow, synovial joint, skeletal muscle), ileum, jejunum, kidney, liver, lungs, mammary tissue, mesenteric lymph node, ovaries, oviducts, pancreas, penis, preputial gland, prostate glands, salivary glands and regional lymph nodes, seminal vesicles, skin, spinal cord, spleen, stomach, tail, testes, Thymus, thyroids, trachea, uterus, vagina. The full pathology report and genotypes is available in Supplemental Dataset 1.

Additional pathology was performed on 13-19 week old brain, liver, kidney and spleen isolated from wild-type (bred and housed in the same facility, not littermates of mutant animals; n=3), Adar1 \(^{P195A/+}\) (n=3), Adar1 \(^{P195A/-}\) (n=4), Adar1 \(^{P195A/-}/Ifih1^{-/-}\) (n=3). Tissue was fixed overnight in 2% paraformaldehyde, transferred to 70% ethanol and then processed, sectioned and stained by the Phenomics Australia Histopathology and Slide Scanning Service, University of Melbourne. The wild-type animals were identified to the pathologists as “controls”, the remaining samples were genotype blinded to the staff and pathologist assessors. The full pathology report and genotypes is available in Supplemental Dataset 2.

**Mouse Embryonic Fibroblasts**

Mouse embryonic fibroblasts (MEFs) were generated from E13.5 embryos of the indicated genotypes. The embryos were dissected, removing the head (used for genotyping), heart and foetal liver, and the remaining tissue was used to generate MEFs. The tissue was placed in 1mL of 0.025% Trypsin-EDTA (Gibco/Thermo Fisher) then drawn through an 18G needle/1mL syringe, then placed at 37°C in a 10cm\(^2\) tissue culture plate for 30 minutes. After 30 minutes 10mL of media (High glucose DMEM (Sigma), 10% FBS (not heat inactivated, Assay Matrix), 1% Penicillin/Streptomycin (Gibco), 1% Glutamax (Gibco), 1% amphotericin B (Sigma;
250µg/mL stock) was added to the plate and the contents dispersed. The MEFs were incubated in a hypoxia chamber flushed with 5% oxygen/5% carbon dioxide in nitrogen at 37°C. Once the cells were 70% confluent, the cells were trypsinized and passaged onto 10cm plates in normoxic conditions for all further culture. MEFs of the indicated genotypes were treated with recombinant murine interferon beta (PBL Assay Science; PBL-12405) at 250U/mL for 24hrs in normal growth media. After 24 hours, cells were collected by trypsinization and pellets washed in cold PBS and resuspended in RIPA buffer (20mM Tris·HCl, pH8.0, 150mM NaCl, 1mM EDTA, 1% sodium deoxycholate, 1% Triton X-100, 0.1% SDS) supplemented with 1x HALT protease inhibitor and 1x PhosSTOP phosphatase inhibitor (Thermo). Lysates were used for western blotting as described below.

**Protein extraction and Western blotting**

Protein was quantified using the Pierce BCA protein assay kit (Thermo Fisher Scientific) on an Enspire multimode plate reader (Perkin Elmer). Lysates from MEFs of the indicated genotypes +/- interferon treatment were used. 40µg of protein extract per sample was loaded on pre-cast NuPAGE™ 10%, Bis-Tris (1.5 mm, 10-well) polyacrylamide gels (Invitrogen) and transferred onto Immobilon-P PVDF membranes (Merck Millipore). Membranes were blocked with 5% milk in Tris-buffered saline with tween (TBST) and incubated at 4°C overnight with rat monoclonal anti-mouse ADAR1 antibody (clone RD4B11) (Liddicoat et al., 2015), mouse anti-Actin (Sigma Aldrich, A1978). Membranes were then probed with HRP-conjugated goat anti-rat (Thermo Fisher Scientific, 31470) or anti-mouse (Thermo Fisher Scientific, 31444) secondary antibodies and visualized using ECL Prime Reagent for chemiluminescent detection on Hyperfilm ECL (Amersham).

**Peripheral blood analysis**

Peripheral blood (approximately 100µl) was obtained via retro-orbital bleeding. The blood was red blood cell-depleted using hypotonic lysis buffer (150mM NH₄Cl, 10mM KHCO₃, 0.1mM Na₂EDTA, pH7.3) and resuspended in 50µl of FACS buffer for flow cytometry analysis.

**Flow cytometry analysis**

Antibodies against murine B220 (conjugated to APC-eF780), Mac-1 (PE), Gr1 (PE-Cy7), F4/80 (APC), CD4 (eF450), and CD8 (PerCP-Cy5.5) were all obtained from eBioscience, BioLegend or BD Pharmingen (Heraud-Farlow et al., 2017; Liddicoat et al., 2015; Singbrant et al., 2011; Smeets et al., 2014). Cells were acquired on a BD LSRIIFortessa and analyzed with FlowJo software Version 9 or 10.0 (Treestar).
RT-qPCR

Whole bone marrow and myeloid cells +/- tamoxifen treatment were collected, and RNA isolated using the RNeasy kit with on-column DNase digestion (Qiagen). Mouse tissues (one brain hemisphere, liver, kidney; additional histology performed on tissue from these same animals) from independent biological replicates from 14-19 week old C57BL/6 (n=3; 1F/2M; bred and housed in same facility), Adar1P195A/+ (n=3; 1F/2M), Adar1P195A-/ (n=4; 2F/2M), Adar1P195A+/Ifih1+/- (n=3; 2F/1M) and Adar1P195A-/Ifih1-/- (n=3; 2F/1M). Tissues were collected and immediately snap frozen in liquid nitrogen and stored at -80°C. Frozen tissues were homogenized in Trisure reagent (Bioline) using IKA T10 basic S5 Ultra-turrax Disperser. RNA was extracted using Direct-Zol columns (Zymo Research) as per manufacturer’s instruction. cDNA was synthesized using Tetro cDNA synthesis kit (Bioline; used for all cDNA described). Real-time PCR was performed using two methods: SYBR green method (for Ifit1, Irf7 and Ifnb when measured in myeloid cell lines and in vivo R26-CreER experiments, normalised to Ppia; Fig 5 and Fig 6) as previously described (Heraud-Farlow et al., 2017). Alternatively predesigned Taqman probes for Oas1a, Ifi27, Irf7, Asns, Cdkn1a, Hmox1 were used and normalised to Ppia and Hprt (for mouse organs from germ-line mutant animals in Fig 4). Duplicate reactions per sample were measured using an AriaMx Real-time PCR machine (Agilent) using TaqMan Fast Advanced Master Mix (Applied Biosystems, ThermoFisher Scientific) and predesigned/inventory FAM conjugated Taqman primer/probe sets (Applied Biosystems, ThermoFisher Scientific) against murine genes: Oas1a (assay ID Mm00836412_m1); Ifi27 (Mm00835449_g1); Irf7 (Mm00516788_m1); Asns (Mm00803785_m1); Cdkn1a (Mm04205640_g1); Hmox1 (Mm00516005_m1) and control gene Hprt (Mm00446968_m1). Hprt was used as reference genes for relative quantification using the ∆∆Ct method.

Immortalized myeloid cells

HOXA9 immortalized myeloid cell lines (Wang et al., 2006) were established by retroviral infection of ficoll depleted bone marrow isolated from three independent adult (>8week old) R26-CreERT2 Adar1+/ and R26-CreERT2 Adar1P195A donor animals. The cells were cultured in IMDM (Sigma) containing 10% FBS (Assay Matrix; non-heat inactivated), 1% Penicillin/Streptomycin (Gibco/Thermo Fisher), 1% Glutamax (Gibco/Thermo Fisher) supplemented with 50ng/mL recombinant mouse stem cell factor (rmSCF, Peprotech), 10ng/mL recombinant mouse interleukin 3 (rmIL-3, Peprotech) and 10ng/mL recombinant human interleukin 6 (rhIL-6, Amgen) for 48 hrs. After 48hr in culture, 1x10⁶ cells were spin-infected at 1,100g for 90 minutes with ecotropic packaged HOXA9 retrovirus (HOXA9 plasmid was generously provided by Dr Mark Kamps, University of California San Diego) and 8ug/mL hexadimethrine bromide (Polybrene; Sigma). At 48 hrs post-infection, the cells were
passaged into IMDM containing 10% FBS, 1% Penicillin/Streptomycin, 1% Glutamax supplemented with 1% granulocyte-macrophage colony-stimulating factor (GM-CSF) conditioned medium (from BHK-HM5 cell conditioned medium). Cells were maintained in GM-CSF containing media after this point. Cell lines established after 3-4 weeks of culture.

Once stable cell lines were established, they were treated for with 200nM 4-hydroxy tamoxifen (Merck Millipore) to activate CreER recombination which results in cells becoming Δ/+ (fl/+ cells become Adar1 heterozygous) or ΔP195A (fl/P195A cells only retain expression of P195A after tamoxifen treatment). Cells were counted with Trypan blue using a Countess II automated counter (Thermo Fisher Scientific) and then passaged every 2-3 days. Viability was assessed by trypan blue staining and counted using a Countess II and gene expression on cDNA made from cells of the indicated genotypes and assessed by SYBR green based qPCR as described previously (Chalk et al., 2019; Heraud-Farlow et al., 2017).

To test if there is an aberrant response in P195A mutant cells toward non-cellular cytosolic dsRNA, Adar1Δ/+ and Adar1ΔP195A cells were treated with high molecular weight polyinosinic:polycytidylic acid (poly(I:C); InvivoGen). R26-CreER Adar1Δ/+ and R26-CreER Adar1ΔP195A cells were treated for 14 days with tamoxifen, genotyped, and then Adar1Δ/+ and Adar1ΔP195A cells were transferred to non-tamoxifen supplemented IMDM media for poly(I:C) testing. Three independent cell lines were used for each genotype. The cells were treated with a dose range of high molecular weight poly(I:C) (1.5-8kb; 1mg/mL stock) by nucleofection following manufacturer’s instructions (Lonza 4D-Nucleofector™ Kits). Gene expression of the indicated genotypes was assessed by SYBR green based qPCR as described previously.

Statistical analysis
To determine statistical significance, Kaplan-Meier survival plots and ordinary one-way ANOVA tests were conducted in GraphPad Prism software version 9 (GraphPad; San Diego, CA, USA). Throughout this study, significance is indicated using the following convention: *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001, and data is presented as mean ± S.E.M. The number of samples used for each experiment is described in the corresponding figure legends.
Results

Generation of an Adar1 P195A knock-in mutant allele

The P193A mutation in human ADAR1 maps to the Za domain that is unique to the ADAR1 p150 isoform. The proline 193 residue of human ADAR1 is homologous to murine proline 195 (Fig 1A). We introduced a C to G point mutation to generate a proline to alanine substitution at amino acid 195 (p.P195A) in murine Adar1 using CRISPR/Cas9 on a C57BL/6 background (Fig 1B). This resulted in the desired mutation that was confirmed by Sanger sequencing and through restriction digest of genomic PCR products, utilising a silent unique restriction site that was introduced during targeting to the modified locus (Fig 1C). After identification of heterozygous founder mice (Adar1P195A/+), these were bred to C57BL/6 mice to confirm germ-line transmission of the mutant allele. Second generation animals were subsequently bred for all experiments.

To assess expression of the ADAR1p150 protein from the mutant allele, we isolated E13.5 mouse embryonic fibroblasts (MEFs) and treated these with murine interferon beta (IFNβ). We tested MEFs from two independent Adar1+/+, Adar1P195A/+ and Adar1P195A/P195A littermates. This demonstrated that the induction of the p150 isoform by interferon was intact and equivalent between Adar1+/+, Adar1P195A/+ and Adar1P195A/P195A cells (Fig 1D).

ADAR1 P195A mutation is well tolerated.

The heterozygous Adar1P195A/+ mice were intercrossed and we recovered viable homozygous animals at the expected frequency, consistent with a previous report (Fig 2A, Supplemental Figure 1B)(Maurano et al., 2021). To understand the consequences of a P195A mutation compounded with a second mutation, more similar to the mutational spectrum reported in AGS (Rice et al., 2017), we crossed the Adar1P195A/+ mice to an Adar1-/+ (Adar1 null allele, exon 2-13 deleted) (Hartner et al., 2004) and in parallel to an editing deficient point mutant model (Adar1E861A/) (Liddicoat et al., 2015) (Supplemental Fig 1). We recovered mice with all possible genotypes from these intercrosses (Fig 2B and data not shown).

We assessed the weaning weights of the different mutants compared to controls genotypes. The Adar1P195A/P195A animals had a higher average weaning weight than wild-type Adar1+/+ mice (Adar1P195A/P195A mean weight = 12.2g, n=19; Adar1+/+ mean weight = 9.9g, n=27; Fig 2C) and higher weaning weights compared to the Adar1P195A- (mean = 8.12g), Adar1E861A/+ (mean = 9.13g), and Adar1P195A/E861A (mean = 8.72g) cohorts (Fig 2C). The Adar1P195A- had significantly lower weaning weights compared to the Adar1-/, Adar1P195A/+ and Adar1P195A/P195A. The Adar1P195A/E861A mice, where the P195A mutation is paired with the editing deficient E861A
mutant p110 and p150 protein, were not significantly different from the $Adar1^{P195A/-}$ cohort and were lighter than the $Adar1^{+/+}$, $Adar1^{P195A/+}$ and $Adar1^{P195A/P195A}$ animals (Fig 2C).

Maurano et al., reported fully penetrant runting of their $Adar1^{P195A/-}$ animals, unlike the variability in weights that we observed in our $Adar1^{P195A/-}$ cohort (Maurano et al., 2021). While the $Adar1^{P195A/-}$ cohort as a whole had significantly lower weaning weights than most other genotypes, they appeared to be clustered into two distinct groups of low or approximately normal weaning weight (Fig 2C). The mean weaning weights across all genotypes in our study (mean range from 8.12 to 12.2g; sexes combined) are consistent with the reference data for 3-4 week old animals of C57BL/6 background mice from both the Jackson Labs and the International Mouse Phenotyping Consortium (Dickinson et al., 2016) (Supplemental Table 1). The wildtype controls reported by Maurano et al. weighing more than 20g at 23 days of age, indicating that they differed from strains we and others have utilised. Upon further analysis, the $Adar1^{P195A/-}$ animals derived from crosses where a $Adar1^{P195A/-}$ was used as a parent were consistently runted (mean weaning weight of $Adar1^{P195A/-}$: 4.23g; n=8; Fig 2E, 2G). In contrast, this was largely ameliorated when a $Adar1^{P195A/P195A}$ was bred to a $Adar1^{+/+}$ animal (mean weaning weight of $Adar1^{P195A/-}$: 9.76g; n=19; Fig 2E, 2G). This result suggests that the breeding pair genotype influences the weaning weight of the compound mutant pups.

Next, we compared the long-term survival of the different genotypes. Cohorts of mice were allowed to age and monitored for any signs of illness or change in health status. The $Adar1^{P195A/+}$ and $Adar1^{P195A/P195A}$ mice did not have any change in long term survival (Fig 2D), consistent with that reported (Maurano et al., 2021). Similarly, when we assessed the survival of the $Adar1^{P195A/E861A}$ animals we found that these mutants survived normally long term (>400 days). The initially reported $Adar1^{P195A/-}$ model used a P195A allele crossed with a germ-line $Adar1$ allele (derived from the germ-line deletion of the floxed Exon 7-9 allele) that was originally reported as null, but later found to express a truncated, unstable, mislocalized editing deficient protein (Bajad et al., 2020; Hartner et al., 2004; Hartner et al., 2009; Pestal et al., 2015). Around 90% of these compound heterozygote $Adar1^{P195A/-}$ mice died by one month of age and only a single animal survived to day 84 (Maurano et al., 2021). In our facility and with our $Adar1$ null allele (exon 2-13 deletion), the $Adar1^{P195A/-}$ mice have survived long term, with >50% of the $Adar1^{P195A/-}$ animals surviving past 120 days of age, and the current oldest >300 days of age (Fig 2D). Since it appeared that there were two distinct groups of $Adar1^{P195A/-}$ mice based on both weaning weight (low or normal, Fig 2C) and survival (die by 60 days or long-term survival, Fig 2D) we further assessed survival based on the breeding pair genotypes. We noted poor survival and outcomes for pups bred from breeding pairs including a $Adar1^{P195A/-}$ parent (2 female and one male $Adar1^{P195A/-}$ used for breeding to date; Fig 2F, 2H). This result
suggested that a significant contributor to the phenotype of the pups was the genotype of the parent, with a $Adar1^{P195A/-}$ parent resulting in runting and poor survival of $Adar1^{P195A/-}$ pups. The $Adar1^{P195A/-}$ genotype more closely approximates that of an AGS patient, and we cannot find literature to indicate if humans with similar genotypes can successfully rear children. Collectively these data indicate that the P195A mutation is well tolerated and is compatible with long term survival, even when compounded with either an ADAR1 null allele or editing deficient mutation. The data further indicate that the increased post weaning mortality of the $Adar1^{P195A/-}$ animals can be fully prevented by provision of an editing deficient, protein expressing allele.

When assessed as adult mice (>8 weeks old), both the $Adar1^{P195A/-}$ and $Adar1^{P195A/E861A}$ were macroscopically normal. To determine if there were any microscopic changes, a histopathological assessment of a cohort of $Adar1^{+/+}$ (wild-type littermates; n=3), $Adar1^{P195A/+}$ (n=3) and $Adar1^{P195A/E861A}$ (n=4) at 6-7 months of age was undertaken as we have previously described (Chalk et al., 2019; Heraud-Farlow et al., 2017). From this analysis there were no genotype specific differences between the groups, nor evidence for pathological changes within the tissues assessed (Supplemental Dataset 1). We also assessed the brain, kidney, liver and spleen from a cohort of 13-19 week old animals containing C57BL/6 mice (n=3; bred and housed in the same facility), $Adar1^{P195A/+}$ (n=3), $Adar1^{P195A/-}$ (n=4), and $Adar1^{P195A/Ifh1/-}$ (n=3; described below, Supplemental Dataset 2). This did not find any genotype specific defects or pathological changes.

**Loss of MDA5 is sufficient to prevent innate immune activation.**

Based on the demonstrated role of MDA5 in sensing and responding to unedited cellular dsRNA (Liddicoat et al., 2015), we sought to determine if loss of MDA5 (gene $Ifh1$) would prevent the lethality we had observed in the $Adar1^{P195A/-}$ in the first 100 days of life. We generated cohorts of animals on both an $Ifh1^{+/+}$ or $Ifh1^{-/-}$ background. Strikingly, both heterozygosity for MDA5 or full deletion was able to prevent the death we had observed in a subset of $Adar1^{P195A/-}$ animals on an $Ifh1^{-/-}$ background (Fig 3A-3B, Supplemental Fig 2A-B). Furthermore, heterozygous or homozygous loss of MDA5 normalised the weaning weights across all genotypes assessed. This was especially the case for the $Adar1^{P195A/-}$ where the $Adar1^{P195A/Ifh1^{-/-}}$ weights were less variable than on an $Ifh1^{+/+}$ background ($Adar1^{P195A/-}$ mean weight = 8.12g, n=27; $Adar1^{P195A/Ifh1^{-/-}}$ mean weight = 10.63g, n=16; Fig 3A-3B, Supplemental Fig 2A-B). The $Adar1^{P195A/Ifh1^{-/-}}$ and $Adar1^{P195A/Ifh1^{-/-}}$ weights were not significantly different from any other cohort on an MDA5 heterozygous or null background. (Fig 3A-3B, Supplemental Fig 2A-B). This demonstrates that loss of MDA5 alone, even as a
heterozygous mutation, is sufficient to restore viability and long-term survival of the $\text{Adar1}^{P195A/-}$ and $\text{Adar1}^{P195A/E861A}$ animals.

**Compound $\text{Adar1}^{P195A/-}$ and $\text{Adar1}^{P195A/E861A}$ have a mild activation of ISGs but not of ISRs**

We next sought to determine if the P195A mutation, which is specific to the cytoplasmic p150 isoform of ADAR1, altered the cellular innate immune response. We isolated RNA from brain, liver and kidney tissues from 13-19 week old C57BL/6 mice (n=3; bred and housed in the same facility; same cohort subjected to histopathology), $\text{Adar1}^{P195A/-}$ (n=3), $\text{Adar1}^{P195A/-}$ (n=4), $\text{Adar1}^{P195A/E861A}$ (n=3), $\text{Adar1}^{P195A/-/Ifih1-/-}$ (n=3) and $\text{Adar1}^{P195A/-/Ifih1-/-}$ (n=3) and used the same Taqman based qPCR assays as described in the original P195A mouse model description (Maurano et al., 2021) to assess the expression of the ISGs $\text{Oas1a}$, $\text{Ifi27}$ and $\text{Irf7}$ (Fig 4). We found that the $\text{Adar1}^{P195A/-}$ and the $\text{Adar1}^{P195A/E861A}$ had mildly elevated expression of all three ISGs with variability in the level of increased expression between individuals and tissues (Fig 4), consistent with that reported by Maurano and colleagues (Maurano et al., 2021) and in other Zα domain mutants (De Reuver et al., 2021; Nakahama et al., 2021; Tang et al., 2021). The induction of the ISGs was completely prevented by the deletion of MDA5 (Fig 4), consistent with the known function of MDA5 as the primary sensor of under-edited/unedited cellular dsRNA (Liddicoat et al., 2015; Mannion et al., 2014; Pestal et al., 2015). Based on the proposed activation of the integrated stress response (ISR) in the previously reported P195A model (Maurano et al., 2021), we assessed the expression of the ISR genes $\text{Asns}$, $\text{Cdkn1a}$, $\text{Hmox1}$ using Taqman based qPCR on the same samples assessed for ISGs. In contrast to that reported (Maurano et al., 2021), we do not find evidence for an increase or activation of the ISR as determined from the expression of these genes across the genotypes tested (Fig 4). In summary, the compound mutation of P195A with either the null allele ($\text{Adar1}^{P195A/-}$) or the editing dead allele ($\text{Adar1}^{P195A/E861A}$) leads to the activation of an MDA5 dependent ISG response. We do not find evidence activation of the integrated stress response and see complete genetic rescue by the loss of MDA5 in vivo.

**Acute expression of ADAR1 P195A alone is well tolerated in vitro**

To assess the consequence of somatic restricted P195A expression, we established HOXA9 immortalised myeloid cell lines (Wang et al., 2006) using bone marrow from adult (>8 week old) $\text{R26-CreER}^{T2} \text{Adar1}^{+/+}$ and $\text{R26-CreER}^{T2} \text{Adar1}^{BP195A}$ animals (Fig 5A). These cell lines become ADAR1 heterozygous ($fl/+)$ or ADAR1 P195A only expressing ($fl/P195A$ becomes $\mathcal{N}P195A$) after treatment with tamoxifen (Fig 5B). Notably, these cells employ the same floxed $\text{Adar1}$ exon 7-9 allele used by Maurano and colleagues. We used this model to assess the consequences of acute deletion of the floxed $\text{Adar1}$ allele and expression of only
P195A. We cultured the cells with and without tamoxifen over a 14-day time course. We achieved complete deletion of the Adar1 floxed allele as assessed by genotyping (Fig 5C). The proliferation and viability of the P195A cells was not affected by treatment with tamoxifen and the cells behaved in a manner comparable to the control cells (Fig 5D-5E). We further assessed if there were changes in ISG expression as cells transitioned to being Adar1^−/P195A. We did not see any change in expression of the ISGs Ifit1 or Irf7 by qPCR, indicating that there was not an activation of the innate immune sensing system in these cells (Fig 5F). We further challenged the Adar1^−/P195A cells with transfection with high molecular weight polyinosinic-polycytidylic acid (polyI:C), a synthetic dsRNA, to determine if the P195A mutation altered the response to non-cellular dsRNA. The Adar1^−/P195A cells did not have a different response to a dose range of polyI:C compared to control cells when assessing induction of ISGs (Fig 5G). Therefore, at least in an immortalized myeloid cell, the expression of P195A alone is not sufficient to spontaneously activate an innate immune response to cellular dsRNA nor induce a different response to control cells to exogenous dsRNA. As we see activated ISGs in vivo, this result suggests that different cell types may respond differently to the P195A mutation as was reported for a biochemical ADAR1 Zα mutant model (Tang et al., 2021).

In vivo expression of P195A alone activates an ISG signature but is well tolerated

To directly test if there was a difference between the in vitro and in vivo response to the P195A mutation, we completed an in vivo tamoxifen treatment of the R26-CreER^T2 Adar1^fl/+ and R26-CreER^T2 Adar1^fl/P195A animals as we previously described (Heraud-Farlow et al., 2017). Adult (>8 week old) animals were fed tamoxifen containing diet for 4 weeks (Fig 6A). Analysis of the genomic DNA derived from bone marrow cells at day 28 demonstrated efficient and comparable recombination of the Adar1 floxed allele (Fig 6B). We did not see evidence for selection against efficient deletion, evidenced by retention of the floxed allele, as we had previously reported with either the R26-CreER^T2 Adar1^fl/+ or R26-CreER^T2 Adar1^fl/E861A in the same experimental model (Heraud-Farlow et al., 2017). All animals tolerated the diet well, with no significant difference in weight change between the cohorts compared to the weight at Day 0 (Fig 6C-6D).

At day 28 of tamoxifen diet, we collected peripheral blood (Fig 6E), bone marrow (Fig 6F-6H), spleen (Fig 6I-6J) and thymus (Fig 6K-6L) and assessed haematopoiesis. We did not see any substantive changes across these organs in terms of cellularity or lineage distribution/differentiation. We also assessed activation of the ISG response by assessing the expression of Sca-1, a known ISG induced in response to a loss of ADAR1, and expression of Ifit1 and Irf7 (Essers et al., 2009; Hartner et al., 2009; Heraud-Farlow et al., 2017) (Fig 6N-
We saw increased expression of Sca-1 on the cell surface of the Adar1^P195A within the lineage negative fraction of the bone marrow. Prompted by the increased Sca-1 expression, qPCR for the ISGs Ifit1 and Irf7 demonstrated a 6-8 fold increased expression of these ISGs in the Adar1^P195A animals bone marrow (Fig 6N-6M). Collectively, these analyses demonstrate that in vivo Adar1^P195A animals, but not in vitro myeloid cells, have an activation of an innate immune response following somatic restricted expression of P195A. This ISG activation is of a relatively low level, compared to the levels seen when animals and cells are engineered to be ADAR1 null or editing deficient (Heraud-Farlow et al., 2017; Liddicoat et al., 2016b; Liddicoat et al., 2015), and is well tolerated without any effects on the animals overall well-being or survival.
Discussion

Understanding how disease associated ADAR1 mutations affect the proteins’ function will be critical to the ultimate goal of developing effective treatments for patients with mutations in ADAR. To this end, the development of preclinical models that mirror human genetics is an important step. The P193A mutation, which specifically impacts the cytoplasmic ADAR1p150 isoform, is the most commonly reported ADAR mutations in humans with AGS and BSN (Livingston et al., 2014; Rice et al., 2012; Rice et al., 2017). In AGS, the P193A mutation is reported as a compound heterozygous mutation with a second mutation, most often one that either compromises expression of the second allele or is predicted to compromise A-to-I editing by the protein product of the second allele. Intriguingly, the P193A mutation is also present in the general human population. As this is the most common human ADAR1 mutation it will be important to understand its effect on both ADAR1’s canonical function in A-to-I RNA editing and in other protein dependent functions of ADAR1.

Herein we describe the independent generation and phenotyping of a murine model of the human P193A mutation. The homologous murine mutation, P195A, is well tolerated and compatible with adult homeostasis when either heterozygous or homozygous. This is consistent with that reported using independently generated P195A knock-in alleles by both Maurano et al., (Maurano et al., 2021) and in the work of Guo et al., (co-submitted). Whilst homozygosity for P193A is a rare occurrence in the general human population (Karczewski et al., 2020) and has not been reported in AGS patients (Rice et al., 2012; Rice et al., 2017), all groups have established viable and ostensibly normal Adar1<sup>P195A/P195A</sup> mice. The present data, together with the observations from the independent murine P195A alleles, demonstrate that this mutation is well tolerated and does not significantly compromise ADAR1p150 function in vivo under normal laboratory conditions. This proline is not conserved in the Zα family. It is present in the domain wing and affects the kinetics of binding to left-handed nucleic acids that differs between species (Subramani et al., 2016). Interestingly, Zα mutation to highly conserved residues involved in binding Z-DNA (N175A/Y179A) (de Reuver et al., 2021; Tang et al., 2021) was well tolerated in vivo when homozygous, while the W197A mutant (W195 in human), essential to stabilizing the wing, was reported to have a more severe in vivo phenotype (Nakahama et al., 2021). Collectively, these data indicate that specific mutations within the Zα domain of ADAR1p150, such as the W197A, reveal specific functions for Zα in vivo and this warrants further detailed understanding.

The most striking difference between our findings and those reported by Maurano et al., (Maurano et al., 2021) are apparent once the P195A allele was crossed to an Adar1<sup>+/−</sup> allele.
The Adar1^{P195A/-} genotype more closely approximates that reported in AGS patients, where the P193A allele is reported with a second mutation predicted to be deleterious (Rice et al., 2017). Maurano et al., reported that the majority of their Adar1^{P195A/-} mice were significantly runted and die by 30 days of age, with a single animal surviving to 84 days of age (Maurano et al., 2021). In contrast, we see long term (>250 days) survival of the majority of the compound mutant Adar1^{P195A/-} mice. The basis for this profound difference is not immediately apparent but is important to understand. Whilst differences in vivariums may contribute, it seems unlikely that this is the primary driver of the differences. We believe that two factors may be significant contributors to the differences. Firstly, we observe significant differences in survival and weights of the Adar1^{P195A/-} mice based on the breeding pair genotype (Fig 3). At present we have results from the breeding of two female and one male Adar1^{P195A/-} mice crossed to Adar1^{P195A/+} mice. When an Adar1^{P195A/-} was used for breeding we see significantly reduced weaning weights and poor survival of the Adar1^{P195A/-} pups, not dissimilar to that reported by Maurano et al., (Maurano et al., 2021). It should be noted that the Adar1^{P195A/-} genotype is most similar to that of an AGS patient, and to the best of our knowledge there is no available information as to the breeding of these human genotypes. Maurano et al., also tested the effects of the P195A mutation and concurrent deletion of the other ADAR1p150 isoform (Adar1^{P195A/p150-}) which we have not undertaken (Maurano et al., 2021). We did, however, test the effect of the combination of the P195A mutation with the editing dead E861A mutation (Liddicoat et al., 2015). The Adar1^{P195A/E861A} did not have any early lethality and had more consistent weaning weights than the Adar1^{P195A/-} mice in our cohorts. This demonstrates that A-to-I editing by the non P195A p150 protein is not essential to suppress the observed phenotypes but is required to prevent MDA5 activation, based on ISG expression. Therefore, runting and early lethality of Adar1^{P195A/-} mice is due to a protein-dependent editing-independent function of ADAR1p150.

A more significant contributor to the dissimilarities is likely to be the different Adar1 deficient alleles used in the respective studies. Maurano et al, utilised an Adar1 deficient allele derived by the germ-line deletion of the conditional Adar1^{fl} allele that deletes exon 7 to 9 (Hartner et al., 2004; Hartner et al., 2009; Maurano et al., 2021; Pestal et al., 2015). This allele has recently been demonstrated to yield a truncated, editing deficient and mislocalized protein product impacting both p110 and p150 isoforms (Bajad et al., 2020). It is possible that the defective ADAR protein from the exon 7-9 deleted allele could stimulate the integrated stress response reported, as misfolded proteins are known to activate this pathway (Subramani et al., 2016). We have used this same genotype in both the immortalised myeloid cells (Fig 4) and somatic deletion models (Fig 5) and in both acute deletion settings we see tolerance of this genotype. Alternatively, the truncated p110 and p150 proteins, whilst non-functional for A-
to-I editing, harbour the potential to interfere with the activity of the P195A p150 protein in the cytoplasm and the wild-type p110 expressed from the P195A allele through substrate competition or dimerization (Cho et al., 2003; Valente and Nishikura, 2007). This could potentially result in a less functional, effectively hypomorphic, P195A p150 and WT p110 when heterozygous. In our study we have used an exon 2-13 deletion of Adar1, demonstrated to be a null allele and not known to express any protein product (Hartner et al., 2004). In a complimentary independent report by Guo et al., (co-submitted), they used an independently generated P195A knock-in and an exon 12-15 deletion allele (Wang et al., 2004), also demonstrated to be a protein null, and they have also seen long term survival of the Adar1<sup>P195A−/−</sup> mice they generated with even better survival rates than we observed. The data from these two independently generated P195A alleles, crossed to two separate Adar1 null alleles, strongly argue that the runting and completely penetrant post-natal lethality of the Adar1<sup>P195A−/−</sup> mice reported by Maurano et al., is most like confounded by the effects of the Adar1<sup>ΔEx7-9</sup> allele (Maurano et al., 2021). When crossed to true null alleles, the majority of Adar1<sup>P195A−/−</sup> mice survive with no pathological differences in adult animals compared to controls. There may also be other strain differences such as those immune deficiencies previously reported in some lines of C57BL/6 mice (Koehler et al., 2020) or accounting for the high birth weight in the Maurano et al. study. The ultimate cause of death of the runted Adar1<sup>P195A−/−</sup> mice from our cohorts remains to be determined. Moreover, the P195A mutation can be complemented by the provision of the editing deficient E861A allele. This demonstrates that provision of a full length, editing deficient p150 protein is sufficient to prevent the early lethality and runting we see in a subset of our Adar1<sup>P195A−/−</sup> cohort that depends on parental genotype. In summary the P195A mutation, homologous to the most common human ADAR1<sup>E861A/E861A</sup> mice, is well tolerated in vivo and the loss of MDA5 is sufficient to completely rescue the Adar1<sup>P195A−/−</sup> mice.

The physiologically most important role of A-to-I editing by ADAR1 is to edit cellular/endogenous dsRNA to prevent MDA5 mediated innate immune sensing (Liddicoat et al., 2016a; Liddicoat et al., 2015). This has been demonstrated in both mouse models and human cells (Liddicoat et al., 2015; Mannion et al., 2014; Pestal et al., 2015). The P195A mutation is associated with reduced editing of some substrates in cellulo (Maurano et al., 2021). In both human and mouse, the loss of MDA5-MAVS prevents innate immune activation following the loss of ADAR1 or loss of A-to-I editing by ADAR1. Previous work has not demonstrated rescue from embryonic lethality of Adar1<sup>−/−</sup> animals by concurrent loss of PKR (Eif2ak2<sup>−/−</sup>) (Wang et al., 2004) or by loss of IFNAR (Liddicoat et al., 2016b; Mannion et al., 2014), unlike the rescue to birth or adulthood by loss of MDA5 (Ifih1<sup>−/−</sup>) or MAVS (Mavs<sup>−/−</sup>) of the Adar1<sup>−/−</sup> or ADAR1 editing deficient (Adar1<sup>E861A/E861A</sup>) mice, respectively (Heraud-Farlow et
Based on the ISR signature and genetic crosses it was proposed that PKR activation was important in the pathology of the P195A mouse models generated by Maurano et al., and this was genetically tested (Maurano et al., 2021). We see a modest activation of the downstream transcriptional markers of MDA5 signalling (interferon stimulated genes) in tissues isolated from adult Adar1<sup>P195A/-</sup> and Adar1<sup>P195A/E861A</sup> mice. Importantly, this can be completely prevented by loss of MDA5, and interestingly there is a dosage dependent effect of MDA5 with a degree of genetic rescue afforded by being Ifih1<sup>+/−</sup> (Supplemental Fig 2B). We see complete rescue of the Adar1<sup>P195A/-</sup> and Adar1<sup>P195A/E861A</sup> mice by loss of MDA5 and a full suppression of ISG expression to baseline levels in tissues, consistent with a model where MDA5 is the primary, and initiating, sensor of unedited self dsRNA. We do not see evidence in the adult mouse tissues of transcriptional changes consistent with activation of PKR, as was reported by Maurano et al., in the Adar1<sup>P195A/p150</sup>- animals (Maurano et al., 2021). All of the ADAR1 Zα mutants described to date demonstrate complete rescue by loss of either MDA5 (Maurano et al., 2021; Nakahama et al., 2021) or MAVS (de Reuver et al., 2021; Tang et al., 2021). Collectively, these results demonstrate that the absence of MDA5, and subsequent innate immune response, is sufficient to rescue the effects of ADAR1 P195A Zα mutants. In summary the P195A mutation, homologous to the most common human ADAR1 mutation reported in AGS, is well tolerated <i>in vivo</i> and the loss of MDA5 is sufficient to completely rescue the Adar1<sup>P195A/-</sup> mice. These findings are consistent with a role for p150 in down-regulating the ISG response but do not explain the pathology observed in AGS patients.
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Author Contribution Statement

J.H-F and C.R.W conceptualized the study. Z.L, J.H-F and C.R.W designed the experiments. Z.L, S.T, A.G, J.H-F and C.R.W performed the experiments. Z.L, J.H-F and C.R.W wrote the original manuscript, and all authors reviewed and edited the manuscript; J.H-F and C.R.W were responsible for funding acquisition. J.H.F and C.R.W provided supervision.

Declaration of Interests

All authors declare no competing financial interests
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**Figure Legends**

**Figure 1. Generation of an Adar1<sup>P195A</sup> knock-in allele.**

A) Schematic of the P195A mutation.

B) Genomic alignment and translation of the WT and P195A allele. During introduction of the P195A mutation, two additional silent mutations were introduced immediately upstream and downstream, respectively. These were to allow restriction digestion of the mutant allele and to prevent re-cutting by CAS9 during targeting.

C) Sanger sequencing traces and alignments of genomic DNA isolated from animals of the indicated genotypes.

D) Western blot analysis of ADAR1 expression in E13.5 mouse embryonic fibroblasts (MEFs) of the indicated genotypes +/- interferon-β (IFNβ) treatment. IFNβ was used to induce expression of the p150 isoform of ADAR1.

**Figure 2. Long term survival of Adar1<sup>P195A/−</sup> animals.**

A) Results from inbreeding of Adar1<sup>P195A/+</sup> animals.

B) Results from breeding of Adar1<sup>P195A/+</sup> animals with Adar1<sup>P195A/−</sup> animals.

C) Weaning weights of mice of the indicated genotypes. *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001; Ordinary one-way ANOVA with Tukey’s multiple comparison test (adjusted P value).

D) Survival analysis of mice of the indicated genotypes; number as indicated for each genotype. Statistical difference determined by Log-rank (Mantel-Cox) test.

E) Weaning weights and F) survival analysis of pups derived from breeding of a Adar1<sup>P195A/P195A</sup> to a Adar1<sup>+/−</sup> (Adar1 germ-line deficient heterozygous) animal; number as indicated for each genotype. Statistical analysis of weight using unpaired t test (two sided); of survival using Log-rank (Mantel-Cox) test.

G) Weaning weights and H) survival analysis of pups derived from breeding of a Adar1<sup>P195A/−</sup> to a Adar1<sup>P195A/+</sup> animal; number as indicated for each genotype. Inset photo – 35 day old Adar1<sup>P195A/−</sup> male bred from Adar1<sup>P195A/P195A</sup> x Adar1<sup>+/−</sup> parents (data in Panel E/F); sibling 29 day old Adar1<sup>P195A/+</sup> and Adar1<sup>P195A/−</sup> male bred from an Adar1<sup>P195A/−</sup> x Adar1<sup>P195A/+</sup> (data in Panel G/H). ****P<0.0001; for weight analysis Ordinary one-way ANOVA with Tukey’s multiple comparison test (adjusted P value); survival using Log-rank (Mantel-Cox) test.

Data displayed as individual animals with mean indicated. Weaning weights not available if animal was found dead prior to weaning; all animals where genotype was confirmed are included in the survival analysis (where possible any found dead prior to weaning were genotyped post-mortem).

**Figure 3. Loss of MDA5 rescues both weight and viability of Adar1<sup>P195A/−</sup>.**
A) Weaning weights of mice of the indicated genotypes on an MDA5 heterozygous (Ifih1\(^{+/−}\)) or null (Ifih1\(^{−/−}\)) background. No statistically significant difference across any comparison (Ordinary one-way ANOVA with Tukey’s multiple comparison test).

B) Survival analysis of mice of the indicated genotypes (all Ifih1\(^{−/−}\)); number as indicated for each genotype. Survival analysis for all Ifih1\(^{+/−}\) and Ifih1\(^{−/−}\) genotypes in Supplemental Fig 2B. No statistical difference between genotypes by Log-rank (Mantel-Cox) test.

Data displayed as individual animals with mean indicated. Weaning age was determined by animal facility staff independently of investigators based on animal welfare and facility SOPs. Animals typically weaned at 20-25 days of age; number as indicated for each genotype. Weaning weights not available if animal was found dead prior to weaning; all animals where genotype was confirmed are included in the survival analysis (where possible any found dead prior to weaning were genotyped post-mortem).

**Figure 4.** Mild tissue-specific upregulation of interferon stimulated gene expression in Adar1\(^{P195A/−}\) and Adar1\(^{P195A/E861A}\) is MDA5 dependent.

A) Taqman based qPCR for the interferon stimulated genes Oas1a, Ifi27 and Irf7 in adult brain, liver and kidney of mice of the indicated genotypes; number of independently derived samples as indicated for each genotype.

B) Taqman based qPCR for the integrated stress response genes Asns, Cdkn1a and Hmox1 in adult brain, liver and kidney of mice of the indicated genotypes; number of independently derived samples as indicated for each genotype.

Data expressed as mean +/- SEM gene expression relative to Hprt expression. N=3 per genotype; Statistical analysis for all qPCR: Ordinary one-way ANOVA with Dunnett’s multiple comparison test; *P<0.05; **P<0.01 or P as indicated. No P value indicates not statistical significance.

**Figure 5.** Expression of P195A is well tolerated in myeloid cells.

A) Schematic outline of how the HOXA9 immortalised myeloid cell lines are derived.

B) Experimental outline. When tamoxifen is added to the cells this activates deletion of the floxed Adar1 allele leaving the cells either heterozygous (Controls; \(Δ/+\)) or P195A only expressing (\(Δ/P195A\)).

C) Genomic DNA genotyping demonstrating efficient recombination of the floxed Adar1 allele following tamoxifen treatment.

D) Proliferation of cell lines of the indicated genotypes with and without tamoxifen treatment (isogenic pairs). Cells were counted on a Countess II cell counter.
E) Cell viability of cell lines of the indicated genotypes with and without tamoxifen treatment (isogenic pairs) assessed by trypan blue staining and on a Countess II cell counter.

F) qPCR (SYBR green) based analysis of *Ifit1* and *Irf7* expression in the cell lines collected at day 14 of analysis. Data expressed as mean +/- SEM gene expression relative to *Ppia* expression.

G) qPCR (SYBR green) based analysis of *Ifit1*, *Irf7* and *Ifnb* expression in the cell lines transfected with the indicated dose of high molecular weight Polyinosinic-polycytidylic acid (polyI:C). Data expressed as mean +/- SEM gene expression relative to *Ppia* expression. Statistical analysis by t-test using individually calculated AUC per sample. Cell lines had been treated for 14 days with tamoxifen, tamoxifen withdrawn, genotyped, then used for polyI:C dose response.

Data from three independently derived cell lines (different donor bone marrow) for each genotype.

**Figure 6. In vivo expression of P195A is well tolerated and results in a modest induction of interferon regulated gene expression.**

A) Schematic outline of experiment.

B) Representative genotyping of recombination of the *Adar1* floxed allele at day 28 using genomic DNA isolated from whole bone marrow cells. Recombination percentage was calculated using LabChip (PerkinElmer) based quantitation of band intensity compared to the WT/P195A allele and known standard/marker.

C) Survival analysis of mice of the indicated genotypes; number as indicated for each genotype. No statistical difference between genotypes by Log-rank (Mantel-Cox) test using Prism.

D) Percentage change in body weight of each cohort based on comparison of the weight at day 28 of tamoxifen food compared to day 0 (prior to initiation of tamoxifen containing diet).

E) Peripheral blood leukocyte populations (by lineage) between genotypes at day 0 and day 28.

F) Bone marrow cellularity at day 28 between genotypes.

G) Differential analysis of nucleated cell populations in the bone marrow at day 28.

H) Erythroid cell populations in the bone marrow at day 28.

I) Splenic cellularity at day 28 between genotypes.

J) Differential analysis of nucleated cell populations in the spleen at day 28.

K) Thymic cellularity at day 28 between genotypes.

L) Differential analysis of thymocyte populations in the thymus at day 28.

M) qPCR (SYBR green) based analysis of *Ifit1* and *Irf7* expression in whole bone marrow at day 28. Data expressed as mean +/- SEM gene expression related to *Ppia* expression.
N) Representative flow cytometry histogram of Sca-1 expression between a control (Δ/++; grey) and P195A only (Δ/P195A; blue) expressing bone marrow sample. Quantitation of Sca-1 mean fluorescence intensity within the lineage negative fraction of whole bone marrow. Unless otherwise stated data expressed as mean +/- SEM; n=3 R26-CreER<sup>ki/+</sup> Adar1<sup>F/+</sup> (control) and n=4 R26-CreER<sup>ki/+</sup> Adar1<sup>F>P195A</sup> (test).
Supplemental Information:

Figures:

Supplemental Figure 1. Schematic outline of the different alleles used in this study and weaning weights and survival of pups from Adar1\(^{P195A/+}\) inbreeding.

A) Outline of the different alleles used and expected consequences on ADAR1 p110 and p150 expression/ function.

B) Weaning weights and survival analysis of pups derived from inbreeding of Adar1\(^{P195A/+}\) animals; number as indicated for each genotype. For weights: Ordinary one-way ANOVA with Tukey’s multiple comparison test (adjusted P value); survival using Log-rank (Mantel-Cox) test - no significant difference when tested. Data displayed as individual animals with mean indicated. Weaning weights not available if animal was found dead prior to weaning; all animals where genotype was confirmed are included in the survival analysis (where possible any found dead prior to weaning were genotyped post-mortem).

Supplementary Figure 2. Loss of MDA5 prevents Adar1\(^{P195A/-}\) lethality.

A) Comparison of survival and weaning weights of Adar1\(^{P195A/+}\) and Adar1\(^{P195A/-}\) Ifih1\(^{-/-}\) (datasets replotted from Fig 2C-2F). For survival analysis: *P<0.05 (Log-rank (Mantel-Cox) test); For weaning weights: *P<0.05; **P<0.01 (One-way ANOVA with Tukey’s multiple comparison test; adjusted P Value).

B) Comparison of survival of all genotypes on an MDA5 heterozygous (Ifih1\(^{+/+}\)) and homozygous null (Ifih1\(^{-/-}\)) background; complete dataset for cohorts presented in Figure 2E.

Tables:

Supplementary Table 1. Average weights of C57Bl/6 background mice.

A) Data from The Jackson Laboratory.

B) Data from the International Mouse Phenotyping Consortium

Datasets:

Supplementary Dataset 1: Full histopathology report of adult Adar1\(^{P195A/E861A}\) and control mice.

Supplementary Dataset 2: Full histopathology report of brain, kidney and liver samples from adult Adar1\(^{P195A/-}\) and control mice.
**Figure 1** Liang et al. CW250522

A. 

WT

<table>
<thead>
<tr>
<th>ADAR1 p150</th>
<th>ADAR1 p110</th>
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<td>Deaminase domain</td>
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P195A mutation

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B. 

Genomic DNA sequence

WT

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Protein sequence

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C. 

WT

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P195A+/+

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<td>Protein</td>
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P195A/+

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D. 

IFNβ

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<th>P195A/195A</th>
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ADAR1

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<tr>
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ACTIN

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<tr>
<td>+</td>
<td>-</td>
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ADAR1 (long exp.)

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<th>-/-</th>
<th>+/+</th>
<th>-/+</th>
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<tbody>
<tr>
<td>p150</td>
<td>p110</td>
<td>p150</td>
<td>p110</td>
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A. **Adar1P195A+/+ x Adar1P195A+/+**

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<tr>
<th></th>
<th>Expected (%)</th>
<th>Expected (n)</th>
<th>Actual (n)</th>
<th>Actual (%)</th>
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<tbody>
<tr>
<td>Adar1+/+</td>
<td>25%</td>
<td>26.25</td>
<td>28</td>
<td>26.7%</td>
</tr>
<tr>
<td>Adar1P195A/+</td>
<td>50%</td>
<td>52.5</td>
<td>59</td>
<td>56.2%</td>
</tr>
<tr>
<td>Adar1P195A/P195A</td>
<td>25%</td>
<td>26.25</td>
<td>18</td>
<td>17.0%</td>
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<tr>
<td><strong>Total</strong></td>
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Chi-square P=0.1725

B. **Adar1P195A/- x Adar1P195A/+**

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<th>Expected (%)</th>
<th>Expected (n)</th>
<th>Actual (n)</th>
<th>Actual (%)</th>
</tr>
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<tbody>
<tr>
<td>Adar1+/+</td>
<td>25%</td>
<td>11.75</td>
<td>10</td>
<td>21.3%</td>
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<tr>
<td>Adar1P195A/+</td>
<td>25%</td>
<td>11.75</td>
<td>14</td>
<td>29.8%</td>
</tr>
<tr>
<td>Adar1P195A/P195A</td>
<td>25%</td>
<td>11.75</td>
<td>14</td>
<td>29.8%</td>
</tr>
<tr>
<td><strong>Total</strong></td>
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<td>47</td>
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</table>

Chi-square P=0.6224

E. **Weaning Weights (g)**

- **P195A/E861A**
- **P195A/-**
- **P195A/P195A**
- **P195A/+**
- **E861A/+**
- **Adar1+/+**
- **WT**

*P=0.049

F. **Probability of Survival**

- **P195A/+**
- **P195A/-**
- **P195A/P195A**
- **P195A/E861A**

P=0.066

H. **Weaning Weights (g)**

- **Adar1+/+**
- **P195A/+**
- **P195A/-**

*P=0.0001

Chi-square P=0.6224

Figure 2 Liang et al_CW240622
A. 

<table>
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<tr>
<th>Genotype</th>
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<tr>
<td>P195A/E861A Ifih1-/-</td>
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<tr>
<td>P195A/E861A Ifih1+/-</td>
<td>n=6</td>
</tr>
<tr>
<td>P195A/ Ifih1-/-</td>
<td>n=16</td>
</tr>
<tr>
<td>P195A/ Ifih1+/-</td>
<td>n=8</td>
</tr>
<tr>
<td>P195A/P195A Ifih1-/-</td>
<td>n=12</td>
</tr>
<tr>
<td>P195A/P195A Ifih1+/-</td>
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</tr>
<tr>
<td>P195A/ Ifih1+/-</td>
<td>n=22</td>
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<tr>
<td>P195A/ Ifih1+/-</td>
<td>n=19</td>
</tr>
<tr>
<td>Adar1+/+ Ifih1-/-</td>
<td>n=11</td>
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</tbody>
</table>

B. 

Weaning Weights (g) vs. Days elapsed

Weaning Weight Graph:

- P195A/+ Ifih1-/- (n=15)
- P195A/ Ifih1+/- (n=1)
- P195A/P195A Ifih1-/- (n=12)
- P195A/E861A Ifih1-/- (n=16)

Probability of Survival Graph:

- P195A/+ Ifih1-/- (n=15)
- P195A/ Ifih1+/- (n=11)
- P195A/P195A Ifih1-/- (n=12)
- P195A/E861A Ifih1-/- (n=16)
A.

**Brain**

- **Oas1a**
- **Ifi27**
- **Irf7**

**Liver**

- Normalised expression (to Hprt)
- P=0.08

**Kidney**

- Normalised expression (to Hprt)

B.

- **Asns**
- **Cdkn1a**
- **Hmox1**
GM-CSF

HOXA9 retrovirus becomes Adar1\(^{\Delta/+}\) - ADAR1 protein expressed

Adar1\(^{\Delta/\text{P195A}}\) - only express ADAR1 P195A

Tamoxifen

Days + Tamoxifen

HOXA9 immortalised myeloid cells

R26-CreER Adar1\(^{\text{fl/+}}\) (Ctrl)

R26-CreER Adar1\(^{\text{fl/P195A}}\)

Cell Viability (%)

Days + Tamoxifen

Cumulative cell number

fl/+-Tam

fl/+ +Tam

fl/P195A-Tam

fl/P195A

fl/+ -Tam

fl/+ +Tam

fl/P195A-Tam

fl/P195A + Tam

Ifit1: Ppia mRNA

Irf7: Ppia mRNA

Ifnb: Ppia mRNA

Hmw polyI:C μg/mL

mock

Δ/+ vs Δ/P195A

P=0.34

Δ/+ vs Δ/P195A

P=0.0015
A. R26-CreER $Adar^{f1}$ (Ctrl)
B. Genotype
C. Probability of Survival (%)
D. Day 28 body weight (% of Day 0)
E. PB Leukocytes (x10^6)
F. Cells/Femur (x10^6)
G. Cells/Femur (x10^6)
H. Cells/Femur (x10^6)
I. Cells/Spleen (x10^6)
J. Cells/Spleen (x10^6)
K. Cells/Thymus (x10^6)
L. Cells/Thymus (x10^6)
M. Irf7: Ppia mRNA
N. Sca-1 Mean Fluorescence Intensity
Supplemental Figure 1. Schematic outline of the different alleles used in this study and weaning weights and survival of pups from Adar1\(^{P195A/+}\) x \(P195A/\) inbreeding.

**A.** Outline of the different alleles used and expected consequences on ADAR1 p110 and p150 expression/function.

**B.** Weaning weights and survival analysis of pups derived from inbreeding of Adar1\(^{P195A/+}\) animals; number as indicated for each genotype. For weights: Ordinary one-way ANOVA with Tukey’s multiple comparison test (adjusted P value); survival using Log-rank (Mantel-Cox) test - no significant difference when tested. Data displayed as individual animals with mean +/- sem indicated. Weaning weights not available if animal was found dead prior to weaning; all animals where genotype was confirmed are included in the survival analysis (where possible any found dead prior to weaning were genotyped post-mortem).
Supplemental Figure 2. Loss of MDA5 prevents Adar1P195A/- lethality.
(A) Comparison of weaning weights and survival of Adar1P195A/- and Adar1P195A/- Ifih1/- (datasets replotted from Fig 2C-2F). For survival analysis: P=0.0007 (Log-rank (Mantel-Cox) test); For weaning weights: *P<0.05; **P<0.01 (One-way ANOVA with Tukey’s multiple comparison test; adjusted P Value).
(B) Comparison of survival of all genotypes on an MDA5 heterozygous (Ifih1+/−) and homozygous null (Ifih1−/−) background; complete dataset for cohorts presented in Figure 2E.

n as indicated in legends.
Supplementary Table 1. Average mouse weights by age for C57Bl/6 background mice.

A. The Jackson Laboratory Data for mouse weights by age

Data downloaded 24-March-2022
BODY WEIGHT INFORMATION FOR C57BL/6J (000664)
JAX® Mice Strain - C57BL/6J
Stock Number 000664

Male and female mice were weighed the same day each week in nine mouse rooms representing all three breeding facilities. Weights were measured from 40 males and 40 females in each room; the values shown here represent averages of up to 360 males and 360 females. Mice were fed a diet containing 6% fat (LabDiet 5K52 formulation). Values represent mean and one standard deviation. Ages are ± 3 days.

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<thead>
<tr>
<th>Age (Weeks)</th>
<th>Females</th>
<th>Males</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>10.1 ± 1.7</td>
<td>10.6 ± 1.9</td>
</tr>
<tr>
<td>4</td>
<td>14.7 ± 1.8</td>
<td>16.5 ± 2.6</td>
</tr>
<tr>
<td>5</td>
<td>17.8 ± 1.1</td>
<td>20.7 ± 1.8</td>
</tr>
<tr>
<td>6</td>
<td>18.5 ± 0.9</td>
<td>21.9 ± 1.8</td>
</tr>
</tbody>
</table>

B. International Mouse Phenotyping Consortium (www.mousephenotype.org)

Data downloaded 24-March-2022
Data obtained from Gene: Adar (allele Adar<sup>tm1b(EUCOMM)Wtsi</sup>) entry; MGI:1889575;
Phenotyping Centre: Jax; involves: C57BL/6NJ
Control cohort data (aggregate)
Age rounded to the nearest week

Body Weight (grams; st. dev range; n)

<table>
<thead>
<tr>
<th>Age (Weeks)</th>
<th>Females</th>
<th>Males</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>12.69; SD 11.06-14.34 (n=139)</td>
<td>13.74; SD 11.61-15.87 (n=140)</td>
</tr>
<tr>
<td>4</td>
<td>14.74; SD 12.91-16.57 (n=2612)</td>
<td>16.59; SD 13.94-19.24 (n=2621)</td>
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<tr>
<td>5</td>
<td>16.10; SD 14.37-17.82 (n=2374)</td>
<td>19.00; SD 16.62-21.37 (n=2376)</td>
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### 9.1 Histopathology Report

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<tr>
<th>Case Number</th>
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<th>Animal Details</th>
<th>Death</th>
<th>Origin</th>
<th>Treatment</th>
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<td>Wed 08/09/2021</td>
<td>#20, WT (Genotype: Adar1+/+)</td>
<td>CO2</td>
<td>St Vincent's Institute</td>
<td>Mutation</td>
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<tr>
<td></td>
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<td>DOB: 27/02/2021, 6 months old, Male, 32.0g, Black</td>
<td></td>
<td></td>
<td>ADAR1 mutations are associated with a paediatric leukodystrophy called</td>
</tr>
<tr>
<td></td>
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<td>Strain: Adar1P195A/E861A</td>
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<td>Strain: Adar1P195A/E861A</td>
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<td>DOB: 05/02/2021, 7 months old, Female, 23.3g, Black</td>
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<tr>
<td></td>
<td></td>
<td>Strain: Adar1P195A/E861A</td>
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<tr>
<td></td>
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<td>DOB: 05/02/2021, 7 months old, Female, 22.7g, Black</td>
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<td>Strain: Adar1P195A/E861A</td>
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Thu, 28 Oct 2021 01:07:44 AM
Aicardi Goutières’s Syndrome (AGS; AGS6 is the subcategory of AGS caused by ADAR1 mutations). These patients have a constitutively elevated interferon response (ISGs) causes by self-sensing of endogenous RNA. The major phenotypes in ADAR1 mutant humans are (from OMIM database): Clinical details included intracranial calcification and severe developmental delay in all, leukodystrophy in most, and markedly elevated cerebrospinal fluid (CSF) interferon-alpha in all in whom it was measured. Six of the 9 children presented between 9 and 18 months with acute onset of severe generalized dystonia in the context of a nonspecific febrile illness, such as ear infection, respiratory illness, or viral illness with rash. Clinical features included limb tremor and stiffening, rigidity, loss of previous motor and other developmental skills, and severe dystonia affecting all 4 limbs. These neurologic abnormalities developed and progressed rapidly within several weeks, and were refractory to pharmacologic management. Two additional patients had a slower course, with progressive dystonia and neurologic deterioration over several months.

Can also get chilblain type lesions on skin and feet in patients (but we haven't noticed any on our mice).

The mutations in these mice are:
ADAR1 P195A (homologue to human P193A) – the most common ADAR1 mutation in humans and undergoing positive selection in human. This is also the most common human mutation in AGS patients
ADAR1 E861A (homologue to human E912A) – this is a biochemical mutation which causes the specific loss of editing activity by the ADAR1 enzyme. Patients with AGS6 most often have a P195A mutation with a deaminase compromising mutation similar but not the same as E912A.

Only compound mutants would be expected to be pathologic/phenotypic. Within the cohort of mice sent there are WT (indicated); some single heterozygous mutants and some compound mutants.

Request for particular attention to the brain.

(C. Walkley 17/08/2021).

**Species / Breed / Strain**
Adar1P195A/E861A

**Animal Health Facility**
St. Vincent's Bioresources Centre
POSITIVE for Mouse Norovirus, Helicobacter spp, Helicobacter hepaticus, Chilomastix bettencourti, Entamoeba muris, Protozoa, Tritrichomonas muris

**Macropathology:**
1. Cardiac epicardial mineralisation, a background finding.
2. Unilateral or bilateral inflammation of the upper and/or lower eyelids.

**Organs Examined**
Adrenal glands, Bladder, Bone marrow, Brain, Cecum, Cervix, Clitoral gland, Colon, Duodenum, Epididymes, Eyes, Gall bladder, Harderian glands, Head, Heart, Hind leg (Long bone, Bone marrow, Synovial joint, Skeletal muscle), Ileum, Jejunum, Kidney, Liver, Lungs, Mammary tissue, Mesenteric lymph node, Ovaries, Oviducts, Pancreas, Penis, Preputial gland, Prostate glands, Salivary glands and Regional lymph nodes, Seminal vesicles, Skin, Spinal cord, Spleen, Sternum, Stomach, Tail, Testes, Thymus, Thyroids, Trachea, Uterus, Vagina

**Macroscopic Observations**
Date of transport to the APN: 9th September 2021
Courier/Transportation details: Jetpets REF# 1415798

Body Condition Scoring (BCS): on a scale of 1-5, animals scored a BCS of 3.
5: The mouse is obese, and bones cannot be felt at all
4: The mouse is well-fleshed, and bones are barely felt
3: The mouse is in optimal condition. The bones are palpable but not prominent
2: The mouse is thin, and bones are prominent
1: Muscle wasting is advanced; fat deposits are gone, and bones are very prominent.

At the time of necropsy, the animals appeared well nourished, well groomed, active/curious and healthy with normal movement and gait. There were no observable dermal lesions and no nasal/ocular discharges. The gastrointestinal tract contained ample ingesta and the thoracic and abdominal viscera showed no macroscopic abnormalities. Noted that animal #18 weighed significantly less than the other male mice in this cohort. Please refer to individual animals for more macroscopic details.

X-Rays: No discernible neoformations of the bone. No readily discernible thinning or atrophic lesions in the skeletal structures. Please see accompanying APN21/036SVI (C. Walkley) X-Ray Images document.

Blood report:
Blood results showed most readings within the normal mouse reference intervals.
Female mice showed no obvious differences between Adar1P195A/E861A mice and WT controls.
Notable findings in the male Adar1P195A/E861A mice:
#3:
1. Elevated neutrophils (%NEUT, AbsNeut)
2. Low lymphocytes (%LYM, AbsLymph)
3. Low monocytes (%MONO, AbsMono)
#18:
1. Low lymphocytes (%LYM, AbsLymph)
2. Elevated eosinophils (%EOS, AbsEos)
Note: haematology values can vary with mouse strain/stock, age, sex, blood sampling method, fasting and environmental conditions, pathogen status, and laboratory. The reference intervals used for this report are based on published values of adult mice (16 weeks or older).
For more details please see the accompanying APN21/036SVI (C. Walkley) Blood Report.

### Microscopic Observations

**SUMMARY**
Both the Adar1P195A/E861A mutant animals and the WT controls of this cohort showed features indicative of increased cerebrospinal fluid in the lateral ventricles of the brain (hydrocephalus). These features were seemingly more prominent in the Adar1P195A/E861A mutants.
Note: hydrocephalus is a background lesion seen in some inbred mouse strains.
Various tissues showed multiple other incidental and/or background lesions - no obvious micromorphological differences were identified between the Adar1P195A/E861A mutant animals and WT controls of this cohort.
Please see individual animals for more micromorphological details.

**Notes:**
1. Reactive lymph nodes are defined as mild follicular hyperplasia, germinal centre formation and occasional sinus histiocytosis - a common finding in mice.
2. Hyperplastic lymph node follicles are identified by an increase in number and size of follicles and conversion to secondary follicles. Hyperplasia of the paracortex is characterized by an increase in the cell density and, depending on the degree of hyperplasia, an increase in the paracortical area.
3. Accumulation of leukocytes and other cells are common nonneoplastic lesions in many tissues. The term “inflammation” is used when the cell (leukocyte) accumulations are part of an active inflammatory process (typified by concurrent features such as vascular changes, necrosis, fibrosis, and/or tissue...
In contrast, cell "infiltration" is used when the cell (e.g., lymphocyte) accumulations are present in tissue without other disruption or pathology.

4. Focal inflammatory cell aggregates consisting of mononuclear, polymorphonuclear, and/or histiocytic cells are frequently observed in ageing mice (Maranpot RR. 1999. Pathology of the Mouse.). These can be present as lymphoid aggregates found in various tissues including the renal pelvis, bladder, lungs, liver (Pettan-Brewer C and Treuting PM. 2011. Practical pathology of aging mice.) and salivary glands (Haines DC, Chattopadhyay S and Ward JM. 2001. Pathology of Aging B6;129 Mice).

5. Mild extramedullary haematopoiesis (EMH) in the red pulp of the spleen is a common finding in the mouse. EMH consists of erythroid precursors, myeloid precursors, megakaryocytes or all three. While some degree of extramedullary haematopoiesis is present in normal rodents, especially in mice, increased extramedullary haematopoiesis can result from haematotoxin insult, systemic anaemia, and infections elsewhere in the body. (Suttie AW. 2006. Histopathology of the spleen).

6. The appearance of pigment in the red pulp is a common background lesion in rodents (Suttie AW. 2006. Histopathology of the spleen).

7. Thymic cysts in the rodent represent either a dilatation of thymic tubular structures or remnants of the thymopharyngeal duct. They are common findings in the involuted and/or atrophied thymus glands of rats and mice. Thymic cyst formation becomes more prominent with age and is associated with involution. (Hobbie K, Elmore SA and Kolenda-Roberts HM. 2015. Thymus – Cyst. In: National Toxicology Program Nonneoplastic Lesion Atlas).

8. Cytoplasmic inclusions of homogeneous eosinophilic hyaline-like material may be seen in older mice in intrahepatic biliary epithelial cells, as well as epithelial cells in the gallbladder. In marked cases, there is hyperplasia of the glandular epithelium, and crystalline forms of the eosinophilic inclusion material may be present both intracellularly and extracellularly (Maronpot RR. 1999. Liver and gallbladder. In: National Toxicology Program Nonneoplastic Lesion Atlas).

9. Focal fatty change of the liver can be a spontaneous lesion and may be more common in some strains than others (Maronpot RR. 2014. Liver-Fatty Change. In: National Toxicology Program Nonneoplastic Lesion Atlas).

10. Fatty change of the liver may occur in mice as a response to a toxicant. It is also seen in old obese controls and is more common in male than in female mice. The degree of fatty metamorphosis may vary and usually starts with a centriflobular distribution (Frith CH, Ward JM. 1988. Digestive System. In: Color Atlas of Neoplastic and Non-neoplastic Lesions in Aging Mice).


12. Focal pancreatic infiltrates of lymphocytes, plasma cells, and macrophages are uncommon in aged mice. The infiltrates are generally minimal and may be associated with atrophy (Maranpot RR. 1999. Exocrine and Endocrine Pancreas. In: Pathology of the Mouse: Reference and Atlas).

13. Yellow to brown pigment is often seen as an aging change in mouse and rat ovaries. Ceroid is the pigment most frequently seen, and this accumulates mainly in the cytoplasm of interstitial cells. Golden brown to darker brown pigment is sometimes seen in the uterus. Hemosiderin may be seen secondary to haemorrhage. Other types of pigment may also be seen, such as lipofuscin or ceroid. (Willson G, Cimon KY. 2015. Uterus, Endometrium – Hyperplasia, Cystic. In: National Toxicology Program Nonneoplastic Lesion Atlas).

14. The early histopathologic changes of nephropathy are characterised by the presence of focal to multifocal cortical tubules which have slight cytoplasmic basophilia and nuclear crowding. The basement membrane of these tubules are often variably thickened....As the disease progresses more affected tubules become evident and occasional eosinophilic proteinaceous tubular casts are noted (Maranpot RR. 1999. Kidney – Nephropathy. In: Pathology of the Mouse).

15. Lymphocyte apoptosis is characterized by cell shrinkage, nuclear pyknosis and fragmentation with apoptotic bodies. This type of cell death normally occurs within the germinal centres of secondary follicles where it is an important homeostatic mechanism (Hobbie K, Elmore SA and Kolenda-Roberts HM. 2015. Lymph Node – Apoptosis, Lymphocyte. In: National Toxicology Program Nonneoplastic Lesion Atlas).

16. Valvular myxomatous changes or degeneration can be an age-related spontaneous or chemical-induced change. The lesion is characterized by focal or segmental thickening of the subendocardium in the valve leaflets and expansion of the spongiosa of the valve leaflet with extracellular fibromyxoid material composed predominantly of glycosaminoglycans. Occasionally, fibrin deposits or thrombi and collections of neutrophils or mononuclear cells are seen (Johnson CL, Nyska A. 2017. Heart, Valve – Degeneration. In: National Toxicology Program Nonneoplastic Lesion Atlas).

17. A number of intestinal parasites may be seen within the lumen of the large intestine, the caecum, and less commonly the small intestine. The significance and presence of protozoan organisms is questionable since infected animals are normally asymptomatic. The presence of Protozoa such as flagellates, ciliates etc. have not been associated with any microscopic lesions of clinical significance in mice (Maranpot RR. 2014. Pathology of the Mouse: Reference and Atlas).

18. Necrosis of the exocrine pancreas, a naturally occurring lesion in mice usually only involving a few acini. The spontaneous lesion is both uncommon and very limited within the pancreas (Maronpot RR. 1999. Exocrine and Endocrine Pancreas. In: Pathology of the Mouse: Reference and Atlas).

19. Epidermal ulcers are among the most common spontaneous findings in mice and rats but may also be associated with dermal application of test agents. Ulcers are characterized by segmental or more extensive loss of the epidermis, including the basement membrane, with exposure of the underlying dermis. Erosion is characterized by the partial loss of the epithelium, with the basement membrane left intact. (Boyle MH, Hill GD. 2014. Skin - Ulcer and Erosion. NTP Nonneoplastic Lesion Atlas).

20. Focal hepatic necrosis is a non-specific entity quite often encountered as an incidental finding in the liver of mice. It can be the result of viruses (mouse hepatitis), bacteria (Clostridium piliforme), toxicants, and ischemia while the etiology is often unknown. It may involve single cells, single or multiple lobules, and it may vary in distribution. Coagulation necrosis with distinct eosinophilic cytoplasm and pyknotic or absent nuclei is the typical morphologic feature (Frith CH, Ward JM. 1988. Color Atlas of Neoplastic and Non-neoplastic Lesions in Aging Mice).

21. Ovarian cysts are a common finding in rats and mice. Ovarian cysts may be unilateral or bilateral, single or multiple, and may become quite large... size and number of cysts increase with age. (Willson G, Cimon KY. 2015. Ovary - Cyst. In: National Toxicology Program Nonneoplastic Lesion Atlas).

22. Hydrocephalus may be communicating or noncommunicating; that is, the former has no apparent obstructive process, whereas the latter has an obstructive cause somewhere in the ventricular connections. Most commonly, communicating hydrocephalus is considered to result from an idiopathic increase in cerebrospinal fluid production or deceased resorption. (Little P, Rao DB. 2014. Brain – Hydrocephalus. In: National Toxicology Program Nonneoplastic Lesion Atlas).

23. Germ cell degeneration of the testes is a nonspecific term that generally includes a number of degenerative features, such as tubular vacuolation, partial depletion of germ cells, degenerating (multinucleated or apoptotic) germ cells, and disordered arrangement of the germ cell layers. Chemically induced germ cell degeneration can be multifocal in distribution, but it is most often a bilateral lesion that affects most of the seminiferous tubules to varying degrees. It can also be an incidental background finding in rats and mice of any age, but the incidence increases with age. (Wilson G and Cimon K Y. 2015. Testis, Germ cell – Degeneration. In: National Toxicology Program Nonneoplastic Lesion Atlas).

24. Small focal clusters of inflammatory cells in the interstitium are common incidental findings in the Harderian gland of rats and mice. These infiltrates are most commonly mononuclear cells (lymphocytes), but other inflammatory cells may also be present (Gruebbel MM. 2014. Harderian Gland – Infiltration Cellular, Mononuclear Cell. In: National Toxicology Program Nonneoplastic Lesion Atlas).

25. Exocrine acinar atrophy is the most common spontaneous degenerative change in the pancreas of both rats and mice. Acinar atrophy may range from focal atrophic exocrine acini with no inflammation or fibrosis to diffuse atrophy of exocrine acini with replacement by adipose tissue and residual ducts, vasculature and islets of Langerhans. Acinar atrophy frequently represents the sequel of chronic inflammation, and as such, is often accompanied by infiltrates of mononuclear cells and fibrosis. (Nolte T, Brander-Weber P and Dangler C. 2016. Pancreas (Exocrine) – Atrophy, acinar cell. In: Nonproliferative and Proliferative Lesions of the Gastrointestinal Tract, Pancreas and Salivary Glands of the Rat and Mouse. J Toxicol Pathol 2016; 29 (1 Suppl): 1S–124S).

**#20 (control)**

Macro Observations

Tail suspension test for neurological defects - negative.

Dentition, tongue and oral cavity was unremarkable.

BCS: 3

Testes: 8x5x4mm, symmetrical

Spleen: 15x5x3mm

Kidneys: 13x7x6mm, symmetrical

Thymus: 7x5x2mm

Lungs inflated.

Heart: 10x8x7mm

Brain: 15x11x5mm, symmetrical

Pituitary gland identified, macroscopically normal

Tail length: 83mm (straight)

Head harvested for evaluation of auditory and vestibular structures.

Bone marrow smear taken from left hind leg.
No macroscopic lesions identified.
Excessive adipose tissue: abdominal viscera (+).

**Micro Observations**

Animal #20 was used as a male histological control for this cohort.

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.

Organs examined: Testes/Epididymes (85198), Seminal vesicles (85199, 85200), Prostate glands (85199, 85200), Penis/Preputial gland (85201), Urinary Bladder (85199, 85200), Liver/Gall Bladder (85202), Stomach (85203), Duodenum/jejunum/ileum/GALT (85203, 85204, 85205), Cecum/Colon/GALT (85205), Mesenteric Lymph Node (85197), Spleen (85199, 85200), Pancreas (85199, 85200), Kidneys/Adrenal Glands (85206), Salivary Glands/regional lymph nodes (85197), Thyroids (85208), Trachea/Lungs (85207, 85208), Thymus (85207, 85208), Heart (85207, 85208), Skin (85209), Tail (85279, 85280, 85281), Eyes/Harderian Glands (85210), Brain (85211), Spinal cord (85283, 85284, 85285), Hind leg (85279, 85280, 85281), Head (85312, 85313, 85314), Sternum (85282).

**Micromorphological changes** -

- Testes: few tubules showing tubular degeneration/atrophy (85198).
- Liver: multiple small foci of hepatocyte cell loss/necrosis and associated leucocyte infiltrates (85202).
- Liver: minimal multifocal perivascular mononuclear cell infiltrates (85202).
- Gall bladder: epithelial hyperplasia, cytoplasmic hyaline droplet accumulation, and neutrophilic infiltrates in the lamina propria (85202).
- Large intestine: increased mononuclear cell infiltrates in the lamina propria of the colon (85205).
- Pancreas: focal necrosis and associated leukocyte infiltrates (85203).
- Pancreas: minimal multifocal perivascular mononuclear cell infiltrates (85199, 85200).
- Kidneys: minimal multifocal perivascular/peripelvic mononuclear cell infiltrates (85206).
- Kidneys: single granular cast-like structure in cortex (85206).
- Salivary glands: mild multifocal perivascular mononuclear cell infiltrates, also seen in the associated lacrimal gland (85197).
- Lung: minimal multifocal perivascular mononuclear cell infiltrates (85207, 85208).
- Thymus: few small sized cysts (85208).
- Heart: few foci of cardiomyocyte vacuolation, indicative of degeneration (85207, 85208).
- Eye: query thinning/loss of the corneal epithelial cell layer, unilateral (85210). No other indicators of inflammation or damage.
- Brain: some flattening/stretching of the ependymal cells in the lateral ventricles (85211), indicative of increased cerebrospinal fluid (hydrocephalus).
- Hind leg: two foci of myodegeneration and necrosis (85280, 85281). Some regenerating myofibres identified.
- Spinal cord: few foci of myodegeneration (85284).
- Head: few foci of cytoplasmic hyaline droplet accumulation in the nasal epithelium with scattered mucosal neutrophils (85313).
- Head: inflammation of a single hair follicle in the upper lip (85314).

**#5 (control)**

**Macro Observations**

Tail suspension test for neurological defects - negative.
Dentition, tongue and oral cavity was unremarkable.
BCS: 3
Spleen: 18x5x2mm
Kidneys: 12x8x6mm, symmetrical
Thymus: 6x6x2mm
Lungs inflated.
Heart: 12x6x5mm
Brain: 16x11x5mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 85mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Animal #5 was used as a female histological control for this cohort.

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.

(85084)

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.

(85085)

Organs examined: Mammary glands (85092, 85096), Ovaries/oviducts (85086, 85087), Uterus/cervix/vagina (85086, 85087), Urinary Bladder (85086, 85087), Liver/Gall Bladder (85088), Stomach (85089), Duodenum/jejenum/ileum/GALT (85089, 85090, 85091), Cecum/Colon/GALT (85091), Mesenteric Lymph Node (85092), Spleen (85086, 85087), Pancreas (85086, 85087), Kidneys/Adrenal Glands (85093), Salivary Glands/regional lymph nodes (85092), Thyroids (85094, 85095), Trachea/Lungs (85094, 85095), Thymus (85094, 85095), Heart (85094, 85095), Skin (85096), Tail (85247, 85248, 85249), Eyes/Harderian Glands (85097), Brain (85098), Spinal cord (85245, 85246), Hind leg (85247, 85248, 85249), Head (85292, 85293), Sternum (85244).

Micromorphological changes-
Ovaries: occasional pigment laden interstitial cells (likely ceroid-lipofuscin) (85086, 85087).
Oviducts: epithelial cytoplasmic vacuolation, bilateral (85086, 85087).
Uterus: cystic endometrial hyperplasia (85086, 85087).
Liver: mild multifocal perivascular mononuclear cell infiltrates (85088).
Liver: small foci of hepatocyte cell loss/necrosis and associated leucocyte infiltrates (mixed) (85088).
Gall bladder: scattered neutrophils in the lamina propria (85088).
Large intestine: of the cecum, abundant luminal protozoa, and increased mononuclear cell infiltrates in the lamina propria (85091).
Pancreas: minimal multifocal perivascular mononuclear cell infiltrates (85086, 85087).
Pancreas: focal loss/necrosis of the exocrine pancreas (acini) with mainly mononuclear cell infiltrates (85089).
Kidneys: several protein casts within the cortex and medulla regions. Few in the medulla are mildly dilated (85093).
Kidney: minimal multifocal perivascular and parenchymal mononuclear cell infiltrates (85093).
Salivary glands: mild multifocal perivascular mononuclear cell infiltrates within the submandibular gland (85092).
Thymus: single small sized cyst (85094).
Lungs: few small foci of intra-alveolar haemorrhage, judged to be artefactual (85094, 85095).
Heart: few foci of mononuclear cell infiltrates and cardiomyocyte degeneration (85094).
Heart: myxomatous valvular changes (thickened leaflets) (85094).
Brain: some flattening/stretching of the ependymal cells in the lateral ventricles (85098), indicative of increased cerebrospinal fluid (hydrocephalus).
Spinal: minimal focal eosinophilic infiltrate in the surrounding connective tissue (85245).
Head: minimal multifocal peri-glandular mononuclear cell infiltrates within the cervical mammary tissue (85292, 85293).
Head: few foci of mixed leucocyte infiltrates within the neck muscle (85292).
Multiple small to parge sized perivascular mononuclear cell aggregates within the surrounding peri-adipose tissues of the lungs and reproductive organs.

#6 (control)

Macro Observations

Tail suspension test for neurological defects - negative.
Dentition, tongue and oral cavity was unremarkable.
BCS: 3
Spleen: 13x4x2mm
Kidneys: 11x6x4mm, symmetrical
Thymus: 7x7x2mm
Lungs inflated.
Heart: 9x8x5mm
Brain: 15x10x5mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 84mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified
Notable head shape- arch of snout (tip of nose) to crown (top of head between the ears) was prominently rounded/convex.

Micro Observations

Animal #6 was used as a female histological control for this cohort.

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.
(85099)

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.
(85100)

Organs examined: Mammary glands (85107, 85111), Ovaries/oviducts (85101, 85102), Uterus/cervix/vagina (85101, 85102), Urinary Bladder (85101, 85102), Liver/Gall Bladder (85103), Stomach (85104), Duodenum/jejenum/ ileum/GALT (85104, 85105, 85106), Cecum/Colon/GALT (85106), Mesenteric Lymph Node (85107), Spleen (85101, 85102), Pancreas (85101, 85102), Kidneys/Adrenal Gland (85108), Salivary Glands/regional lymph nodes (85107), Thyroids (85109), Trachea/Lungs (85109, 85110), Thymus (85109, 85110), Heart (85109, 85110), Skin (85111), Tail (85234, 85235, 85236), Eyes/Harderian Glands (85112), Brain (85113), Spinal cord (85231, 85232), Hind leg (85234, 85235, 85236), Head (85294, 85295, 85296), Sternum (85233).

Micromorphological changes-
Ovaries: occasional pigment laden interstitial cells (likely ceroid-lipofucsin) (85101, 85102).
Oviducts: focal epithelial cytoplasmic vacuolation, bilateral (85101, 85102).
Uterus: cystic endometrial hyperplasia (85101, 85102).
Liver: multiple small parenchymal leukocyte aggregates with some hepatocyte cell loss/necrosis (85103).
Liver: minimal multifocal perivascular mononuclear cell infiltrates (85103).
Gall bladder: focal epithelial hyperplasia, and underlying neutrophils in the lamina propria (85103).
Small intestine: of the distal half, increased number of eosinophils in the lamina propria (85105, 85106).
Large intestine: in mainly the cecum, abundant luminal protozoa (85106).
Pancreas: minimal focal perivascular mononuclear cell infiltrate (85101).
Salivary glands: of the parotid gland, small cluster of enlarged basophilic acini cells nearby a small focus of glandular atrophy (ducts and acini) (85107).
Salivary glands: another larger focus of glandular atrophy (ducts and acini), likely of the parotid gland (85107).
Kidney: multiple mildly dilated protein casts within the medulla (85108).
Thymus: few small sized cysts (85109, 85110).
Heart: myxomatous valvular changes (thickened leaflets) (85109).
Heart: focal myocardiacyte degeneration of the ventricular wall (85109).
Lungs: single small sized parenchymal mononuclear cell aggregate (85110).
Lungs: few small mononuclear cell aggregates within the surrounding peri-adipose tissue (85109, 85110).
Brain: some flattening/stretching of the ependymal cells in the lateral ventricles (85113), indicative of increased cerebrospinal fluid (hydrocephalus).
Hind leg: multifocal mixed leucocyte inflammation within the digits of the foot (85235, 85236).
Hing leg: oedematous changes within the subcutis above the foot (85234, 85235, 85236).
Head: some hyaline droplet accumulation within the nasal epithelium (85294, 85295, 85296).
Head: multiple small inflammatory foci in the skin of the outer ear (85295, 85296), query focal inflammation/necrosis of the outer surface of the tympanic membrane (85295).

#11
Macro Observations

Tail suspension test for neurological defects - negative.
Dentition, tongue and oral cavity was unremarkable.
BCS: 3
Spleen: 13x4x3mm
Kidneys: 11x6x5mm, symmetrical
Thymus: 8x7x2mm
Lungs inflated.
Heart: 8x8x6mm
Brain: 14x10x5mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 81mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.
(85148)
Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.
(85149)

Mammary glands

Typical mammary fat pad with developing lactiferous ducts, blood vessels, and nerve bundles.
(85156)
No lesions of significance
Ovaries/Oviducts

Unremarkable ovaries containing follicles at various stages of development (primary through to antral) and several corpora lutea.
Clusters of pigment laden interstitial cells (likely ceroid-lipofuscin), an age-related change - feature seen in the controls.
Mostly unremarkable oviduct micromorphology with typical columnar epithelium and mucosal folds.
Focal epithelial cytoplasmic vacuolation, an age-related change - feature seen in the controls.
(85150, 85151)
No lesions of significance

Uterus/Cervix/Vagina/Clitoral gland

Cystic endometrial hyperplasia, an age-related change - feature seen in the controls.
Unremarkable architecture myometrium and adventitia.
The micromorphology of the uterus and vagina places the animal at proestrus.
(85150, 85151)
No lesions of significance

Urinary Bladder

Unremarkable bladder with typical urothelium and detrusor muscle.
(85150, 85151)
No lesions of significance

Liver/Gall bladder

Liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Areas of hydropic degeneration of hepatocytes, an age-related finding in mice.
Multiple small foci of parenchymal leukocyte aggregates and some hepatocyte cell loss/necrosis - common incidental/background finding - feature seen in the controls.
Gall bladder with epithelial hyperplasia, cytoplasmic hyaline droplet accumulation, and underlying neutrophils in the lamina propria, common incidental/background finding - feature seen in the controls.
(85152)

Comments:
Pathology to comment

Stomach

Unremarkable fore and glandular portions of the stomach with limiting ridge.
Includes pyloric sphincter and duodenal bulb.
(85153)
No lesions of significance

Small Intestine (Duodenum, Jejunum & Ileum)/GALT

Typical mucosal villi and submucosal layers.
Peyer's patches display typical reactive nodal histology. Occasional, typical lymphoid cluster (cryptopatches).
(85153, 85154, 85155)
No lesions of significance

Cecum/Colon/GALT

Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion cells of the plexuses.
Occasional, typical lymphoid cluster (Peyer's patch).
(85155)
No lesions of significance
Mesenteric lymph node

Typical reactive nodal histology including enlarged size, mild follicular hyperplasia, an expansive paracortical area with some lymphocyte apoptosis, and sinus histiocytosis.

(85156)
No lesions of significance

Spleen

Mild follicular hyperplasia with germinal centre formation and some lymphocyte apoptosis. Mild coalescence of the white pulp.
Expansive red pulp with marked extramedullary haematopoiesis and conspicuous haemosiderin laden macrophages.

(85151)
No lesions of significance

Pancreas

Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).

(85150, 85151)
No lesions of significance

Kidney

Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable.
Minimal perivascular/peripelvic mononuclear cell infiltrates, incidental/age-related change - feature seen in the controls.
Several protein casts within the medulla and papilla, moderately dilated within the medulla, incidental/age-related change - feature seen in the controls.
Renal lymph nodes with typical reactive nodal histology and mild sinus histiocytosis.

(85157)
No lesions of significance

Adrenal glands

Adrenal glands with typical cortex/medulla micromorphology.

(85157)
No lesions of significance

Salivary glands and Regional lymph nodes

Unremarkable submandibular, sublingual and parotid glands.
Regional lymph nodes display typical reactive nodal histology.

(85156)
No lesions of significance

Thyroids

Normal lateral lobes of the thyroid gland with typical colloid secreting follicles lined by cuboidal epithelium.
Small sheet-like mass of polygonal cells, characteristic of the parathyroid gland.

(85158, 85159)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, blood vessels and parabronchial lymph node.
Minimal multifocal perivascular mononuclear cell infiltrates, an incidental/age-related change - feature seen in the controls.
Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.

(85158, 85159)
No lesions of significance
Thymus

Typical medulla/cortex distribution and micromorphology.
Single small sized cyst (85158), common incidental/background finding - feature seen in the controls.

(85158, 85159)
No lesions of significance

Heart/chambers/vessels/valves

Mostly typical micromorphology observed in cardiac muscle, chambers, valves and great vessels of the heart.
Myxomatous valvular changes (thickened leaflets), an age-related change - feature seen in the controls.
The cardiac muscle fibres demonstrated typical features including central nuclei, branching fibres and striations.

(85158, 85159)
No lesions of significance

Skin

Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle (panniculus carnosus).
Minimal focal mononuclear cell infiltrate surrounding one hair follicle, common incidental finding in mice.

(85160)
No lesions of significance

Tail

Typical tail components including keratinized squamous epithelium, dense regular connective tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and blood vessels.

(85267, 85268, 85269)
No lesions of significance

Eyes/Harderian glands

Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
Typical branched tubuloalveolar formation of the Harderian gland.
Includes portion of unremarkable optic nerve and extraocular muscles.

(85161)
No lesions of significance

Brain

Sections were prepared from the standard levels of the brain:

Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third ventricles.
Level III hindbrain: includes the cerebellum, pons and fourth ventricle.

Sections of brain appear symmetrical with unremarkable meninges and typical lamination.
The cerebellum appears symmetrical with typical architecture and Purkinje cells.
There was no obvious neuronal loss and the myelination appears normal.

Some flattening/stretching of the ependymal cells and increased size of the lateral ventricular space, features indicative of increased cerebrospinal fluid (hydrocephalus), a known background lesion in mice - similar feature seen in the controls.

(85162)

Comments:
**Neuropathology to comment**

**Spinal cord**

Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc, striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.

(85271, 85272)

No lesions of significance

**Comments:**

**Neuropathology to comment**

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle

Mostly unremarkable long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow. The skeletal muscle shows consistent fibre size with peripheral nuclei.

Focal myodegeneration adjacent the hip joint (85269) - feature seen in the controls.

(85267, 85268, 85269)

**Comments:**

**Pathology to comment**

**Head**

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85303). The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments (85304- 85306). Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outer hair cells and typical tectorial membrane (85304- 85306). The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85306).

Focal and bilateral inflammation of the nasal epithelium, seemingly due to inhaled foreign material.

(85303- 85306)

No lesions of significance

**Sternum**

Representative sternebrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat.

Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85270)

No lesions of significance

**#12**

**Macro Observations**

Tail suspension test for neurological defects - negative.

Dentition, tongue and oral cavity was unremarkable.

BCS: 3

Testes: 8x5x4mm, symmetrical

Spleen: 15x5x2mm
Kidneys: 14x8x5mm, symmetrical
Thymus: 6x6x2mm
Lungs inflated.
Heart: 10x8x6mm
Brain: 16x10x5mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 85mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.

Mammary glands

Typical mammary fat pad with developing lactiferous ducts, blood vessels, and nerve bundles.

No lesions of significance

Ovaries/Oviducts

Unremarkable ovaries containing follicles at various stages of development (primary through to antral) and several corpora lutea.
Scattered pigment laden interstitial cells (likely ceroid-lipofuscin), an age-related change - feature seen in the controls.
Single ovarian cyst, an incidental/background finding in mice.
Mononuclear cell infiltration of the bursa membrane, unilateral, an age-related finding in mice.
Mostly unremarkable oviduct micromorphology with typical columnar epithelium and mucosal folds.
Focal epithelial cytoplasmic vacuolation, an age-related change - feature seen in the controls.

No lesions of significance

Uterus/Cervix/Vagina/Clitoral gland

Endometrial hyperplasia, an age-related change - feature seen in the controls.
Unremarkable architecture myometrium and adventitia.

No lesions of significance

Urinary Bladder

Unremarkable bladder with typical urothelium and detrusor muscle.

No lesions of significance

Liver/Gall bladder

Liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Areas of hydropic degeneration of hepatocytes, an age-related finding in mice.
Several small parenchymal leukocyte aggregates - common incidental/background finding - feature seen in the controls.
Section does not include gall bladder.
Comments:
Pathology to comment

Stomach

Unremarkable fore and glandular portions of the stomach with limiting ridge.
Small focal basophilic deposit within the gland lumen (?mineralisation) - likely an incidental finding.
Includes pyloric sphincter and duodenal bulb.

(85168)
No lesions of significance

Small Intestine (Duodenum, Jejunum & Ileum)/GALT

Typical mucosal villi and submucosal layers.
Peyer's patches display typical reactive nodal histology. Occasional, typical lymphoid cluster (cryptopatches).
(85168, 85169, 85170)
No lesions of significance

Cecum/Colon/GALT

Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion cells of the plexuses.
Occasional, typical lymphoid cluster (Peyer's patch).
In mainly the cecum, abundant luminal protozoa - feature seen in the controls.
(85170)
No lesions of significance

Mesenteric lymph node

Typical reactive nodal histology with enlarged size, representative cortex including the occasional follicle, an expansive paracortical area with lymphocyte apoptosis, and sinus histiocytosis.
(85171)
No lesions of significance

Spleen

Mild follicular hyperplasia with germinal centre formation and some lymphocyte apoptosis. Mild coalescence of the white pulp.
Expansive red pulp with marked extramedullary haematopoiesis.
(85166)
No lesions of significance

Pancreas

Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).
(85166)
No lesions of significance

Kidney

Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable.
Minimal perivascular/peripelvic mononuclear cell infiltrates, incidental/age-related change - feature seen in the controls.
Few protein casts within the medulla, incidental/age-related change - feature seen in the controls.
Renal lymph nodes with typical reactive nodal histology.
(85172)
No lesions of significance
Adrenal glands

Adrenal glands with typical cortex/medulla micromorphology.
(85172)
No lesions of significance

Salivary glands and Regional lymph nodes

Unremarkable submandibular, sublingual and parotid glands.
Mild multifocal perivascular mononuclear cell infiltrates within the submandibular gland, an age-related change - feature seen in the controls.
Regional lymph nodes display typical reactive nodal histology.
(85171)
No lesions of significance

Thyroids

Normal lateral lobes of the thyroid gland with typical colloid secreting follicles lined by cuboidal epithelium.
Small sheet-like mass of polygonal cells, characteristic of the parathyroid gland.
(85173, 85174)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, blood vessels and parabronchial lymph node.
Mild multifocal perivascular/peribronchial mononuclear cell infiltrates, an incidental/age-related change - feature seen in the controls. Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.
Several small to medium sized mononuclear cell aggregates in the surrounding peri-adipose tissue, common finding - feature seen in the controls.
(85173, 85174)
No lesions of significance

Thymus

Typical medulla/cortex distribution and micromorphology.
(85173, 85174)
No lesions of significance

Heart/chambers/vessels/valves

Mostly typical micromorphology observed in cardiac muscle, chambers, valves and great vessels of the heart.
Myxomatous valvular changes (thickened leaflets), an age-related change - feature seen in the controls.
The cardiac muscle fibres demonstrated typical features including central nuclei, branching fibres and striations.
(85173, 85174)
No lesions of significance

Skin

Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle (panniculus carnosus).
(85175)
No lesions of significance
Tail
Typical tail components including keratinized squamous epithelium, dense regular connective tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and blood vessels.
(85261, 85262, 85263)
No lesions of significance

Eyes/Harderian glands
Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
Typical branched tubuloalveolar formation of the Harderian gland.
Includes portion of unremarkable optic nerve and extraocular muscles.
(85176)
No lesions of significance

Brain
Sections were prepared from the standard levels of the brain:
Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third ventricles.
Level III hindbrain: includes the cerebellum, pons and fourth ventricle.
Sections of brain appear symmetrical with unremarkable meninges and typical lamination.
The cerebellum appears symmetrical with typical architecture and Purkinje cells.
There was no obvious neuronal loss and the myelination appears normal.
Prominent flattening/stretching of the ependymal cells and increased size of the lateral ventricular space, features indicative of increased cerebrospinal fluid (hydrocephalus), a known background lesion in mice - similar feature seen in the controls.
Query vacuolation of the adjacent white matter, possible artefact.
(85177)
Comments:
Neuropathology to comment

Spinal cord
Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc, striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.
(85265, 85266)
No lesions of significance
Comments:
Neuropathology to comment

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle
Unremarkable long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow. The skeletal muscle shows consistent fibre size with peripheral nuclei.
Small focus of accumulated mononuclear cell infiltrates in the dermis (85263), common incidental finding.
(85261, 85262, 85263)
No lesions of significance
Head

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85307). The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments (85308, 85309).

Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outer hair cells and typical tectorial membrane (85308, 85309). The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85308, 85309).

(85307-85309)

No lesions of significance

Sternum

Representative sternebrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat.

Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85264)

No lesions of significance

#18

Macro Observations

Tail suspension test for neurological defects - negative.
Dentition, tongue and oral cavity was unremarkable.

BCS: 3
Testes: 8x6x4mm, symmetrical
Spleen: 18x6x2mm
Kidneys: 15x8x6mm, symmetrical
Thymus: 7x7x2mm
Lungs inflated.
Heart: 12x8x6mm
Brain: 16x11x6mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 80mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.

(85178)

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.

(85179)
Testes/Epididymes
Mostly typical convoluted seminiferous tubules at various stages of cycle surrounded by the tunica albuginea. Within the tubules, unremarkable spermatogenic cells including, Sertoli cells, spermatogonia, developing spermatocytes and spermatids. Typical interstitial Leydig cells. Multifocal tubular degeneration/atrophy, an age-related change - feature seen in the control. Unremarkable vas deferens with typical intraluminal sperm. Architecture of the epididymis is mostly typical, with numerous intraluminal elongated spermatozoa. Focal interstitial oedema-like changes in an epididymis, possible plane of section artefact. (85180)

Seminal vesicles
Unremarkable tall columnar epithelium and folded mucosa. Presence of typical intraluminal eosinophilic secretions. (85181, 85182) No lesions of significance

Prostate glands
Unremarkable dorsal lateral/ventral/coagulating glands with typical intraluminal secretions. Portion of unremarkable vas deferens. (85181, 85182) No lesions of significance

Penis/Preputial gland
Typical penile structures including prepuce, glans, corpus cavernosum and urethra. Typical preputial glands including basal and secretory cells. Skin shows focal accumulation of dermal melanocytes, a common incidental/background finding in mice. (85183) No lesions of significance

Urinary Bladder
Unremarkable bladder with typical urothelium and detrusor muscle. (85181, 85182) No lesions of significance

Liver/Gall bladder
Liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins. Several small parenchymal leucocyte aggregates, incidental/background finding - feature seen in the controls. Unremarkable gall bladder/section does not show gall bladder. (85184) No lesions of significance

Stomach
Mostly unremarkable fore and glandular portions of the stomach with limiting ridge. Segmental eosinophil infiltrates in the submucosa and lamina propria, common incidental finding in mice. Includes pyloric sphincter and duodenal bulb. (85185) No lesions of significance
Small Intestine (Duodenum, Jejunum & Ileum)/GALT

- Typical mucosal villi and submucosal layers.
- Peyer's patches display typical nodal histology. Occasional, typical lymphoid cluster (cryptopatches).
- No lesions of significance

Cecum/Colon/GALT

- Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion cells of the plexuses.
- Occasional, typical lymphoid cluster (Peyer's patch).
- No lesions of significance

Mesenteric lymph node

- Typical reactive nodal histology with enlarged size, representative cortex including the occasional follicle, an expansive paracortical area with some lymphocyte apoptosis, and sinus histiocytosis.
- No lesions of significance

Spleen

- Mild follicular hyperplasia with germinal centre formation and some lymphocyte apoptosis. Mild coalescence of the white pulp.
- Extramedullary haematopoiesis in the red pulp.
- No lesions of significance

Pancreas

- Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).
- No lesions of significance

Kidney

- Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable.
- Minimal perivascular/peripelvic mononuclear cell infiltrates, incidental/age-related change - feature seen in the controls.
- Few protein casts within the medulla, incidental/age-related change - feature seen in the controls.
- Focal cluster of irregular tubules showing tubule basophilia and shrunken cytoplasm/nuclear crowding, an incidental/background finding in mice.
- Renal lymph nodes with typical reactive nodal histology and mild sinus histiocytosis.
- No lesions of significance

Adrenal glands

- Adrenal glands with typical cortex/medulla micromorphology.
- No lesions of significance

Salivary glands and Regional lymph nodes

- Unremarkable submandibular, sublingual and parotid glands.
- Minimal multifocal perivascular mononuclear cell infiltrates within the submandibular gland, an age-related change - feature seen in the controls.
- Regional lymph nodes display typical reactive nodal histology.
- No lesions of significance
Thyroids

Typical colloid secreting follicles lined by cuboidal epithelium.
(85190, 85191)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, and blood vessels.
Minimal perivascular mononuclear cell infiltrates, an incidental/age-related change - feature seen in the controls.
Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.
(85190, 85191)
No lesions of significance

Thymus

Typical medulla/cortex distribution and micromorphology.
(85190, 85191)
No lesions of significance

Heart/chambers/vessels/valves

Mostly typical micromorphology observed in cardiac muscle, chambers, valves and great vessels of the heart.
Myxomatous valvular changes (thickened leaflets) (85191), incidental/age-related finding - feature seen in the controls.
Several foci of cardiomyocyte vacuolation and degeneration, likely incidental/age-related - feature seen in the controls.
Remaining cardiac muscle fibres demonstrated typical features including central nuclei, branching fibres and striations.
(85190, 85191)

Skin

Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle (panniculus carnosus).
(85192)
No lesions of significance

Tail

Typical tail components including keratinized squamous epithelium, dense regular connective tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and blood vessels.
(85273, 85274, 85275)
No lesions of significance

Eyes/Harderian glands

Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
Typical branched tubuloalveolar formation of the Harderian gland.
Includes portion of unremarkable optic nerve and extraocular muscles.
(85193)
No lesions of significance

Brain

Sections were prepared from the standard levels of the brain:

Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third ventricles.
Level III hindbrain: includes the cerebellum, pons and fourth ventricle.
Sections of brain appear symmetrical with unremarkable meninges and typical lamination. The cerebellum appears symmetrical with typical architecture and Purkinje cells. There was no obvious neuronal loss and the myelination appears normal.

Some flattening/stretching of the ependymal cells, indicative of increased cerebrospinal fluid (hydrocephalus), a known background lesion in mice - similar feature seen in the controls.

(85194)

**Comments:**
Neuropathology to comment

**Spinal cord**

Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc, striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.

(85277, 85278)

No lesions of significance

**Comments:**
Neuropathology to comment

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle

Representative long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow. The skeletal muscle shows consistent fibre size with peripheral nuclei.

Query focal necrosis in the foot (?bone) (85273).

(85273, 85274, 85275)

**Comments:**
Pathology to comment

**Head**

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85310). The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments.

Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outter hair cells and typical tectorial membrane.

The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85311).

Inflammation of the middle ear characterised by oedema, luminal cellular debris, multinucleated giant cells, and basophilic deposit/mineralisation, unilateral - an incidental/background finding in mice.

(85310, 85311)

**Sternum**

Representative sternebrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat.

Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85276)
No lesions of significance

**#3**

**Macro Observations**

- Tail suspension test for neurological defects - negative.
- Dentition, tongue and oral cavity was unremarkable.
- BCS: 3
- Testes: 8x5x4mm, symmetrical
- Spleen: 16x5x3mm
- Kidneys: 13x7x4mm, symmetrical
- Thymus: 7x6x2mm
- Lungs inflated.
- Heart: 11x8x7mm
- Brain: 15x10x5mm, symmetrical
- Pituitary gland identified, macroscopically normal
- Tail length: 84mm (straight)
- Head harvested for evaluation of auditory and vestibular structures.
- Bone marrow smear taken from left hind leg.

- Fur was slightly ruffled.
- No macroscopic lesions identified.

**Micro Observations**

- Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts. (85050)

- Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites. (85051)

**Testes/Epididymes**

- Mostly typical convoluted seminiferous tubules at various stages of cycle surrounded by the tunica albuginea. Within the tubules, unremarkable spermatogenic cells including, Sertoli cells, spermatogonia, developing spermatocytes and spermatids. Typical interstitial Leydig cells. Focal cluster of tubules showing degeneration/atrophy, an age-related change - feature seen in the control.
- Unremarkable vas deferens with typical intraluminal sperm.
- Architecture of the epididymis is typical, with numerous intraluminal elongated spermatozoa. (85052)
- No lesions of significance

**Seminal vesicles**

- Unremarkable tall columnar epithelium and folded mucosa.
- Presence of typical intraluminal eosinophilic secretions. (85053, 85054)
- No lesions of significance

**Prostate glands**

- Unremarkable dorsal lateral/ventral/coagulating glands with typical intraluminal secretions.
- Portion of unremarkable vas deferens. (85053, 85054)
- No lesions of significance
Penis/Preputial gland
   Typical penile structures including prepuce, glans, corpus cavernosum and urethra.
   Typical preputial glands including basal and secretory cells.
   (85055, 85056)
   No lesions of significance

Urinary Bladder
   Unremarkable bladder with typical urothelium and detrusor muscle.
   (85053, 85054)
   No lesions of significance

Liver/Gall bladder
   Typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
   Unremarkable gall bladder.
   (85057)
   No lesions of significance

Stomach
   Unremarkable fore and glandular portions of the stomach with limiting ridge.
   Includes pyloric sphincter and duodenal bulb.
   (85058)
   No lesions of significance

Small Intestine (Duodenum, Jejunum & Ileum)/GALT
   Typical mucosal villi and submucosal layers.
   Peyer's patches display typical reactive nodal histology. Occasional, typical lymphoid cluster
   (cryptopatches).
   (85058, 85059, 85060)
   No lesions of significance

Cecum/Colon/GALT
   Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion
   cells of the plexuses.
   Occasional, typical lymphoid cluster (Peyer's patch).
   (85060)
   No lesions of significance

Mesenteric lymph node
   Typical reactive mesenteric lymph node with representative cortex including the occasional
   follicle, an expansive paracortical area with lymphocyte apoptosis, and sinus histiocytosis.
   (85061)
   No lesions of significance

Spleen
   Mild follicular hyperplasia with germinal centre formation and some lymphocyte apoptosis. Mild
   coalescence of the white pulp.
   Extramedullary haematopoiesis in the red pulp.
   Haemosiderin laden macrophages identified in both the red and white pulp.
   (85053, 85054)
   No lesions of significance

Pancreas
   Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).
   (85053, 85054)
   No lesions of significance
Kidney

Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable. Minimal multifocal peripelvic mononuclear cell infiltrates, incidental/age-related finding - feature seen in the controls.
Few vacuolated glomeruli, change considered too mild to be significant.
Few protein casts within the cortex and medulla regions, incidental/age related finding - feature seen in the controls.
Sections do not include renal lymph nodes.
(85062)
No lesions of significance

Adrenal glands

Adrenal glands with typical cortex/medulla micromorphology.
(85062)
No lesions of significance

Salivary glands and Regional lymph nodes

Unremarkable submandibular, sublingual and parotid glands.
Minimal multifocal mononuclear cell infiltrates within the submandibular and parotid glands, an age-related change - feature seen in the controls.
Focal necrosis of the parotid gland, likely an incidental/spontaneous change (85061).
Regional lymph nodes display typical reactive nodal histology.
(85061)

Comments:
Pathology to comment

Thyroids

Normal lateral lobes of the thyroid gland with typical colloid secreting follicles lined by cuboidal epithelium.
Small sheet-like mass of polygonal cells, characteristic of the parathyroid gland.
(85063, 85064)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, blood vessels and parabronchial lymph node.
Focal parenchymal congestion and collapse judged to be artefactual.
Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.
Few small sized mononuclear cell aggegrates within the surrounding peri-adipose tissue, common finding - feature seen in the controls.
(85063, 85064)
No lesions of significance

Thymus

Typical medulla/cortex distribution and micromorphology.
Few small sized cysts (85064), common incidental/background finding - feature seen in the controls.
(85063, 85064)
No lesions of significance

Heart/chambers/vessels/valves

Mostly typical micromorphology observed in cardiac muscle, chambers, valves and great vessels of the heart.
Myxomatous valvular changes (thickened leaflets), incidental/age-related finding - feature seen in the controls.
Few foci of cardiomyocyte vacuolation and degeneration, likely incidental/age-related - feature
seen in the controls.
Remaining cardiac muscle fibres demonstrated typical features including central nuclei,
branching fibres and striations.
(85063, 85064)

Comments:
Pathology to comment

Skin

Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle
(panniculus carnosus).
Multifocal epidermal hyperplasia with underlying dermal infiltrates that are mainly neutrophilic,
common incidental/background finding in mice.
(85065)
No lesions of significance

Tail

Typical tail components including keratinized squamous epithelium, dense regular connective
tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and
blood vessels.
(85237, 85238, 85239)
No lesions of significance

Eyes/Harderian glands

Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
Typical branched tubuloalveolar formation of the Harderian gland.
Unremarkable optic nerve and extraocular muscles.
(85066)
No lesions of significance

Brain

Sections were prepared from the standard levels of the brain:
Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third
ventricles.
Level III hindbrain: includes the cerebellum, pons and fourth ventricle.

Sections of brain appear symmetrical with unremarkable meninges and typical lamination.
The cerebellum appears symmetrical with typical architecture and Purkinje cells.
There was no obvious neuronal loss and the myelination appears normal.

Prominent flattening/stretching of the ependymal cells and increased size of the lateral
ventricular space, features indicative of increased cerebrospinal fluid (hydrocephalus), a known
background lesion in mice - similar feature seen in the controls.
Query vacuolation of the adjacent white matter, possible artefact.
(85067)

Comments:
Neuropathology to comment

Spinal cord

Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc,
striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.
Focal loss of the epidermis (ulcer) and multifocal inflammation affecting mainly the subcutis.
Segmental degeneration of the adjacent panniculus muscle (85243).
(85241, 85242, 85243)

Comments:

Neuropathology to comment

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle

Representative long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow.

Focal mixed leucocyte inflammation within one digit of the foot (85237).

Focal myodegeneration adjacent the hip joint (85237). Displays some features of regeneration.

Considered to be incidental changes - features seen in the controls.

Minimal focal perivascular eosinophilic infiltrate (85238, 85239).

Query scattered necrotic myofibres (85238, 85239).

Remaining skeletal muscle shows consistent fibre size with peripheral nuclei.

(85237, 85238, 85239)

Comments:

Pathology to comment

Head

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85286).

The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments (85287, 85288).

Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outer hair cells and typical tectorial membrane (85287, 85288).

The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85288).

(85286-85288)

No lesions of significance

Sternum

Representative sternebrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat.

Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85240)

No lesions of significance

#4

Macro Observations

Tail suspension test for neurological defects - negative.

Dentition, tongue and oral cavity was unremarkable.

BCS: 3

Spleen: 20x8x2mm

Kidneys: 14x8x5mm, symmetrical

Thymus: 7x7x2mm

Lungs inflated.

Heart: 12x10x6mm

Brain: 16x11x5mm, symmetrical

Pituitary gland identified, macroscopically normal
Tail length: 81mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.
(85068)

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.
(85069)

Mammary glands

Typical mammary fat pad with developing lactiferous ducts, blood vessels, and nerve bundles.
No lesions of significance

Ovaries/Oviducts

Unremarkable ovaries containing follicles at various stages of development (primary through to antral) and several corpora lutea.
Few small to medium sized cysts in one ovary, common incidental/background finding in the mouse.
Mostly unremarkable oviduct micromorphology with typical columnar epithelium and mucosal folds.
Focal epithelial cytoplasmic vacuolation (85072), an age-related change - feature seen in the controls.
(85070, 85071, 85072)
No lesions of significance

Uterus/Cervix/Vagina/Clitoral gland

Cystic endometrial hyperplasia, an age-related change - feature seen in the controls.
Unremarkable myometrium and adventitia.
Minimal multifocal perivascular mononuclear cell infiltrates within the surrounding peri-adipose tissue, common background finding - feature seen in the controls.
The micromorphology of the uterus and vagina places the animal at estrus.
(85070, 85071, 85072)
No lesions of significance

Urinary Bladder

Portion of unremarkable detrusor muscle.
(85072)
No lesions of significance

Liver/Gall bladder

Liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Multiple small foci of parenchymal inflammation and necrosis, common incidental/background finding - similar feature seen in the controls.
Unremarkable gall bladder.
(85073)

Comments:
Pathology to comment
Stomach

Unremarkable fore and glandular portions of the stomach with limiting ridge.
Includes pyloric sphincter and duodenal bulb.
(85074)
No lesions of significance

Small Intestine (Duodenum, Jejunum & Ileum)/GALT

Typical mucosal villi and submucosal layers.
Peyer's patches display typical reactive nodal histology. Occasional, typical lymphoid cluster (cryptopatches).
(85074, 85075, 85076)
No lesions of significance

Cecum/Colon/GALT

Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion cells of the plexuses.
Occasional, typical lymphoid cluster (Peyer's patch).
In mainly the cecum, abundant luminal protozoa - feature seen in the controls.
(85076)
No lesions of significance

Mesenteric lymph node

Highly reactive micromorphology including enlarged size, follicular hyperplasia with germinal centre formation, an expansive paracortical area containing lymphocyte apoptosis, and sinus histiocytosis.
(85077)
No lesions of significance

Spleen

Mild follicular hyperplasia with germinal centre formation and some lymphocyte apoptosis. Mild coalescence of the white pulp.
Expansive red pulp with marked extramedullary haematopoiesis.
Haemosiderin laden macrophages identified in both the red and white pulp.
(85070, 85071, 85072)
No lesions of significance

Pancreas

Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).
(85070, 85071, 85072)
No lesions of significance

Kidney

Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable.
Minimal focal perivascular/peripelvic mononuclear cell infiltrate, incidental/age-related change - feature seen in the controls.
Few mildly dilated protein casts within the medulla, incidental/age-related change - feature seen in the controls.
Renal lymph node with typical reactive nodal histology.
(85078)
No lesions of significance
Adrenal glands

Adrenal glands with typical cortex/medulla micromorphology.
(85078)
No lesions of significance

Salivary glands and Regional lymph nodes

Unremarkable submandibular, sublingual and parotid glands.
Mild multifocal perivascular mononuclear cell infiltrates within the submandibular gland, an age-related change - feature seen in the controls.
Regional lymph nodes display typical reactive nodal histology.
(85077)
No lesions of significance

Thyroids

Normal lateral lobes of the thyroid gland with typical colloid secreting follicles lined by cuboidal epithelium.
Small sheet-like mass of polygonal cells, characteristic of the parathyroid gland.
(85079, 85080)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, blood vessels and parabronchial lymph node.
Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.
Mild multifocal perivascular mononuclear cell infiltrates within the surrounding peri-adipose tissue, common finding - feature seen in the controls.
(85079, 85080)
No lesions of significance

Thymus

Typical medulla/cortex distribution and micromorphology.
(85079, 85080)
No lesions of significance

Heart/chambers/vessels/valves

Mostly typical micromorphology observed in cardiac muscle, chambers, valves and great vessels of the heart.
Myxomatous valvular changes (thickened leaflets) (85080), incidental/age-related finding - feature seen in the controls.
Occasional foci of cardiomyocyte vacuolation and degeneration, likely incidental/age-related - feature seen in the controls.
Remaining cardiac muscle fibres demonstrated typical features including central nuclei, branching fibres and striations.
(85079, 85080)

Comments:
Pathology to comment

Skin

Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle (panniculus carnosus).
(85081)
No lesions of significance
Tail

- Typical tail components including keratinized squamous epithelium, dense regular connective tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and blood vessels.
- No lesions of significance

Eyes/Harderian glands

- Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
- Typical branched tubuloalveolar formation of the Harderian gland.
- Includes portion of unremarkable optic nerve and extraocular muscles.
- No lesions of significance

Brain

- Sections were prepared from the standard levels of the brain:
  - Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
  - Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third ventricles.
  - Level III hindbrain: includes the cerebellum, pons and fourth ventricle.
- Sections of brain appear symmetrical with unremarkable meninges and typical lamination.
- The cerebellum appears symmetrical with typical architecture and Purkinje cells.
- There was no obvious neuronal loss and the myelination appears normal.
- Some flattening/stretching of the ependymal cells, indicative of increased cerebrospinal fluid (hydrocephalus), a known background lesion in mice - similar feature seen in the controls.
- Query vacuolation of the adjacent white matter, possible artefact.
- No lesions of significance

Spinal cord

- Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc, striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.
- No lesions of significance

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle

- Unremarkable long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow. The skeletal muscle mostly shows consistent fibre size with peripheral nuclei.
- Focal myodegeneration adjacent the hip joint, likely incidental - feature seen in the controls.
- Marked focal mixed leukocyte inflammation of the foot (85257), likely incidental - feature seen in the controls.

- No lesions of significance

Comments:

Neuropathology to comment
Head

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85289). The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments (85290, 85291). Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outer hair cells and typical tectorial membrane (85290, 85291). The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85291).

Inflammation of the middle ear characterised by oedema, luminal cellular debris, and some formation of cholesterol clefts (85291), unilateral - a known incidental/background finding in mice.
Few foci of cytoplasmic hyaline droplet accumulation within the nasal epithelium (85290, 85291), incidental - feature seen in the controls.

(85289-85291)

Comments:
Pathology to comment

Sternum

Representative sternebrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat. Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85258)

No lesions of significance

#7

Macro Observations

Tail suspension test for neurological defects - negative.
Dentition, tongue and oral cavity was unremarkable.
BCS: 3
Spleen: 20x8x2mm
Kidneys: 14x8x5mm, symmetrical (left kidney bisected coronally)
Thymus: 7x7x2mm
Lungs inflated.
Heart: 12x10x6mm
Brain: 16x10x6mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 82mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.

(85114)
Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites. (85115)

Mammary glands

Typical mammary fat pad with developing lactiferous ducts, blood vessels, and nerve bundles. Minimal multifocal perivascular mononuclear cell infiltrates (85128), incidental/age-related - feature seen in the controls. (85128)

No lesions of significance

Ovaries/Oviducts

Unremarkable ovaries containing follicles at various stages of development (primary through to antral) and several corpora lutea.
Moderate mononuclear cell infiltration of the bursa membrane, bilateral, an age-related change in mice.
Mostly unremarkable oviduct micromorphology with typical columnar epithelium and mucosal folds.
Focal epithelial cytoplasmic vacuolation, bilateral, an age-related change - feature seen in the controls.
Mild multifocal perivascular mononuclear cell infiltrates in the surrounding peri-adipose tissue, an age-related change - feature seen in the controls. (85116-85119)

No lesions of significance

Uterus/Cervix/Vagina/Clitoral gland

Endometrial hyperplasia, an age-related change - feature seen in the controls.
Unremarkable architecture of the myometrium and adventitia.
The micromorphology of the uterus and vagina places the animal at metestrus. (85116-85119)

No lesions of significance

Urinary Bladder

Portion of unremarkable bladder with typical urothelium and detrusor muscle. (85119)

No lesions of significance

Liver/Gall bladder

Liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Minimal multifocal perivascular mononuclear cell infiltrates, an incidental/age-related change - feature seen in the controls.
Multiple small parenchymal leucocyte aggregates, and focal parenchymal necrosis, common incidental/background findings in mice - similar feature seen in the controls.
Several small foci of hepatocyte vacuolation resembling fatty change (steatosis), an incidental/age-related finding in mice.
Unremarkable gall bladder. (85120)

Comments: Pathology to comment

Stomach

Unremarkable fore portion of the stomach.
Of the glandular portion, marked and mainly submucosal eosinophilic inflammatoin (transmural), common incidental finding in mice.
Sections includes pyloric sphincter and duodenal bulb.
Small Intestine (Duodenum, Jejunum & Ileum)/GALT

Mostly typical mucosal villi and submucosal layers.
Marked and mainly eosinophilic infiltration of the distal ileum (transmural) (85122), likely an incidental finding.
Peyer’s patches display typical reactive nodal histology. Occasional, typical lymphoid cluster (cryptopatches).
(85121, 85131, 85122)

Cecum/Colon/GALT

Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion cells of the plexuses.
Occasional, typical lymphoid cluster (Peyer’s patch).
Abundant luminal protozoa in the cecum and proximal colon - feature seen in the controls.
(85123)
No lesions of significance

Mesenteric lymph node

Highly reactive micromorphology including enlarged size, follicular hyperplasia with germinal centre formation, an expansive paracortical area, scattered lymphocyte apoptosis, and sinus histiocytosis.
(85124)
No lesions of significance

Spleen

Mild follicular hyperplasia with germinal centre formation and lymphocyte apoptosis. Mild coalescence of the white pulp.
Expansive red pulp with marked extramedullary haematopoiesis.
(85116-85119)
No lesions of significance

Pancreas

Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).
(85116-85119)
No lesions of significance

Kidney

Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable. Renal lymph nodes with typical reactive nodal histology.
(85120, 85125)
No lesions of significance

Adrenal glands

Adrenal glands with typical cortex/medulla micromorphology.
(85120, 85125)
No lesions of significance

Salivary glands and Regional lymph nodes

Mild multifocal perivascular mononuclear cell infiltrates within the submandibular gland, an age-related change - feature seen in the controls.
Unremarkable sublingual and parotid glands.
Regional lymph nodes display typical reactive nodal histology.
(85124)
No lesions of significance
Thyroids

Normal lateral lobes of the thyroid gland with typical colloid secreting follicles lined by cuboidal epithelium.
(85126, 85127)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, and blood vessels.
Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.
Mild multifocal perivascular mononuclear cell infiltrates in the surrounding periadipose tissue, common background finding - feature seen in the controls.
(85126, 85127)
No lesions of significance

Thymus

Typical medulla/cortex distribution and micromorphology.
Single small sized cyst (85216), common incidental/background finding - feature seen in the controls.
(85126, 85127)
No lesions of significance

Heart/chambers/vessels/valves

Representative cardiac muscle, chambers, valves and great vessels of the heart.
Some myxomatous valvular changes (thickened leaflets) (85126), an age-related change - feature seen in the controls.
The cardiac muscle fibres demonstrated typical features including central nuclei, branching fibres and striations.
(85126, 85127)
No lesions of significance

Skin

Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle (panniculus carnosus).
(85128)
No lesions of significance

Tail

Typical tail components including keratinized squamous epithelium, dense regular connective tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and blood vessels.
(85250, 85251)
No lesions of significance

Eyes/Harderian glands

Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
Typical branched tubuloalveolar formation of the Harderian gland.
Includes portion of unremarkable optic nerve and extraocular muscles.
(85129)
No lesions of significance

Brain

Sections were prepared from the standard levels of the brain:
Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third ventricles.
Level III hindbrain: includes the cerebellum, pons and fourth ventricle.

Sections of brain appear symmetrical with unremarkable meninges and typical lamination. The cerebellum appears symmetrical with typical architecture and Purkinje cells. There was no obvious neuronal loss and the myelination appears normal.

Prominent flattening/stretching of the ependymal cells and increased size of the lateral ventricular space, features indicative of increased cerebrospinal fluid (hydrocephalus), a known background lesion in mice - similar feature seen in the controls.

(85130)

Comments:
Neuropathology to comment

Spinal cord

Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc, striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.

(85253, 85254)
No lesions of significance

Comments:
Neuropathology to comment

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle

Mostly unremarkable long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow. The skeletal muscle shows consistent fibre size with peripheral nuclei.
Focal mixed leucocyte inflammation surrounding a tendon of the distal leg, likely incidental - similar feature seen in the controls.

(85250, 85251)

Comments:
Pathology to comment

Head

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85297). The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments (85298, 85299).
Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outer hair cells and typical tectorial membrane (85298, 85299).
The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85299).

(85297-85299)
No lesions of significance

Sternum

Representative sternabrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat.
Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85252)
Macro Observations

Tail suspension test for neurological defects - negative.
Dentition, tongue and oral cavity was unremarkable.
BCS: 3
Spleen: 14x5x3mm
Kidneys: not measured, symmetrical
Thymus: 8x8x2mm
Lungs inflated.
Heart: 9x7x5mm
Brain: 15x10x5mm, symmetrical
Pituitary gland identified, macroscopically normal
Tail length: 80mm (straight)
Head harvested for evaluation of auditory and vestibular structures.
Bone marrow smear taken from left hind leg.

No macroscopic lesions identified.

Micro Observations

Marrow smear: Examination of the smear showed representative cells from the myeloid and erythroid series. Occasional cells from the lymphoid series. Occasional and unremarkable megakaryoblasts.
(85132)

Peripheral blood smear: Examination of the smear showed red blood cells (majority of cells shown), occasional white blood cells including lymphocytes, segmented neutrophils, monocytes and platelets (clumps). No discernible morphological changes or detectable parasites.
(85133)

Mammary glands

Typical mammary fat pad with developing lactiferous ducts, blood vessels, and nerve bundles.
(85141, 85145)
No lesions of significance

Ovaries/Oviducts

Unremarkable ovaries containing follicles at various stages of development (primary through to antral) and several corpora lutea.
Few pigment laden interstitial cells (likely ceroid-lipofuscin), an age-related change - feature seen in the controls.
Mononuclear cell infiltration of the bursa membrane, bilateral, an age-related finding in mice.
Mostly unremarkable oviduct micromorphology with typical columnar epithelium and mucosal folds.
Focal epithelial cytoplasmic vacuolation, an age-related change - feature seen in the controls.
(85134, 85135, 85136)
No lesions of significance

Uterus/Cervix/Vagina/Clitoral gland

Cystic endometrial hyperplasia, an age-related change - feature seen in the controls.
Unremarkable architecture myometrium and adventitia.
The micromorphology of the uterus and vagina places the animal at estrus.
(85134, 85135, 85136)
No lesions of significance
Urinary Bladder

Unremarkable bladder with typical urothelium and detrusor muscle.
(85134, 85135, 85136)
No lesions of significance

Liver/Gall bladder

Liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Multiple small parenchymal leukocyte aggregates with some hepatocyte cell loss/necrosis - common incidental/background finding - feature seen in the controls.
Mild multifocal perivascular mononuclear cell infiltrates, an incidental/age-related change - feature seen in the controls.
Gall bladder with epithelial hyperplasia, cytoplasmic hyaline droplet accumulation, and underlying neutrophils in the lamina propria, common incidental/background finding - feature seen in the controls.
(85137)
No lesions of significance

Stomach

Unremarkable fore and glandular portions of the stomach with limiting ridge.
Includes pyloric sphincter and duodenal bulb.
(85138)
No lesions of significance

Small Intestine (Duodenum, Jejunum & Ileum)/GALT

Typical mucosal villi and submucosal layers.
Peyer's patches display typical reactive nodal histology. Occasional, typical lymphoid cluster (cryptopatches).
(85138, 85139, 85140)
No lesions of significance

Cecum/Colon/GALT

Typical mucosal folds and submucosal layers. Unremarkable muscularis and discernible ganglion cells of the plexuses.
Occasional, typical lymphoid cluster (Peyer's patch).
Clusters of luminal protozoa in the cecum - feature seen in the controls.
Increased mononuclear cell infiltrates in the lamina of the proximal colon, incidental - feature seen in the controls.
(85140)
No lesions of significance

Mesenteric lymph node

Typical reactive nodal histology with representative cortex including the occasional follicle, an expansive paracortical area with lymphocyte apoptosis, and sinus histiocytosis.
(85141)
No lesions of significance

Spleen

Mild follicular hyperplasia with germinal centre formation and some lymphocyte apoptosis. Mild coalescence of the white pulp.
Expansive red pulp with marked extramedullary haematopoiesis.
(85136)
No lesions of significance
Pancreas

Representative exocrine tissue (serous acini) and endocrine tissue (islets of Langerhans).
(85134, 85135, 85136)
No lesions of significance

Kidney

Sections show a cortex, medulla, and papilla. There is a uniform distribution of glomeruli and accompanying nephron components and the micromorphology of the tubules is unremarkable. Minimal perivascular/peripelvic mononuclear cell infiltrates, incidental/age-related change - feature seen in the controls.
Several mildly dilated protein casts within the medulla, incidental/age-related change - feature seen in the controls.
Renal lymph nodes with typical reactive nodal histology and mild sinus histiocytosis.
(85142)
No lesions of significance

Adrenal glands

Adrenal glands with typical cortex/medulla micromorphology.
(85142)
No lesions of significance

Salivary glands and Regional lymph nodes

Mild multifocal perivascular mononuclear cell infiltrates within the submandibular gland, an age-related change - feature seen in the controls.
Unremarkable sublingual and parotid glands.
Regional lymph nodes display typical reactive nodal histology.
(85141)
No lesions of significance

Thyroids

Normal lateral lobes of the thyroid gland with typical colloid secreting follicles lined by cuboidal epithelium.
(85143, 85144)
No lesions of significance

Trachea/Lungs

Typical lung parenchyma/alveoli, bronchioles, and blood vessels.
Focal parenchymal congestion and collapse judged to be artefactual.
Mild multifocal perivascular mononuclear cell infiltrates, an incidental/age-related change - feature seen in the controls.
Trachea with unremarkable mucosal epithelial lining and hyaline cartilage.
Oesophagus with typical features including stratified squamous epithelium.
(85143, 85144)
No lesions of significance

Thymus

Typical medulla/cortex distribution and micromorphology.
(85143, 85144)
No lesions of significance

Heart/chambers/vessels/valves

Mostly typical micromorphology observed in cardiac muscle, chambers, valves and great vessels of the heart.
Myxomatous valvular changes (thickened leaflets), an age-related change - feature seen in the controls.
The cardiac muscle fibres demonstrated typical features including central nuclei, branching fibres and striations.
Focal basophilic deposit in the ventricular muscle resembling mineralisation, likely incidental.
(85143, 85144)

Comments:
Pathology to comment

Skin
Typical dermal appendages and distribution. Unremarkable thin layer of striated muscle (panniculus carnosus).
(85145)
No lesions of significance

Tail
Typical tail components including keratinized squamous epithelium, dense regular connective tissue, tendons, caudal vertebra, bone marrow, intervertebral disc, skeletal muscle, nerves and blood vessels.
(85224, 85225, 85226)
No lesions of significance

Eyes/Harderian glands
Unremarkable retina, cornea, iris, ciliary body, lens, sclera and choroid.
Typical branched tubuloalveolar formation of the Harderian gland.
Minimal multifocal glandular mononuclear cell infiltrates - an incidental/age-related finding in mice.
Includes portion of unremarkable optic nerve and extraocular muscles.
(85146)
No lesions of significance

Brain
Sections were prepared from the standard levels of the brain:
Level I forebrain: including cortex, corpus callosum, caudate putamen and lateral ventricles.
Level II midbrain: including the hippocampus, thalamus, hypothalamus and lateral and third ventricles.
Level III hindbrain: includes the cerebellum, pons and fourth ventricle.

Sections of brain appear symmetrical with unremarkable meninges and typical lamination. The cerebellum appears symmetrical with typical architecture and Purkinje cells. There was no obvious neuronal loss and the myelination appears normal.

Prominent flattening/stretching of the ependymal cells and increased size of the lateral ventricular space, features indicative of increased cerebrospinal fluid (hydrocephalus), a known background lesion in mice - similar feature seen in the controls.
Query vacuolation of the adjacent white matter, possible artefact.
(85147)

Comments:
Neuropathology to comment

Spinal cord
Representative thoracic and lumbar region of spinal cord, vertebral bone, intervertebral disc, striated muscle, peripheral nerves, brown adipose tissue, and bone marrow.
(85227, 85228, 85229)
No lesions of significance

Comments:
Neuropathology to comment

(Hind leg) Long bone/Bone marrow/Synovial joint/Skeletal muscle

Unremarkable long bone, striated muscle, examples of nerve fascicles, fibrocartilage of the meniscus, synovial joint and bone marrow. The skeletal muscle shows consistent fibre size with peripheral nuclei.

(85224, 85225, 85226)
No lesions of significance

Head

Multiple levels through the head demonstrate dermal appendages, nasal cavity, oral cavity, teeth and tongue including muscle bundles. Sections also show unremarkable pituitary gland including pars intermedia, pars distalis and pars nervosa as well and the trigeminal nerve/ganglia (85300). The outer and middle regions of the ear are discernible. The tympanic membrane is intact and the ossicles are unremarkable and include the stapedial annular ligaments (85301, 85302). Typical components of the inner ear including bony labyrinth, organ of corti, stria vascularis and scala cavities are discernible. Based on multiple levels, the organ of corti is unremarkable with no discernible loss of inner/outer hair cells and typical tectorial membrane (85301, 85302). The cochlear nerve and spiral ganglion is also demonstrated and based on several levels, there is no reduction in the density of the spiral ganglion cells. Examples of otolith organs can be seen with typical features such as the hair cells and mineral otoliths. The ampulla including the crista ridge with hair cells is discernible (85302).

Segmental hyaline droplet accumulation within the nasal epithelium, incidental - feature seen in the controls.

(85300-85302)
No lesions of significance

Sternum

Representative sternebrae, costal cartilage, intersternebral joint, intercostal skeletal muscle and brown fat. Haematopoietic tissue islands surrounded by vascular sinuses interspersed within a meshwork of trabecular bone. The bone marrow morphology demonstrated typical myeloid features including conspicuous megakaryoblasts and lymphoid features.

(85230)
No lesions of significance

Comment / Plan

Case APN21/036SVI (C. Walkley) will be referred to Dr. John Finnie, University of Adelaide for comment.

Aira Nuguid
18th October, 2021

Supplementary Pathology Report

#20 (control)

85203 – Pancreas – focal necrosis of exocrine pancreas with lymphocytic infiltration.

85207 – Heart – cardiomyocyte degeneration with cytoplasmic vacuolation.


85281 – Hind leg – focal skeletal myocyte degeneration and necrosis.
#5 (control)
85088 – Liver – multifocal hepatocellular necrosis with mononuclear cell reaction, likely bacterial, and periportal lymphocytic infiltration, both common incidental findings.

85089 – Pancreas – focal necrosis of exocrine pancreas with lymphocytic infiltration.

85094 – Heart – multifocal cardiomyocyte degeneration and necrosis and lymphocytic infiltration.

#6 (control)
85107 – Salivary gland – acinar atrophy.

85295 – Head/external ear – focal coagulation necrosis of lining epithelium.

85234 – Hind leg – marked subcutaneous oedema (anasarca).

85235 – Hind leg – subcutaneous degeneration with mast cell hyperplasia and eosinophil infiltration (? allergic aetiology).

#11
85152 – Liver – mild diffuse cloudy swelling and hydropic degeneration of hepatocytes.

85269 – Hind leg – focal skeletal myocyte degeneration and necrosis.

#12
85167 – Liver – mild diffuse cloudy swelling and hydropic degeneration of hepatocytes.

#18
85273 – Hind leg – necrotic osteolysis (? traumatic aetiology).

#3
85061 – Salivary gland – large focus of acute acinar necrosis

85063 – Heart – multifocal cardiomyocyte degeneration and necrosis.

85237 – Hind leg – focal skeletal myocyte degeneration and regeneration.

85239 – Hind leg – focal coagulation necrosis of skeletal myocytes.

85243 – Skin – acute focal epidermal ulceration and degeneration and necrosis of panniculus muscle.

#4
85073 – Liver – multifocal hepatocellular necrosis with mononuclear cell reaction, likely bacterial, and periportal lymphocytic infiltration, both common incidental findings.

85079 – Heart – multifocal cardiomyocyte degeneration and necrosis.

85291 – Head/middle ear – luminal macrophages and multinucleated giant cells with cholesterol clefts in wall.

85257 – Hind leg – skeletal myocyte degeneration and necrosis.
#7
85120 – Liver – multifocal hepatocellular necrosis with mononuclear cell reaction, likely bacterial, and periportal lymphocytic infiltration, both common incidental findings.
85120 – Liver – focal micro and macro vesicular steatosis of hepatocytes.
85251 – Hind leg – apparent cellulitis with mixed inflammatory cell infiltration.

#8
85144 – Heart – focal mineralization.

Summary
Please see individual animals in the above report for comments.

Note:
NAD = No abnormalities detected.

Pathologist name and details removed CW230222
27th October, 2021

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Supplementary Neuropathology Report

#20 (control)
85211 – Brain – NAD.
85285 – Spinal cord – NAD.

#5 (control)
85098 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85246 – Spinal cord – NAD.

#6 (control)
85113 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85232 – Spinal cord – NAD.

#11
85162 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85272 – Spinal cord – NAD.

#12
85177 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85266 – Spinal cord – NAD.
#18
85194 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85278 – Spinal cord – NAD.

#3
85067 – Brain – variable dilation of lateral ventricles (internal hydrocephalus) and oedema of surrounding white matter tracts.
85343 – Spinal cord – NAD.

#4
85083 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85260 – Spinal cord – NAD.

#7
85130 – Brain – variable dilation of lateral ventricles (internal hydrocephalus).
85254 – Spinal cord – NAD.

#8
85147 – Brain – variable dilation of lateral ventricles (internal hydrocephalus) and oedema of surrounding white matter tracts.
85229 – Spinal cord – NAD.

Summary
Brains – #4, #5, #6, #7, #11, #12 and #18 – variable dilation of lateral ventricles (internal hydrocephalus).
Brains – #3, #8 – variable dilation of lateral ventricles (internal hydrocephalus) and oedema of surrounding white matter tracts.

Sections of the spinal cord show no significant findings.

Note:
NAD = No abnormalities detected.

(Pathologist name and contact details removed_CW23022)
27th October, 2021
Phenomics Australia advises all research groups that images or results obtained are to be acknowledged in resultant publications. Example acknowledgement: "This study utilised Phenomics Australia Histopathology and Slide Scanning Service, University of Melbourne".

Authorised by (name removed_CW230222), Phenomics Australia and Slide Scanning Service manager.
APN21/036SVI
Macro Images
Carl Walkley

Male #3
Adar1P195A/E861A

Female #4
Adar1P195A/E861A

#3 MUT
Adar1P195A/E861A, Male, 7 months old

#4
Adar1P195A/E861A, Female, 7 months old
APN21/036SVI
Macro Images
Carl Walkley

#5 WT
Adar1P195A/E861A, Female, 7 months old

#6 WT
Adar1P195A/E861A, Female, 7 months old
Macro Images

Female #7
Adar1P195A/E861A, Female, 7 months old

Female #8
Adar1P195A/E861A, Female, 7 months old
APN21/036SVI
Macro Images
Carl Walkley

Male #18
Adar1P195A/E861A, Male, 6 months old

#18 MUT
Adar1P195A/E861A, Male, 6 months old

Male #20
WT

#20 WT
Adar1P195A/E861A, Male, 6 months old
Images are of animal #3 MUT, Adar1P195A/E861A, Male, 7 months old.
Images are of animal #4 MUT, Adar1P195A/E861A, Female, 7 months old
Images are of animal #5 WT, Adar1P195A/E861A, Female, 7 months old
Images are of animal #6 WT, Adar1P195A/E861A, Female, 7 months old
Images are of animal #7 MUT, Adar1P195A/E861A, Female, 7 months old
Images are of animal #8 MUT, Adar1P195A/E861A, Female, 7 months old
Images are of animal #11 MUT, Adar1P195A/E861A, Female, 7 months old.
Images are of animal #12 MUT, Adar1P195A/E861A, Female, 7 months old.
Images are of animal #18 MUT, Adar1P195A/E861A, Male, 6 months old
Images are of animal #20 WT, Adar1P195A/E861A, Male, 6 months old
#5 WT Female Ovary 63x 85087
Pigmented interstitial cells

#5 WT Female Oviduct 10x 85087
Non-vacuolated (A), vacuolated (B)

#6 WT Female Uterus 3x 85102
Cystic endometrial hyperplasia

#5 WT Female Liver 10x 85088
Mononuclear cell infiltrate

#5 WT Female Liver 40x 85088
Hepatocyte cell loss/necrosis and infiltrates (A)

#5 WT Female Gall bladder 63x 85088
Epithelial hyperplasia, and neutrophils (A)
#5 WT Female Large intestine 20x 85091
Cecum - luminal protozoa (A)

#5 WT Female Pancreas 40x 85087
Mononuclear cell infiltrate

#5 WT Female Pancreas 20x 85089
Acini necrosis (A), normal acini (B)

#5 WT Female Kidney 20x 85093
Protein cast

#5 WT Female Kidney 40x 85093
Mononuclear cell infiltrate

#5 WT Female Salivary gland 20x 85092
Mononuclear cell infiltrate
#5 WT Female Thymus 40x 85094

Cyst

#5 WT Female Heart 40x 85094

Cardiomyocyte degeneration

For comparison

#5 WT Female Heart 40x 85094

Myxomatous changes (thickened leaflet) (A)

#5 WT Female Spinal cord 40x 85245

Connective tissue - eosinophils

#5 WT Female Head 20x 85292

Mammary tissue - mononuclear cell infiltrate (A)
#5 WT Female Head 40x 85292
Neck muscle - leukocyte infiltrate (A)

#6 WT Female Salivary gland 63x 85107
Enlarged and basophilic acini

#6 WT Female Salivary gland 20x 85107
Normal (A), Atrophic (B)

#6 WT Female Lung 30x 85110
Mononuclear cell infiltrate

#6 WT Female Hind leg 20x 85235
Foot - inflammation (A)

#5 WT Female Hind leg 20x 85247
For comparison
#6 WT Female Hind leg 10x 85235
Subcutaneous oedema (A)

#5 WT Female Hind leg 10x 85247
For comparison

#20 WT Male Head 63x 85313
Nasal epithelium - hyaline droplet accumulation

#6 WT Female Head 20x 85295
Tympanic membrane - inflammation/necrosis (A)

#5 WT Female Head 20x 85293
Tympanic membrane - for comparison

#20 WT Male Testis 20x 85198
Tubular degeneration/atrophy (A), normal (B)
#20 WT Male Pancreas 15x 85203
Necrosis and infiltrates (A), normal acini (B)

#20 WT Male Eye 40x 85210
Cornea - epithelial cell loss

For comparison

#20 WT Male Hind leg 40x 85280
Myodegeneration

For comparison

#20 WT Male Hind leg 63x 85280
Myofibre necrosis (A)
#20 WT Male Hind leg 63x 85281
Myofibre regeneration

#3 MUT Male Spinal cord 10x 85243
Skin - ulcer (A), inflammation (B)

#7 MUT Female Ileum 20x 85123
Eosinophilic infiltrates (transmural)

#20 WT Male Head 30x 85314
Hair follicle inflammation (A), normal follicle (B)

#4 MUT Female Ovary 5x 85072
Cyst (A)

#6 WT Female Ileum 20x 85106
For comparison
#8 MUT Female Harderian gland 40x 85146
Mononuclear cell infiltrate

#8 MUT Female Heart 63x 85144
Basophilic deposit

#8 MUT Female Ovary 10x 85136
Bursa membrane - mononuclear cell infiltrate

For comparison

#5 WT Female Ovary 10x 85087

#11 MUT Female Liver 63x 85152
Hydropic degeneration

#5 WT Female Liver 63x 85088
For comparison
#11 MUT Female Head 20x 85304  
Nasal cavity - inflammation

#6 WT Female Head 20x 85296  
For comparison

#18 MUT Male Hind leg 20x 85273  
Foot - inflammation and necrosis

#6 WT Female Hind leg 20x 85236  
Foot - for comparison

#8 MUT Female Brain 5x 85147  
Increased extracellular space (lateral ventricle) (A)

#20 WT Male Brain 5x 85211  
For comparison
# 9.1 Histopathology Report

<table>
<thead>
<tr>
<th>Case Number</th>
<th>Registration Date</th>
<th>Animal Details</th>
<th>Strain</th>
</tr>
</thead>
<tbody>
<tr>
<td>APN22/016 St Vincent’s Institute of Medical Research (Carl Walkley)</td>
<td>Wed 09/03/2022</td>
<td>409, WT Ly5.1C57-CW DOB: null, 17.9 weeks, Male, nullg</td>
<td>C57BL/6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>410, WT Ly5.1C57-CW DOB: null, 17.9 weeks, Male, nullg</td>
<td>C57BL/6</td>
</tr>
<tr>
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<td>C57BL/6</td>
</tr>
<tr>
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<td></td>
<td>104, Adar1P195A/E861A DOB: null, 14.4 weeks, Male, nullg</td>
<td>C57BL/6</td>
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<tr>
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<td>124, Adar1P195A/- DOB: null, 17 weeks, Male, nullg</td>
<td>C57BL/6</td>
</tr>
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<td>125, Adar1P195A/- DOB: null, 17 weeks, Male, nullg</td>
<td>C57BL/6</td>
</tr>
<tr>
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<td>126, Adar1P195A/- DOB: null, 17 weeks, Male, nullg</td>
<td>C57BL/6</td>
</tr>
<tr>
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<td>129, Adar1P195A/- DOB: null, 17 weeks, Female, nullg</td>
<td>C57BL/6</td>
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<tr>
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<td>C57BL/6</td>
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<td>C57BL/6</td>
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<tr>
<td></td>
<td></td>
<td>72, Adar1P195A/- Ifih1/- DOB: null, 16.4 weeks, Male, nullg</td>
<td>C57BL/6</td>
</tr>
</tbody>
</table>
ADAR1 P195A/+  
97, Adar1P195A/E861A  
DOB: null, 18 weeks, Female, nullg  
Strain: C57BL/6

DoD / Necropsy  
Necropsy not conducted at P.A

Death  
Unknown

Origin  
St Vincent’s Institute of Medical Research

Treatment  
Knock-out  
Tissues from each animal - hemisphere of brain, piece of liver, spleen and whole kidney  
Collected and fixed overnight in 2% PFA, then transferred to 70% ethanol for storage.

Adar1 P195A mutants.  
Adar1 P915A is the murine homologue of ADAR1 P193A which is the most common human mutation of ADAR1.

Please embed, section and assess slides.  
General histology and any pathology observed of interest. Humans with ADAR mutation have neuronal calcifications but we didn’t see that in the whole animal histology samples previously assessed (Adar1E861A/P195A) at APN.

Previously reported that P195A mice had (PMID: 34343497): In our initial assessment, the clearest histological defects were found in the kidney and liver. The kidney exhibited glomerular mesangial matrix expansion that increased with age, and the liver exhibited extensive microvesicular cytoplasmic vacuolation (Figure 3A–3C). We found no substantial inflammatory immune cell infiltrates in these tissues (Figures 3A–3C). We additionally identified abnormal architecture of the spleens in AdarP195A/p150 mice, characterized by lymphoid depletion. Note our P195A line is not displaying the early lethality reported in the paper above that reports the kidney and liver pathology.

C. Walkley 8/03/2022

Species / Breed / Strain  
C57BL/6; Adar1 P195A/- strain

Animal Health Facility  
St. Vincent's Bioresources Centre

N/A

Organs Examined  
Brain, Kidney, Liver, Spleen

Macroscopic Observations  
Courier/Transportation details: C. Walkley 9/03/2022  
Number of animals/samples: 13 mice  4 samples per mouse (except mouse 410 with only 3 samples-no spleen included) = 51 fixed samples  
Sample details: Tissues from each animal - hemisphere of brain, piece of liver, spleen and whole kidney.  
Collected and fixed overnight in 2% PFA, then transferred to 70% ethanol for storage C. Walkley 8/03/2022

PA iLab request number: PAHSSS-CW-74

REFERENCES:
(1) National Toxicology Program-Nonneoplastic Lesion Atlas  
https://ntp.niehs.nih.gov  
(3) Histology of the Central Nervous System, R.H. Garman, Toxicologic
Pathology 2009
(4) Proliferative and Nonproliferative Lesions of the Rat and Mouse Hepatobiliary System, B. Thoolen et al, 2010

NOTES
(1) The clear spaces in the cytoplasm of the hepatocyte represent sites of glycogen that has been dissolved in aqueous fixative. Glycogen deposition does not usually displace the hepatocyte nucleus. This is a typical histological feature in normal, non fasted mice and should not be confused with fatty change.
(2) EMH consists of erythroid precursors, myeloid precursors, megakaryocytes or all three. While some degree of extramedullary haematopoiesis is present in normal rodents, especially in mice, increased extramedullary haematopoiesis can result from haematotoxic insult, systemic anaemia, and infections elsewhere in the body. A.W Suttie, 2006
(3) The presence of a small amount of intracytoplasmic pigment within the splenic red pulp macrophages is a common background finding in rodents. Pigment can be hemosiderin, ceroid/lipofuscin, melanin, or test article related. It is typically found in the red pulp but may also be present in the marginal zone and/or periarteriolar lymphatic sheaths of the white pulp, the capsule, or trabeculae.

Microscopic Observations

SUMMARY

70 Brain (87505)-cellularity/lamination of cerebral cortex or plane of section. Kidney (87508)- Tubule protein cast in medulla.

71 kidney (87504)-interstitial inflammatory cell infiltrates

72 Kidney (87500)- inflammatory cell infiltrate (lymphocytic)

97 NLS Liver(87495)-Mild multifocal inflammatory cell aggregate. Brain (87493)-NLS

104 NLS Brain (87489)-NLS

124 NLS Brain(87481)-NLS

125 NLS Brain(87477)-NLS

126 NLS Spleen (87463)-query less white pulp. Likely to be plane of section. Brain (87461, 87465)-NLS

129 Brain (87473)-cellularity/lamination of cerebral cortex, likely to be plane of section. Liver (87474)- inflammatory cell infiltrate. Spleen(87475)-moderate EMH

131 Liver (87487)-periportal inflammatory cell aggregates with hepatocyte loss. Brain -NLS (87485)

409 NLS Brain (87469)-NLS

410 NLS Brain (87466)-NLS

418 NLS Brain (87457)-NLS

NLS= No lesions of significance
409 (control)
Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 11x6x4mm
No macroscopic lesions identified.

Micro Observations

WT Ly5.1C57-CW

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins. A small aggregation of Kupffer cells with trivial hepatocyte loss.
No lesions of significance
(87470)

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology.
No lesions of significance
(87471)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance
(87472)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, olfactory lobe, cerebral cortex with typical lamination, corpus callosum, ventricles and choroid plexi, hippocampus, thalamus, hypothalamus and hind brain (cerebellum, pons). Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. Readily identified dark neuron fixative artifact, a common feature.
No lesions of significance
(87469)

410 (control)
Macro Observations

Fixed liver, brain & kidney
Kidney 11x6x4mm
No macroscopic lesions identified.

Micro Observations

WT Ly5.1C57-CW

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins. Unremarkable Gall bladder
No lesions of significance
(87467)
Kidney

H&E section show cortex, medulla and papilla regions (and ureter) of the kidney. Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli. The interstitium and tubules is unremarkable. No lesions of significance (87468)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, small portion of the olfactory lobe, cerebral cortex with typical lamination, corpus callosum, ventricles and choroid plexi, hippocampus, thalamus, hypothalamus and hind brain (cerebellum). Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. Readily identified dark neuron fixative artifact, a common feature. No lesions of significance (87466)

418 (control)

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 9x5x3mm
No macroscopic lesions identified.

Micro Observations

WT Ly5.1C57-CW

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins. A small aggregation of Kupffer cells with trivial hepatocyte loss. No lesions of significance (87460)

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology. No lesions of significance (87458)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney. Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli. The interstitium and tubules is unremarkable. No lesions of significance (87459)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, olfactory lobe, cerebral cortex with typical lamination, corpus callosum, ventricles and choroid plexi, hippocampus, thalamus, hypothalamus and hind brain (cerebellum, pons). Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. Readily identified dark neuron fixative artifact, a common feature. Section includes small portion of exocrine tissue (pancreas) judged to be cross contamination. No lesions of significance (87457)
104

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 12x7x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/E861A

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
No lesions of significance
(87491)

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology. Mild extramedullary haematopoiesis identified in the red pulp, a common feature in the mouse spleen.
No lesions of significance
(87490)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney. Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli. The interstitium and tubules is unremarkable.
No lesions of significance
(87492)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and representative choroid plexus, hippocampus, subfornical organ, thalamus, hypothalamus and cerebellum. Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal.
No lesions of significance
(87489)

124

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 13x7x5mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/-

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
No lesions of significance
(87482)
Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology.
Mild extramedullary haematopoiesis identified in the red pulp as well as brown pigment, likely to be haemosiderin.
No lesions of significance
(87483)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance
(87484)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, small portion of the olfactory lobe, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and choroid plexi, hippocampus, thalamus, hypothalamus and hind brain (cerebellum). Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. Readily identified dark neuron fixative artifact, a common feature.
No lesions of significance
(87481)

125

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 13x8x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/-

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
No lesions of significance
(87479)

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology.
Moderate extramedullary haematopoiesis identified in the red pulp as well as brown pigment, likely to be haemosiderin.
No lesions of significance
(87478)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance
(87480)
Brain Slide shows sagittal sections of one hemisphere demonstrating, meninges, small portion of the olfactory lobe, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and choroid plexi, hippocampus, thalamus, hypothalamus and hind brain (cerebellum, pons). Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. Readily identified dark neuron fixative artifact, a common feature.
No lesions of significance

(87477)

126

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 13x7x5mm -mechanical artefact
No macroscopic lesions identified.

Micro Observations

Adar1P195A/-

Liver
Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
No lesions of significance

(87462)

Spleen
Discernible red and white pulp micromorphology.
Less white pulp, judged to be plane of section.
Mild extramedullary haematopoiesis identified in the red pulp.

(87463)

Comments:
Pathology to comment

Kidney
H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman’s capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance

(87464)

Brain
Section shows several pieces of non-intact brain. Limited evaluation due to mechanical artefact.
Discernible and unremarkable cerebellum and olfactory bulb region.
No observable lesions. Cellularity and lamination appears unremarkable.

(87461, 87465)

Comments:
Neuropathology to comment
Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 11x6x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/-

Liver

Section shows liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Mild, multifocal inflammatory cell aggregate with some hepatocyte loss.
(87474)

Comments:
Pathology to comment

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology. Moderate extramedullary haematopoeisis identified in the red pulp as well as brown pigment, likely to be haemosiderin.
(87475)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance
(87476)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, olfactory lobe, cerebral cortex, corpus callosum, ventricles and choroid plexi, thalamus, hypothalamus and hind brain (cerebellum, pons). Unremarkable cerebellum with typical architecture and Purkinje cells.
Irregular cellular lamination of the cerebral cortex, judged to be plane of section.
(87473)

Comments:
Neuropathology to comment

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, olfactory lobe, cerebral cortex, corpus callosum, ventricles and choroid plexi, thalamus, hypothalamus and hind brain (cerebellum, pons). Unremarkable cerebellum with typical architecture and Purkinje cells.
Irregular cellular lamination of the cerebral cortex, judged to be plane of section.
(87473)

Comments:
Neuropathology to comment

131

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 12x6x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/-
Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Mild, multifocal centrilobular and periportal inflammatory cell aggregate with some hepatocyte loss.
(87487)

Comments:
Pathology to comment

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology.
Mild extramedullary haematopoiesis identified in the red pulp, a common feature in the mouse spleen.
No lesions of significance
(87486)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance
(87488)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, small portion of the olfactory lobe, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and representative choroid plexus, hippocampus, subfornical organ, thalamus, hypothalamus, cerebellum and pons. Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. Readily identified dark neuron fixative artifact, a common feature.
No lesions of significance
(87485)

70

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 10x5x3mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/- Lfh1-/-

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Unremarkable Gall bladder.
No lesions of significance
(87507)
Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology. Mild extramedullary haematopoiesis identified in the red pulp, a common feature in the mouse spleen. No lesions of significance (87506)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney. Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman's capsule is apparent in many glomeruli. Tubule protein cast in medulla. The remaining interstitium and tubules is unremarkable. (87508)

Comments:
Pathology to comment

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, cerebral cortex, corpus callosum, ventricles and representative choroid plexus, subfornical organ, hippocampus, thalamus, hypothalamus and cerebellum. Query cellularity/lamination of cerebral cortex or plane of section. Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal. (87505)

Comments:
Neuropathology to comment

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 11x6x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/- Ifih1/-

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins. No lesions of significance (87503)

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology. Mild extramedullary haematopoiesis identified in the red pulp, a common feature in the mouse spleen. No lesions of significance (87502)
Kidney

H&E section show cortex, medulla and papilla regions of the kidney. Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman’s capsule is apparent in many glomeruli. Mild, multifocal interstitial inflammatory cell infiltrate. The remaining interstitium and tubules is unremarkable. (87504)

Comments:
Pathology to comment

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and representative choroid plexus, subfornical organ, thalamus, hypothalamus, cerebellum and pons. Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal.
No lesions of significance (87501)

72

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 10x6x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/- Ifih1-/-

Liver

Section shows typical liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
No lesions of significance (87499)

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology. Mild extramedullary haematopoiesis identified in the red pulp, a common feature in the mouse spleen.
Section also includes a small portion of unremarkable pancreatic exocrine tissue.
No lesions of significance (87498)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney. Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman’s capsule is apparent in many glomeruli. Focal inflammatory cell aggregate (lymphocytic).
The remaining interstitium and tubules is unremarkable. (87500)

Comments:
Pathology to comment
Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and representative choroid plexus, hippocampus, subfornical organ, thalamus, hypothalamus, cerebellum and pons. Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal.
No lesions of significance
(87497)

Macro Observations

Fixed spleen, liver, brain & kidney
Kidney 10x5x4mm
No macroscopic lesions identified.

Micro Observations

Adar1P195A/E861A

Liver

Section shows liver parenchyma including hepatocytes, Kupffer cells, portal triads and central veins.
Mild, multifocal inflammatory cell aggregate, resembling extramedullary haematopoiesis.
(87495)

Comments:
Pathology to comment

Spleen

Unremarkable follicular pattern identified with typical red and white pulp micromorphology.
Mild extramedullary haematopoiesis identified in the red pulp, a common feature in the mouse spleen.
No lesions of significance
(87494)

Kidney

H&E section show cortex, medulla and papilla regions of the kidney.
Typical glomeruli, many with open capillary lumens. The expected cuboidal epithelium lining the Bowman’s capsule is apparent in many glomeruli.
The interstitium and tubules is unremarkable.
No lesions of significance
(87496)

Brain

Slide shows sagittal sections of one hemisphere demonstrating, meninges, portion of olfactory lobe, cerebral cortex with typical cellular lamination, corpus callosum, ventricles and representative choroid plexus, hippocampus, subfornical organ, thalamus, hypothalamus and cerebellum. Unremarkable cerebellum with typical architecture and Purkinje cells. No obvious neuronal loss, and the myelination appears normal.
No lesions of significance
(87493)

Comment / Plan

Case APN22/016SVI(C. Walkley) to has been referred to text removed CW090522
Supplementary Pathology Report

**409 (control)**
No abnormalities detected

**410 (control)**
No abnormalities detected

**418 (control)**
No abnormalities detected

**104**
No abnormalities detected

**124**
No abnormalities detected

**125**
No abnormalities detected

**126**
126(87463)- spleen- normal follicular pattern (unusual plane of section)
No other abnormalities detected

**129**
129(87475)- spleen- normal follicular pattern (unusual plane of section)
129(87474)-liver- focal hepatocellular necrosis with PMN and Kupffer cell reaction (common incidental finding); accentuation of acinar pattern due to mild hydropic degeneration in periacinar and mid zonal hepatocytes.
No other abnormalities detected.

**131**
131(87487)-liver- focal hepatocellular necrosis with PMN and Kupffer cell reaction (common incidental finding); accentuation of acinar pattern due to mild hydropic degeneration in periacinar and mid zonal hepatocytes.
No other abnormalities detected
70
70(87508) kidneys - tubular protein casts (common incidental finding)

No other abnormalities detected

71
71(87504) kidneys - Glomerular tuft hypercellular with basement membrane thickening and capsular thickening. Interstitial mononuclear cell infiltrate with mesenchymal-like cells with hyperchromatic, elongated nuclei.
Tubular protein casts (common incidental finding)

No other abnormalities detected

72
72(87500) kidneys - single periarteriolar mononuclear cell cuff.

No other abnormalities detected

97
97(87495)-liver- focal hepatocellular necrosis with PMN and Kupffer cell reaction (common incidental finding); accentuation of acinar pattern due to mild hydropic degeneration in periacinar and mid zonal hepatocytes

No other abnormalities detected

Summary
Please see individual samples in the above report for comments.

PMN=Polymorphonuclear leukocytes
text removed CW090522

30th March, 2022

Supplementary Neuropathology Report

409 (control)
No abnormalities detected

410 (control)
No abnormalities detected

418 (control)
No abnormalities detected
No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

No abnormalities detected

Summary

No abnormalities detected. Cerebrum; often much mechanical damage in the form of numerous dark neurons; cerebellum appeared normal

Brain-The samples presented in this case bear no lesions of significance.

text removed CW090522

30th March, 2022
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