RESEARCH PROGRESS REPORT SUMMARY

Grant 02157-MOU: Genomics of Deafness in the Dalmatian

Principal Investigator: Claire Wade, PhD
Research Institution: University of Sydney
Grant Amount: $120,960.00
Start Date: 1/1/2015 End Date: 12/31/2018
Progress Report: Mid-Year 4
Report Due: 6/30/2018 Report Received: 6/17/2018

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Original Project Description:

Congenital deafness is a health issue that has higher prevalence in certain breeds, including the Dalmatian. Other studies in this breed have found the trait to be inherited in a complex rather than simple Mendelian manner. Using a large number of samples from animals that have been tested for hearing status, Dr. Wade will employ the latest genomic technologies and computational analyses to conduct this study. The ultimate goal is to identify mutations underlying the trait of congenital deafness in the Dalmatian breed and work towards a genetic testing solution for the Dalmatian breeding community.

Funding for the research is provided through the efforts and generosity of the Dalmatian Club of America Foundation. The AKC Canine Health Foundation supports the funding of this effort and will oversee administration of funds and scientific progress reports.

Publications: None at this time.

Presentations:

Honors student Lucy Armstrong has presented the results of her analysis at internal student forums.

We presented a poster on the project at the International conference on Canine and Feline genomics in Minnesota USA in July 2017.

Simone Carter: Honors presentation "Association of external phenotype with deafness risk loci in the Dalmatian" - this presentation was an outline of the work to be conducted for her honors research project.
Report to Grant Sponsor from Investigator:

Our original analysis assaying 170,000 genetic markers in 9 deaf and 24 bilateral hearing dogs identified five candidate regions that showed complete or near concordant association in our whole-genome sequenced (5 bilateral deaf and 2 hearing) Dalmatian samples. In the past year we obtained deep sequence from a 6th bilateral deaf animal to enable superior access to certain parts of the DNA that were not sequenced very well in the earlier technology.

Throughout the study we have continued to collect samples for breed risk analysis via our neurology clinic run by Dr. Georgina Child. Throughout the course of the project we have obtained samples of 37 bilateral deaf, 55 unilateral and 164 hearing pups. These numbers are not representative of population risk frequencies as our sampling is biased to collection of samples of all phenotypes from litters that contain bilateral deaf or unilateral deaf puppies. The majority of the samples collected have been assayed using canine genotyping arrays.

Through 2017 we deeply investigated two of the most promising loci. These loci were chosen because they had been previously implicated in work by other research groups (chromosome 10 and chromosome 20). We believe that we have identified a protective haplotype for bilateral deafness at one of the original loci under investigation (the white markings locus on chromosome 20). We are unprepared to release a test at this time as we are yet to ascertain the impact (if any) that this variant has on the external appearance of the dog. This question will be the subject of a student project to run throughout 2018. For this student project, we have invited dogs to a judging critique by an Australian National Kennel Council judge licensed to judge Dalmatians and suggested by the Dalmatian club of NSW. The judge will not know the genotypes of the dogs. At the end of the assessments, we will compare the genotyped groups of dogs for their critique. Our null hypothesis is that the low-risk variant does not affect external appearance. If this is true, we will have greater confidence in releasing access to the test for the low-risk variant. This component of the work is currently underway. In 2018 we attended the Dalmatian Club of New South Wales specialty show where we collected DNA and judge’s critiques of external phenotype from 3 ANKC judges and 40 dogs. We have also received genotyping arrays on a further 5 dogs from these dogs. The five new dogs are not phenotyped for deafness by us.

The assay developed by Jenny Liu is not working reliably in the laboratory so we have moved to a marker based assay for the protective haplotype. It must be made very clear that deafness in this breed is unlikely to be caused by a single locus, and indeed there are some bilateral deaf dogs (4 of 29 tested thus far in our data) that have one copy of the low-risk variant but just one that has two low-risk genes. Further, with our increasing sample sizes we have identified two new loci that warrant further investigation into the inheritance of this complex trait. That investigation will form the basis of a study to follow.