Evaluation of Cognitive Function in the Dog Aging Project: Associations with Baseline Canine Characteristics

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Abstract

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Background Canine Cognitive Dysfunction (CCD) is a neurodegenerative disease in aging canines. It has been described previously in relatively small cohorts of dogs using multiple different rating scales. This study aimed to use a minimally modified CCD rating scale developed by previous researchers to describe the prevalence of CCD more thoroughly in a large, nationwide cohort of companion dogs known as the Dog Aging Project (DAP). Specifically, associations between various canine characteristics, predicted life stage quartiles, and CCD were examined.

Methods This study used baseline data collected from October 30, 2020 to December 31, 2020 from 15,019 dogs enrolled in the DAP, a nationwide longitudinal study on canine aging and mortality in US companion dogs. Two surveys, the Canine Social and Learned Behavior Survey (CSLB) and the Health and Life Experience Survey (HLES), which provided owner-reported information on various canine characteristics and classified dogs as having probable CCD, were used. Associations were assessed using univariable and multivariable logistic regression
models adjusted for age, sterilization status, history of 15 categories of health problems, breed type, and activity level. Associations between predicted life stage quartile and CCD were also assessed, and life stage quartile was evaluated for its predictive ability through adjusted and unadjusted logistic regression models. Predictive capacity of these models was quantified using the area under the curve of a constructed Receiver Operating Curve (ROC).

**Results** When controlling for all other characteristics, the odds of CCD increased 52% with each additional year of age (OR=1.52, 95% CI 1.44, 1.61). Among dogs of the same age, health status, breed type, and sterilization status, odds of CCD was 6.47 times higher in dogs who were not active compared to those who were very active (OR=6.47, 95% CI 2.93-17.23). When controlling for age, breed type, activity level, and other comorbidities, dogs with a history of neurological, eye, or ear disorders had higher odds of CCD (OR=1.84, 95% CI 1.26, 2.65; OR=2.16, 95% CI 1.57, 2.98; OR=1.96, 95% 1.42, 2.70, respectively). The diagnostic threshold of CSLB score ≥ 50 (which corresponded with positive CCD status) had excellent predictive capacity for CCD (AUC=0.884).

**Conclusions** There existed a positive association between dog age and CCD. Positive associations between neurological, ear, or eye disorders and CCD, as well as an inverse association between physical activity and CCD, were also identified. Finally, canine life stage showed excellent discriminating ability between CCD positive and negative dogs. Quartile estimation could therefore serve as a tool to inform CCD screening by veterinarians.
Introduction

Neurodegenerative diseases associated with aging have become increasingly prevalent among the aging American population. In 2020, approximately 5.8 million Americans were living with Alzheimer’s disease (AD). It is the sixth leading cause of death in the country (1). Despite this, the complex pathological pathways that lead to the development of AD are still not fully understood and can be difficult to study *in vivo* in early phases of disease progression. While transgenic animal models have been extensively used to study the pathophysiology of AD, limitations have been identified that have prompted investigation into non-transgenic animal models (2).

The clinical and histological presentation of human AD and Canine Cognitive Dysfunction (CCD) in dogs have many similarities. As with humans, canine cognitive function declines throughout the course of the animal’s lifespan (3). Clinical signs of this decline appear to be related to learning and memory deficits, loss of spatial awareness, altered social interactions, and disrupted sleeping patterns (4, 5, 6). Further, the presentation of human AD and CCD share certain neuropathological features such as amyloid-β plaque deposition (2, 7, 8, 9, 10).

The observed parallels between CCD and human AD suggests that dogs exhibiting CCD may offer researchers a valuable animal model in which to study characteristics of neurodegenerative diseases that are relevant to, but challenging to study in, human populations (11). Further, dogs with CCD could serve as candidates for AD preventative and/or therapeutic strategies (2). Finally, CCD is also a major health concern of dog owners and veterinarians. An increased understanding of CCD may help to advance treatment of cognitive disease in dogs. These potential benefits highlight the importance of accurate diagnosis of CCD in companion dogs.
A wide range of structured interviews and theory-driven scales for evaluating CCD have been developed within the last 20 years (4, 12, 13, 14). These studies have shown a correspondingly wide range of prevalence estimates in their study populations that nonetheless appear to indicate an increased prevalence of cognitive impairment as a dog ages (7, 14, 15). In order to gain a better understanding of this trend in aging canines, researchers recently have developed assessment tools that are able to distinguish cognitively impaired aged dogs from those who are experiencing healthy aging (11, 14).

These scales provide veterinarians and researchers with tools to better identify the prevalence of CCD in aging dogs but have only been used to describe cognitive function in relatively small populations (<1,000 dogs) (11, 14). The purpose of this study was to describe the range of cognitive function scores and prevalence of CCD in a very large, nationwide sample of companion dogs participating in the Dog Aging Project (DAP) (n = 20,413). DAP used a minimally modified version of the validated canine cognitive dysfunction rating scale (CCDR) developed by Salvin et al. (4). Further, we examined associations between canine life stage and CCD. Classifying a canine’s lifespan into life stage categories and quantifying its predictive ability potentially allows for preventative healthcare measures to be taken by owners and veterinarians. The results could eventually lead to more timely detection and treatment of CCD (16).

Methods

Study Setting: Data were obtained from the National Institute on Aging-funded nationwide longitudinal study on canine aging and mortality in US companion dogs, the Dog Aging Project (DAP) initiated in 2018. Project 1 of the DAP aims to better define aging in dogs based on comorbidities, frailty, and inflammaging (17). Companion dog owners interested in enrolling their dog completed multiple surveys throughout their involvement in the DAP. For the purposes of this study, these included the baseline Health and Life Experience Survey (HLES) and the
Canine Social and Learned Behavior (CSLB) survey. The HLES is an extensive online survey that was distributed to all dog owners upon baseline enrollment, with sections on dog and owner household demographic characteristics, dog physical activity, environment, behavior, diet, medications and preventatives, and health status. The CSLB survey is a 13-item questionnaire based on Salvin et al.’s CCDR scale that aims to assess CCD (4). It is described in further detail below.

**Study Design:** This study analyzed baseline associations between selected characteristics collected through the HLES and dog cognitive characteristics ascertained through the CSLB. HLES data were provided by participants immediately upon enrollment, and CSLB data were provided by participants when that survey was administered. Dates of collection for HLES data used in the study ranged from December 23, 2019 to December 11, 2020. Dates of collection for CSLB data used in the study ranged from October 30, 2020 to December 31, 2020. The median time between completion of each survey was 5 weeks (range: 0–50.3 weeks).

**Study Subjects:** All dogs whose owners did not complete the baseline HLES and CSLB surveys prior to December 31, 2020 were excluded from study. Further, all dogs whose owners indicated that they were not able to confidently report their dog’s age, as assessed in the HLES, were also excluded. We began with a study sample of 27,542 dogs whose owners had completed the baseline HLES; 20,096 of those dogs’ owners had also completed the CSLB survey. Of those 20,096 dogs, 15,019 owners reported being sure of their dog’s age. The final sample for this study consisted of 15,019 dogs.

**Data Sources:** The validated CCDR instrument, used by Salvin et al for the identification of CCD, was minimally modified and renamed the CSLB (Canine Social and Learned Behavior Survey) for this study. In contrast to the items assessed by the CCDR: 1) the CSLB survey assigned a continuous (rather than binary) score for each item (described below); 2) the instrument name was modified so as not to suggest dementia or cognitive dysfunction (identified
in the term CCD) to the owners of enrolled dogs; and 3) it was programmed as an on-line digital tool with American English spelling. The CSLB thus consisted of 13 questions that assessed behaviors such as getting stuck behind objects, pacing, and failing to recognize familiar people. Responses for each question were scored based on frequency of the behavior (1=never, 2=once a month, 3=once a week, 4=once a day, 5=more than once a day). Certain questions had multipliers based on their clinical severity, as identified in CCD diagnosed dogs (4). In addition, some questions asked owners to compare the severity of their dog’s current behavior to the severity of the behavior 6 months prior. Numeric scores were summed from each owner’s responses. A minimum score of 16 and a maximum score of 80 were possible, with higher values indicating more severe levels of cognitive dysfunction. Age, sex, sterilization status, breed group, geographic region, physical activity level, comorbidities, and primary formal dog activity were assessed through the HLES.

For owner-reported purebred dogs, breed was elicited and assigned based on the eight groups defined by the American Kennel Club: herding, hound, toy, non-sporting, sporting, terrier, working, and miscellaneous/Foundation Stock Service (18). Primary formal dog activity refers to a dog’s main job or activity with the owner and was characterized as one of the following: companion animal or pet, obedience, show, breeding, agility, hunting, working, field trials, search and rescue, service dog, or assistance/therapy dog. Physical activity over the past year was classified as not active, moderately active, or very active. Major comorbidities considered were owner-reported in the HLES, including any history of cancer, endocrine disorders, kidney disorders, immune disorders, trauma, toxin consumption, infectious disease, hematologic disorders, neurological disorders, orthopedic disorders, reproductive disorders, liver disorders, gastrointestinal disorders, respiratory disorders, cardiac disorders, sin disorders, oral disorders, eye disorders, or ear disorders. Geographic region was determined based on the owner’s
primary state of residence and categorized as West, Southwest, Midwest, Southeast, or Northeast.

**Data Analyses:** The primary outcome of interest was total CSLB score. Additionally, canine cognitive function was treated as a binary measure such that dogs with a CSLB score ≥50 were classified as having CCD (4). Logistic regression models were fit to assess univariable associations between CCD and age, sex, sterilization status, breed group, geographic region, major comorbidities, and activity variables. Covariates that were determined *a priori* to be associated with CCD, informed by a directed acyclic graph (DAG), were included in a multivariable logistic regression model.

Predicted canine life stage quartile was then estimated for each subject. Projected mean life expectancies were calculated on a separate data set collected by Urfer et al. consisting of private U.S. veterinary hospital medical records for companion dogs (19). Mean life expectancies were then assigned to each dog in the DAP data set as a function of their weight or projected weight class, sterilization status, and sex (16, 19). Each dog was assigned a predicted life stage quartile that was equivalent to the proportion of their current life lived to their projected lifespan. Life stage quartile was then evaluated for its ability to predict CCD using adjusted and unadjusted logistic regression models. The fitted models were used to construct a Receiver Operating Curve (ROC) for two possible diagnostic thresholds (CSLB score ≥50, representative of previously defined thresholds, and >37, which represented the highest quartile of scores in the data). Finally, area under the curve (AUC) was calculated in order to summarize predictive capacity of the fitted models. All statistical analyses were conducted in RStudio (Version 1.3) (20).

**Results**
Of the 15,019 dogs included in analyses, 19.5% were classified as being in their 4th quartile of life, 24.5% were classified as being in their 3rd quartile of life, 27.0% were in their 2nd quartile of life, and 29.1% of dogs in their 1st quartile of life. A total of 1.4% of dogs were classified as having CCD using the binary cut-off of ≥ 50. There were similar distributions of sex, geographic region, primary role, and breed type among each life stage (Table 1). Sterilization status, history of each of 17 categories of health problems ascertained in the HLES, purebred breed group, and activity level were substantially positively associated with life stage. Dogs in a higher life stage were more likely to be sterilized, more likely to have history of the above comorbidities, less likely to be classified as a purebred breed group and were less active.

When evaluating the continuous CSLB score, no meaningful relationship was identified between CSLB score and age (R²=0.08). Associations between various canine characteristics and binary CCD assignment were therefore assessed using univariable and multivariable logistic regression models (Table 2). The odds of being diagnosed with CCD increased almost 70% with each additional year of age (OR=1.68, 95% CI 1.60, 1.77).

A DAG was constructed to highlight relevant covariates to include in the multivariable logistic regression model assessing the association between age and CCD (Figure 1). Age, sterilization status, history of 15 categories of health problems, breed type, and activity level were selected for the final logistic regression model (Table 2). When controlling for all other characteristics, the odds of CCD increased 52% with each additional year of age (OR=1.52, 95% CI 1.44, 1.61). Interestingly, among dogs of the given age, health status, breed type, and activity level, those who were intact had a higher odds of CCD (OR=1.21, 95% CI 0.51-2.51). Although attenuated, an inverse association between activity level and CCD was still found. Among dogs of the same age, health status, breed type, and sterilization status, odds of CCD was 6.47 times higher in dogs who were not active compared to those who were very active (OR=6.47, 95% CI 2.93-17.23). When controlling for age, breed type, activity level, and other comorbidities, dogs with a
history of neurological, eye, or ear disorders had higher odds of CCD (OR=1.84, 95% CI 1.26, 2.65; OR=2.16, 95% CI 1.57, 2.98; OR=1.96, 95% 1.42, 2.70, respectively).

Adjusted and unadjusted logistic regression models were then fit using the four-quartile categorical canine life stage. Adjusted models included all covariates from the multivariable logistic regression model assessing the association between age and CCD (Table 2). Models were evaluated for their ability to predict CCD at two different diagnostic thresholds: ≥ 50, as cited in previous literature, and > 37, which captures the upper 4th quartile of CSLB scores. In total, 4 predictive models were assessed by evaluating the area under the curve (AUC) of an ROC curve (Figure 2). The diagnostic threshold of CSLB score ≥ 50 had the highest predictive capacity. The adjusted model that controlled for health status, breed type, sterilization status, and activity level provided a slightly improved predictive capacity over the unadjusted model (AUC=0.892, 95% CI: 0.854-0.930 vs. AUC=0.862, 95% CI: 0.831-0.893). The threshold of CSLB score > 37 had lower predictive capacity for both the adjusted and unadjusted models (AUC=0.701, 95% CI: 0.682-0.721 vs. AUC=0.644, 95% CI: 0.624-0.665).

Discussion

Results from this study indicate a positive association between age and CCD in companion dogs, even adjusting for other characteristics in a multivariable logistic regression model. There also existed associations between CCD and decreased activity level, as well as CCD and history of an eye, ear, or neurological disorder. The association between age and CCD aligns with the progressive nature of CCD and with previous canine research findings, which have shown an exponential increase in CCD prevalence with increasing age (4).

An inverse association was seen in dogs whose owners indicated higher activity levels over the past year. Previous studies with rodent models have demonstrated that exercise can have protective effects against the development of biological markers and subsequent behavioral
deficits characteristic of AD, and numerous observational human studies have consistently shown inverse associations between exercise and AD (21, 22, 23, 24, 25). These observations may reflect a variety of biologic mechanisms, including a reduction of pro-inflammatory cytokines in the brain that otherwise contribute to neural damage and death, and an increase in neural plasticity (24, 25). The reduced odds of CCD among more active dogs in our cohort may be a result of these same mechanisms.

The association we observed between dogs who have ever had a neurological disorder and their prevalence of CCD is expected. A possible disorder that an owner could have indicated was dementia, which, in humans, is characterized by a decline in cognitive function. Seizure disorders, which also could have been indicated, have been found to occur more commonly in human AD patients than in comparable individuals without AD (26).

There have been human studies demonstrating the potential links between eye and ear disorders and AD. Impairments in visual acuity and visual fields have been observed more frequently in AD patients than in similarly aged individuals without AD (27, 28). Further, the presence of ophthalmic conditions, such as cataracts and age-related macular degeneration has been found to be associated with an increased risk of all-cause dementia and AD (29). Amyloid deposits, which have been linked to the development of AD and have been associated with the acceleration of AD progression, have been found in the lens of the eye in some individuals with age-related cataracts, potentially indicating common neuropathological pathways leading to AD and cataracts (30). Age-related macular degeneration has also been associated with an increased risk of all-cause dementia and AD (30). Similar to the potential relationship between cataracts and AD, amyloid deposits have been detected in the small fatty protein deposits that accumulate under the eye of individuals with age-related macular degeneration (31).
In addition to numerous studies identifying hearing loss as a possible risk factor for cognitive decline and all-cause dementia, a recent study found that individuals with dual sensory impairment (combined visual and hearing impairments) had a substantially increased risk of AD, possibly as a result of reduced neural resources needed for cognitive performance (31, 32, 33, 34). These associations are potentially mirrored in our companion dog cohort.

Alternatively, the associations we observed between sensory impairment and CCD could be due at least in part to misclassification. The CSLB survey, which is closely derived from Salvin et al.’s CCDR, intended to distinguish between behaviors associated with CCD as opposed to those that are part of normal canine aging (4). However, some behaviors ascertained through the rating scale may not arise solely from cognitive impairment (4). Questions in the survey such as, “How often does your dog walk into walls or doors?” and “How often does your dog have difficulty finding food dropped on the floor?” were asked due to their association with CCD. However, they could result in high scores due to blindness or other eye disorders as opposed to CCD.

We observed a dog’s estimated life stage quartile, which is a function of their age, weight, sex, and sterilization status, to have excellent discriminating ability between CCD positive and negative dogs (CSLB score ≥ 50). This improved only slightly in the model adjusted for age, sterilization status, history of 15 categories of health problems, breed type, and activity level. We also considered a lower diagnostic threshold for CCD assignment by using the top quartile of CSLB scores as a cut-off. However, that classification yielded much lower discriminating ability between CCD positive and negative dogs for life stage quartile. Although the diagnostic threshold of 50 had high predictive capacity, it is important to note that this threshold between normal aging and CCD was chosen by previous researchers, and the sensitivity and specificity were not reported (4). It would therefore be beneficial to examine the validity of this diagnostic tool more closely in future studies.
This study had a number of strengths, including a large sample size, standardized data collection methods, high participant response rates, and the inclusion of many potential confounders in analytic models. Important limitations did exist in this study that are worth addressing. Aside from potential concerns with the sensitivity and specificity of the diagnostic threshold chosen for CCD, this study is not longitudinal, and so it is not possible to assess any causal links between the various canine characteristics and CCD. While the maximum time interval between survey completion was 50.3 weeks, the median time between survey completion was only 5 weeks. There was a low likelihood that this timeframe would result in any sort of meaningful changes in CCD status. Future studies using prospective DAP data will be able to explore potential risk factors for CCD as well as cognitive decline over time.

Another limitation is that all data used in this analysis were based on surveys completed by owners. Owner-reported information is potentially susceptible to multiple forms of bias. Questions requiring owners to recall behaviors and medical conditions that could have occurred between 6 months to many years prior could introduce either differential or non-differential misclassification, depending on whether the errors in recall were correlated with the dog’s cognitive status. Non-differential misclassification could arise from social desirability if, for example, owners indicated a higher level of physical activity for their dog, or a lower level of impairment on the CSLB survey. Such error would tend to dampen associations between physical activity and CCD.

It is also important to recognize the drastic changes that have taken place in many households due to the COVID-19 pandemic, and the possibility that these changes influenced our results. For example, depending on when an owner completed their HLES and CSLB surveys, their dog’s activity level may have changed as a result of stay-at-home orders and/or the owner’s ability to work from home. Additionally, owners spending more time at home with their dog may have an impact a dog’s actual health, and it may increase the likelihood of observing specific
health behaviors that would affect a dog’s reported health status. These types of major lifestyle changes could have altered our data in ways that would be difficult to quantify. Finally, there is the potential presence of unmeasured confounders that were not captured by our surveys, as well as survey data that were not considered for analysis, which could result in residual confounding potentially adding bias to our results.

Conclusions

We identified a positive association between companion dog age and CCD. We also identified strong positive associations between history of a neurological, ear, or eye disorder as well as an association between increased physical activity level and having a CSLB score of 50 or higher. Dogs were classified into their predicted quartile of life based on their age, weight, sex, and sterilization status. Using a binary diagnostic threshold of ≥50, a dog’s life stage quartile showed excellent discriminating ability between CCD positive and negative dogs. This quartile estimation could potentially serve as a useful tool to inform whether a dog should be screened for CCD by their veterinarian. Finally, given increasing evidence of the parallels between canine and human cognitive disease, accurate CCD diagnosis in canines may provide researchers with more suitable animal models in which to study aging in human populations.
## Tables and Figures

**Table 1:** Selected canine demographic characteristics by life stage quartile (n = 15,019), Dog Aging Project 2020-2021

| Characteristic          | First (n = 4,368) | Second (n = 4,050) | Third (n = 3,676) | Fourth (n = 2,925) |
|-------------------------|------------------|-------------------|------------------|-------------------|
| **Sex**                 |                  |                   |                  |                   |
| Female                  | 2,207 (50.5)     | 1,987 (49.1)      | 1,865 (50.7)     | 1,425 (48.7)      |
| Male                    | 2,161 (49.5)     | 2,063 (50.9)      | 1,811 (49.3)     | 1,500 (51.3)      |
| **Sterilization Status**|                  |                   |                  |                   |
| Intact                  | 834 (19.1)       | 319 (7.9)         | 191 (5.2)        | 116 (4.0)         |
| Desexed                 | 3,534 (80.9)     | 3,731 (92.1)      | 3,485 (94.8)     | 2,809 (96.0)      |
| **Any Comorbidity History** |                |                   |                  |                   |
| Cancer                  | 24 (0.5)         | 98 (2.4)          | 270 (7.3)        | 456 (15.6)        |
| Endocrine disorder      | 4 (0.1)          | 50 (1.2)          | 155 (4.2)        | 228 (7.8)         |
| Kidney disorder         | 163 (3.7)        | 240 (5.9)         | 296 (8.1)        | 434 (14.8)        |
| Immune disorder         | 7 (0.2)          | 34 (0.8)          | 36 (1.0)         | 44 (1.5)          |
| Trauma                  | 732 (16.8)       | 1,070 (26.4)      | 1,191 (32.4)     | 1,029 (35.2)      |
| Toxic Consumption       | 376 (8.6)        | 400 (9.9)         | 470 (12.8)       | 382 (13.1)        |
| Infectious disease      | 1,170 (26.8)     | 1,093 (27.0)      | 949 (25.8)       | 791 (27.0)        |
| Hematologic disorder    | 13 (0.3)         | 12 (0.3)          | 12 (0.3)         | 36 (1.2)          |
| Neurological disorder   | 34 (0.8)         | 108 (2.7)         | 166 (4.5)        | 356 (12.2)        |
| Orthopedic disorder     | 218 (5.0)        | 489 (12.7)        | 822 (22.4)       | 1,196 (40.9)      |
| Reproductive disorder   | 114 (2.6)        | 117 (2.9)         | 98 (2.7)         | 70 (2.4)          |
| Liver disorder          | 33 (0.8)         | 80 (2.0)          | 164 (4.5)        | 252 (8.6)         |
| Gastrointestinal disorder | 506 (11.6)   | 579 (14.3)        | 572 (15.6)       | 548 (18.7)        |
| Respiratory disorder    | 51 (1.2)         | 62 (1.5)          | 121 (3.3)        | 231 (7.9)         |
| Cardiac disorder        | 49 (1.1)         | 104 (2.6)         | 269 (7.3)        | 411 (14.1)        |
| Skin disorder           | 767 (17.6)       | 1,125 (27.8)      | 1,232 (33.5)     | 1,100 (37.6)      |
| Oral disorder           | 340 (7.8)        | 803 (19.8)        | 1,316 (35.8)     | 1,410 (48.2)      |
| Eye disorder            | 261 (6.0)        | 345 (8.5)         | 493 (13.4)       | 822 (28.1)        |
| Ear disorder            | 328 (7.5)        | 380 (9.4)         | 463 (12.6)       | 751 (25.7)        |
| **Geographic Region**   |                  |                   |                  |                   |
| Midwest                 | 982 (22.5)       | 871 (21.5)        | 762 (20.7)       | 597 (20.4)        |
| Northeast               | 813 (18.6)       | 715 (17.7)        | 698 (19.0)       | 548 (18.7)        |
| Southeast               | 770 (17.6)       | 751 (18.5)        | 658 (17.9)       | 548 (18.7)        |
| Southwest               | 380 (8.7)        | 362 (8.9)         | 365 (9.9)        | 286 (9.8)         |
| West                    | 1,395 (31.9)     | 1,326 (32.7)      | 1,178 (32.0)     | 925 (31.6)        |
| Missing                 | 28 (0.6)         | 25 (0.6)          | 15 (0.4)         | 21 (0.7)          |
| Primary Role          | 4,169 (95.4) | 3,850 (95.1) | 3,498 (95.2) | 2,802 (95.8) |
|----------------------|--------------|--------------|--------------|--------------|
| Companion/pet        | 4,169 (95.4) | 3,850 (95.1) | 3,498 (95.2) | 2,802 (95.8) |
| Agility              | 9 (0.2)      | 16 (0.4)     | 15 (0.4)     | 6 (0.2)      |
| Assistance or therapy| 22 (0.5)     | 32 (0.8)     | 32 (0.9)     | 20 (0.7)     |
| Breeding             | 4 (0.1)      | 5 (0.1)      | 2 (0.05)     | 7 (0.2)      |
| Field trials         | 2 (0.1)      | 2 (0.1)      | 2 (0.05)     | 0 (0)        |
| Hunting              | 6 (0.1)      | 7 (0.2)      | 7 (0.2)      | 4 (0.1)      |
| Obedience            | 39 (0.9)     | 22 (0.5)     | 14 (0.4)     | 15 (0.5)     |
| Search and rescue    | 10 (0.2)     | 7 (0.2)      | 3 (0.1)      | 4 (0.1)      |
| Service              | 42 (1.0)     | 42 (1.0)     | 33 (0.9)     | 17 (0.6)     |
| Show                 | 10 (0.2)     | 6 (0.1)      | 7 (0.2)      | 3 (0.1)      |
| Working              | 9 (0.2)      | 14 (0.3)     | 11 (0.3)     | 7 (0.2)      |
| Other                | 46 (1.1)     | 47 (1.2)     | 52 (1.4)     | 40 (1.4)     |

| Breed Type          | Purebred      | Mixed        | Purebred      | Mixed        |
|---------------------|---------------|--------------|---------------|--------------|
|                     | 2,259 (51.7)  | 1,784 (48.3) | 2,736 (58.7)  | 1,667 (41.3) |
|                     | 2,205 (60.0)  | 1,466 (40.0) | 1,733 (59.2)  | 1,189 (40.8) |

| Purebred Breed Group | Herding | Hound | Non-Sporting | Sporting | Terrier | Toy | Working | Miscellaneous/FSS | Non-AKC |
|----------------------|---------|-------|--------------|----------|---------|-----|---------|------------------|---------|
|                      | 475 (10.9) | 175 (4) | 269 (6.2) | 900 (20.6) | 151 (3.5) | 223 (5.1) | 319 (7.3) | 32 (0.7) | 34 (0.8) |
|                      | 452 (11.2) | 180 (4.4) | 232 (5.7) | 788 (19.5) | 179 (4.4) | 215 (5.3) | 272 (6.7) | 26 (0.6) | 32 (0.8) |
|                      | 331 (9.0)  | 206 (5.6) | 198 (5.4) | 684 (18.6) | 204 (5.5) | 310 (8.4) | 227 (6.2) | 19 (0.5) | 26 (0.7) |
|                      | 255 (8.7)  | 162 (5.5) | 156 (5.3) | 542 (18.5) | 176 (6.0) | 242 (8.3) | 168 (5.7) | 17 (0.6) | 15 (0.5) |

| Activity Level       | Very Active | Moderately Active | Not Active |
|----------------------|-------------|-------------------|------------|
|                      | 1,552 (35.5)| 2,696 (61.7)      | 120 (2.7)  |
|                      | 919 (22.7)  | 2,828 (69.8)      | 303 (7.5)  |
|                      | 528 (14.4)  | 2,630 (71.5)      | 518 (14.1) |
|                      | 236 (8.1)   | 1,914 (65.4)      | 775 (26.5) |
### Table 2: Association between selected dog characteristics and Canine Cognitive Dysfunction, Dog Aging Project 2020-2021

| Characteristic (n) | Univariable Analysis | Multivariable Analysis |
|-------------------|----------------------|------------------------|
|                   | OR 95% CI            | OR 95% CI              |
| Age (integer years) | 1.68 1.60-1.77       | 1.52 1.44-1.61         |
| Sex               |                      |                        |
| Female (reference) | 1.00                 |                        |
| Male (7,535)      | 0.91 0.69-1.19       |                        |
| Sterilization status |                 |                        |
| Desex (reference) | 1.00                 |                        |
| Intact (1,460)    | 0.36 0.16-0.67       | 1.21 0.51-2.51         |
| Activity level    |                      |                        |
| Very Active (reference) | 1.00             |                        |
| Moderately Active | 4.80 2.29-12.33      | 2.19 1.00-5.83         |
| Not Active (1,716) | 40.46 19.42-103.51   | 6.47 2.93-17.23        |
| Breed group       |                      |                        |
| Sporting (reference) | 1.00             |                        |
| Herding (1,513)   | 1.07 0.55-1.99       |                        |
| Hound (723)       | 1.65 0.78-3.26       |                        |
| Terrier (710)     | 3.58 2.02-6.28       |                        |
| Toy (990)         | 3.80 2.29-6.38       | -                      |
| Non-Sporting (855) | 3.49 2.03-6.00      | -                      |
| Working (986)     | 0.76 0.31-1.67       |                        |
| Misc./FSS (94)    | 0.00                 |                        |
| Non-AKC (107)     | 1.01 0.05-4.80       |                        |
| Geographic region |                      |                        |
| West (reference)  | 1.00                 |                        |
| Midwest (3,212)   | 0.82 0.55-1.19       |                        |
| Northeast (2,774) | 0.90 0.61-1.32       | -                      |
| Southeast (2,727) | 0.82 0.55-1.22       | -                      |
| Southwest (1,393) | 0.81 0.47-1.32       |                        |
| History of Cancer |                      |                        |
| No (reference)    | 1.00                 |                        |
| Yes (848)         | 4.09 2.85-5.72       | 1.35 0.90-1.97         |
| History of Kidney disorder |         |                        |
| No (reference)    | 1.00                 |                        |
| Yes (1,133)       | 3.53 2.52-4.87       | 1.21 0.82-1.76         |
| History of Endocrine disorder |     |                        |
| No (reference)    | 1.00                 |                        |
| Yes (437)         | 3.74 2.30-5.80       | 1.09 0.63-1.79         |
| History of Immune disorder |       |                        |
| No (reference)    | 1.00                 |                        |
| Yes (121)         | 1.16 0.19-3.67       |                        |
| History of Trauma |                      |                        |
| No (reference)    | 1.00                 |                        |
| Yes (4,022)       | 1.63 1.23-2.15       | 1.10 0.79-1.50         |
| History of Toxin Consumption |     |                        |
| No (reference)    | 1.00                 |                        |
| Yes (1,628)       | 1.61 1.10-2.29       | 1.13 0.73-1.69         |
| History of Infectious disease |         |                        |
| No (reference)    | 1.00                 |                        |
| Yes (4,003)       | 0.81 0.58-1.11       |                        |
| History of hematologic disorder |       |                        |
| No (reference)    | 1.00                 |                        |
| Yes (73)          | 5.16 1.79-11.71      |                        |
| History of Neurological disorder |        |                        |
| No (reference)    | 1.00                 |                        |
| Condition                        | Yes (n) | OR       | 95% CI       | P-value |
|---------------------------------|---------|----------|--------------|---------|
| History of Orthopedic disorder  |         | 8.22     | 5.96-11.20   | 1.84    | 1.26-2.65 |
| History of Reproductive disorder|         | 1.00     | -            | 1.00    | -         |
| History of Liver disorder       |         | 1.00     | -            | 1.00    | -         |
| History of Gastrointestinal disorder |   | 1.00     | -            | 1.00    | -         |
| History of Respiratory disorder |         | 1.00     | -            | 1.00    | -         |
| History of Cardiac disorder     |         | 1.00     | -            | 1.00    | -         |
| History of Skin disorder        |         | 1.00     | -            | 1.00    | -         |
| History of Oral disorder        |         | 1.00     | -            | 1.00    | -         |
| History of Eye disorder         |         | 1.00     | -            | 1.00    | -         |
| History of Ear disorder         |         | 1.00     | -            | 1.00    | -         |
| Breed                           |         | 1.00     | -            | 1.00    | -         |
| Mixed                           | (6,106) | 1.00     | -            | 1.00    | -         |
| Purebred                        | (8,913) | 1.43     | 1.07-1.91    | 1.24    | 0.90-1.72 |
| Primary activity                |         | 1.00     | -            | 1.00    | -         |
| Pet                             | (14,319) | 1.00     | -            | 1.00    | -         |
| Agility                         | (46)    | 1.49     | 0.08-6.85    |         |
| Assistance or therapy           | (106)   | 0.00     | -            | 0.00    | -         |
| Breeding                        | (18)    | 0.00     | -            | 0.00    | -         |
| Field Trials                    | (6)     | 0.00     | -            | 0.00    | -         |
| Hunting                         | (24)    | 0.00     | -            | 0.00    | -         |
| Obedience                       | (90)    | 0.00     | -            | 0.00    | -         |
| Other                           | (185)   | 0.73     | 0.12-2.30    |         |
| Search and Rescue               | (24)    | 0.00     | -            | 0.00    | -         |
| Service                         | (134)   | 0.50     | 0.03-2.26    |         |
| Show                            | (26)    | 0.00     | -            | 0.00    | -         |
| Working                         | (41)    | 0.00     | -            | 0.00    | -         |

OR: odds ratio; CI: confidence interval
*Sex was not associated with either age or CCD, and so was not included in the DAG.

**These variables were identified as confounders that were associated with age and CCD and were included in the multivariable models.

Dashed lines represent hypothesized associations considered based on relevance
Figure 2: ROC curve generated from predictive models relating selected dog characteristics to Canine Cognitive Dysfunction prevalence, Dog Aging Project 2020-2021
References

1. Alzheimer’s Association. Alzheimer’s Association 2020 Facts and Figures Report. Alzheimer’s Assoc. 2020:1. https://www.alz.org/alzheimers-dementia/facts-figures%0Ahttps://alz.org/alzheimers-dementia/facts-figures%0Ahttps://www.alz.org/alzheimers-dementia/facts-figures%0Ahttps://www.alz.org/media/Documents/alzheimers-facts-and-figures_1.pdf. Accessed February 21, 2021.

2. Schütt T, Helboe L, Pedersen LØ, Waldemar G, Berendt M, Pedersen JT. Dogs with cognitive dysfunction as a spontaneous model for early Alzheimer’s disease: a translational study of neuropathological and inflammatory markers. J Alzheimer’s Dis. 2016;52(2):433-449. doi:10.3233/JAD-151085

3. Adams B, Chan A, Callahan H, Milgram NW. The canine as a model of human cognitive aging: Recent developments. Prog Neuro-Psychopharmacology Biol Psychiatry. 2000;24(5):675-692. doi:10.1016/S0278-5846(00)00101-9

4. Salvin HE, McGreevy PD, Sachdev PS, Valenzuela MJ. The canine cognitive dysfunction rating scale (CCDR): A data-driven and ecologically relevant assessment tool. Vet J. 2011;188(3):331-336. doi:10.1016/j.tvjl.2010.05.014

5. Ruehl WW, Hart BL. Canine cognitive dysfunction. In: Dodman N, Shuster L, eds. Psychopharmacology of animal behavior disorders. Malden, Mass: Blackwell Scientific Publications, 1998;283–304.

6. Ruehl WW, Bruyette DS, De Paoli A, et al. Canine cognitive dysfunction as a model for human age-related cognitive decline, dementia and Alzheimer’s disease: clinical presentation, cognitive testing, pathology and response to l-deprenyl therapy. Prog Brain Res 1995;106:217–225.

7. Azkona G, García-Belenguer S, Chacón G, Rosado B, León M, Palacio J. Prevalence and risk factors of behavioural changes associated with age-related cognitive impairment in geriatric dogs: PAPER. J Small Anim Pract. 2009;50(2):87-91. doi:10.1111/j.1748-5827.2008.00718.x

8. Cummings B, Su JH, Cotman CW, et al. β-Amyloid accumulation in aged canine brain. A model of early plaque formation in Alzheimer’s disease. Neurobiol Aging 1993;14:547–560.

9. Cummings BJ, Satou T, Head E, et al. Diffuse plaques contain C-terminal Aβ42 and not Aβ40: evidence from cats and dogs. Neurobiol Aging 1996;17:653–659.

10. Cummings BJ, Head E, Ruehl WW, et al. The canine as an animal model of human aging and dementia. Neurobiol Aging 1996;17:259–268

11. Madari A, Farbakova J, Katina S, et al. Assessment of severity and progression of canine cognitive dysfunction syndrome using the CAnine DEmentia Scale (CADES). Appl Anim Behav Sci. 2015;171:138-145. doi:10.1016/j.applanim.2015.08.034
12. Colle M-A, Hauw J-J, Crespeau F, et al. Vascular and parenchymal A-beta deposition in the aging dog: correlation with behavior. Neurobiol Aging. 2000;21(5):695-704. doi:10.1016/s0197-4580(00)00113-5

13. Pugliese M, Carrasco JL, Andrade C, Mas E, Mascort J, Mahy N. Severe cognitive impairment correlates with higher cerebrospinal fluid levels of lactate and pyruvate in a canine model of senile dementia. Prog Neuropsychopharmacol Biol Psychiatry. 2005;29(4):603-610. doi:10.1016/j.pnpbp.2005.01.017

14. Salvin HE, McGreevy PD, Sachdev PS, Valenzuela MJ. Under diagnosis of canine cognitive dysfunction: a cross-sectional survey of older companion dogs. Vet J. 2010;184(3):277-281. doi:10.1016/j.tvjl.2009.11.007

15. Neilson JC, Hart BL, Cliff KD, Ruehl WW. Prevalence of behavioral changes associated with age-related cognitive impairment in dogs. J Am Vet Med Assoc. 2001;218(11):1787-1791. doi:10.2460/javma.2001.218.1787

16. Creevy KE, Grady J, Little SE, et al. 2019 AAHA canine life stage guidelines. J Am Anim Hosp Assoc. 2019;55(6):267-290. doi:10.5326/JAAHA-MS-6999

17. Alexander JE, Colyer A, Haydock RM, Hayek MG, Park J. Understanding how dogs age: longitudinal analysis of markers of inflammation, immune function, and oxidative stress. J Gerontol - Ser A Biol Sci Med Sci. 2018;73(6):720-728. doi:10.1093/gerona/glx182

18. The American Kennel Club. List of Breeds by Group. https://www.akc.org/public-education/resources/general-tips-information/dog-breeds-sorted-groups/. Published 2020. Accessed November 12, 2020.

19. Urfer SR, Kaeberlein M, Promislow DEL, Creevy KE. Lifespan of companion dogs seen in three independent primary care veterinary clinics in the United States. Canine Med Genet. 2020;7(1):1-14. doi:10.1186/s40575-020-00086-8

20. R Core Team (2020). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/.

21. Lindsay J, Laurin D, Verreault R, et al. Risk factors for Alzheimer’s disease: a prospective analysis from the Canadian study of health and aging. Am J Epidemiol. 2002;156(5):445-453. doi:10.1093/aje/kwf074

22. Richards M, Hardy R, Wadsworth ME. Does active leisure protect cognition? Evidence from a national birth cohort. Soc Sci Med. 2003;56(4):785-792. doi:10.1016/s0277-9536(02)00075-8

23. Larson EB, Wang L, Bowen JD, et al. Exercise is associated with reduced risk for incident dementia among persons 65 years of age and older. Ann Intern Med. 2006;144(2):73-81. doi:10.7326/0003-4819-144-2-200601170-00004

24. Nascimento CM, Pereira JR, de Andrade LP, et al. Physical exercise in MCI elderly promotes reduction of pro-inflammatory cytokines and improvements on cognition
25. Duzel E, van Praag H, Sendtner M. Can physical exercise in old age improve memory and hippocampal function? Brain. 2016 Mar;139(Pt 3):662-73. doi: 10.1093/brain/awv407. Epub 2016 Feb 11. PMID: 26912638; PMCID: PMC4766381.

26. Amatniek JC, Hauser WA, DelCastillo-Castaneda C, et al. Incidence and predictors of seizures in patients with Alzheimer’s disease. Epilepsia. 2006;47(5):867-872. doi: 10.1111/j.1528-1167.2006.00554.x

27. Sadun AA, Borchert M, DeVita E, Hinton DR, Bassi CJ. Assessment of visual impairment in patients with Alzheimer's disease. Am J Ophthalmol. 1987;104(2):113-120. doi: 10.1016/0002-9394(87)90001-8

28. Lakshminarayanan V, Lagrave J, Kean ML, Dick M, Shankle R. Vision in dementia: contrast effects. Neurol Res. 1996;18(1):9-15. doi:10.1080/01616412.1996.11740369

29. Hwang PH, Longstreth WT Jr, Thielke SM, et al. Ophthalmic conditions associated with dementia risk: The Cardiovascular Health Study [published online ahead of print, 2021 Mar 31]. Alzheimer’s Dement. 2021;10.1002/alz.12313. doi:10.1002/alz.12313

30. Wang S, Mims PN, Roman RJ, Fan F. Is Beta-Amyloid accumulation a cause or consequence of Alzheimer’s disease? J Alzheimer's Parkinsonism Dement. 2016;1(2):007.

31. Loughrey DG, Kelly ME, Kelley GA, Brennan S, Lawlor BA. Association of age-related hearing loss with cognitive function, cognitive impairment, and dementia: a systematic review and meta-analysis [published correction appears in JAMA Otolaryngol Head Neck Surg. 2018 Feb 1;144(2):176]. JAMA Otolaryngol Head Neck Surg. 2018;144(2):115-126. doi:10.1001/jamaoto.2017.2513

32. Lin FR, Metter EJ, O'Brien RJ, Resnick SM, Zonderman AB, Ferrucci L. Hearing loss and incident dementia. Arch Neurol. 2011;68(2):214-220. doi:10.1001/archneur.2010.362

33. Johnson JCS, Marshall CR, Weil RS, Bamiou DE, Hardy CJD, Warren JD. Hearing and dementia: from ears to brain. Brain. 2021;144(2):391-401. doi:10.1093/brain/awaa429

34. Hwang PH, Longstreth WT Jr, Brenowitz WD, et al. Dual sensory impairment in older adults and risk of dementia from the GEM Study. Alzheimer's Dement (Amst). 2020;12(1):e12054. Published 2020 Jul 7. doi:10.1002/dad2.12054