Evaluating Spatial Overlap and Relatedness of White-tailed Deer in a Chronic Wasting Disease Management Zone

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Abstract

Wildlife disease transmission, at a local scale, can occur from interactions between infected and susceptible conspecifics or from a contaminated environment. Thus, the degree of spatial overlap and rate of contact among deer is likely to impact both direct and indirect transmission of infectious diseases such chronic wasting disease (CWD) or bovine tuberculosis. We identified a strong relationship between degree of spatial overlap (volume of intersection) and genetic relatedness for female white-tailed deer in Wisconsin’s area of highest CWD prevalence. We used volume of intersection as a surrogate for contact rates between deer and concluded that related deer are more likely to have contact, which may drive disease transmission dynamics. In addition, we found that age of deer influences overlap, with fawns exhibiting the highest degree of overlap with other deer. Our results further support the finding that female social groups have higher contact among related deer which can result in transmission of infectious diseases. We suggest that control of large social groups comprised of closely related deer may be an effective strategy in slowing the transmission of infectious pathogens, and CWD in particular.

Introduction

Social organization and interactions among individuals play an important role in the transmission and potential management of infectious wildlife diseases [1,2]. Many host characteristics such as sex, age, relatedness, density, social group composition, inter-group movement and isolation can influence the duration and intensity of contacts and disease transmission [1,3]. Understanding how contact rates and social organization influences disease transmission and spread is a challenging issue in disease ecology yet it is critical for disease management [2,4]. Complex social behaviors are typical for wild mammals and can result in disease transmission rates that are not explained by density alone [1,2]. In social species with stable group membership, given that at least one member is infected, individuals from the same group may have a higher rate of infection than non-members. White-tailed deer (Odocoileus virginianus) are one such social species known to associate in stable matrilineal groups [5,6,7]. Female white-tailed deer generally associate more closely with relatives than with non-relatives [8,9,10]. Although the nature and persistence of these interactions are still under study [3,6,7], this general pattern has widespread acceptance and is sometimes referred to as the “Rosepetal theory” [11]. However, deer social behavior, site fidelity, and spatial overlap can vary among different habitats deer density, and hunting pressure [12,13,14,15,16].

Chronic wasting disease (CWD) is a fatal neurodegenerative disease, posing serious and complex challenges for deer management [17,18]. In captive studies, CWD can be transmitted through animal-to-animal contact and indirect environmental contamination [19,20]. The relative importance of these transmission routes is not known in free-ranging deer. Probability of CWD infection in harvested female deer was recently found to be strongly influenced by genetic relatedness and, only incidentally, by spatial proximity to other infected females [21]. Ultimately, local transmission of CWD results from individual deer movements that lead to interactions with conspecifics or the environment. In particular, the degree of spatial overlap and contact among deer is likely to impact both direct and indirect transmission of CWD, and other infectious diseases such as bovine tuberculosis [3,6,22].

Although infection patterns and spatial distribution of CWD in Wisconsin have recently been described, there is limited empirical information on white-tailed deer behavior and interaction related to potential disease transmission and spread. Following discovery of the disease, an important management goal of the Wisconsin Department of Natural Resources (WDNR) was to reduce the deer population in the area where disease prevalence was highest.

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Table 1. Results of AIC model selection procedure to determine the best models predicting white-tailed deer spatial overlap in Wisconsin.

| Dataset | Variables | K | Delta AICc | Weight |
|---------|-----------|---|------------|--------|
| 1 (Pairs with at least one adult) | Age classes | Relatedness (Rxy-cat) | 4 | 0.00 | 0.71 |
| | Age classes (sex effects- fawns) | Relatedness (Rxy-cat) | 4 | 1.95 | 0.27 |
| | Age classes | Relatedness (Rxy) | 4 | 7.66 | 0.02 |
| | Age classes (sex effects- fawns) | Relatedness (Rxy) | 4 | 9.59 | 0.01 |
| | Age classes | Relatedness (Rxy-cat) | 3 | 13.63 | <0.01 |
| | Age classes (sex effects- fawns) | Relatedness (Rxy) | 3 | 15.55 | <0.01 |
| | Relatedness (Rxy) | Relatedness (Rxy-cat) | 3 | 24.62 | <0.01 |
| 2 (Capture groups 1 and 2) | Age classes | Relatedness (Rxy-cat) | 4 | 0.00 | 0.88 |
| | Age classes | Relatedness (Rxy) | 4 | 4.69 | 0.08 |
| | Age classes (sex effects- fawns) | Relatedness (Rxy-cat) | 4 | 7.38 | 0.02 |
| | Age classes | Relatedness (Rxy) | 3 | 8.05 | 0.02 |
| | Age classes (sex effects- fawns) | Relatedness (Rxy-cat) | 3 | 12.23 | <0.01 |
| | Relatedness (Rxy-cat) | Relatedness (Rxy) | 3 | 13.93 | <0.01 |
| | Age classes (sex effects- fawns) | Relatedness (Rxy) | 3 | 15.53 | <0.01 |
| | Relatedness (Rxy) | Relatedness (Rxy-cat) | 3 | 19.74 | <0.01 |

Datasetadult contains only deer-pairs including at least one adult deer, and datasetcapture group contains only capture groups 1 and 2 (see text). All models contain capture group and deer-pair as random effects. Age classes are either in categories of adult, yearling, and fawn (if lacking sex effects), or adult, yearling, male fawn, and female fawn. Relatedness is either represented as a continuous variable (Rxy) or categorical (Rxy-cat), in 3 categories consisting of Rxy of 0–0.25, 0.26–0.5, and 0.5–1.

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termed the Disease Eradication Zone (DEZ, [23]). Population reductions are most likely to be effective in reducing disease prevalence if disease transmission is density-dependent. Information on deer movement, space use, social structure and potential interaction are necessary to understand local-scale infectious contacts that generate the emergent dynamics of disease transmission on the landscape.

Although previous studies on deer spatial overlap have been conducted [3,6], we provide the first study evaluating links between deer spatial overlap, measured using VHF telemetry data, and deer relatedness based on microsatellite genetic markers. Based on female social structure, we hypothesize that related deer have greater spatial overlap, and thus more direct contact, than unrelated deer in the same areas. We also hypothesize that fawns, who are dependent on their mothers, will overlap with adjacent deer more strongly than yearlings or adults. These overlap areas, where multiple deer share space, are the most likely regions for either direct or indirect transmission of CWD [3]. As such, understanding the factors that determine the degree of overlap, and consequently the amount of direct contact, between deer will be critical to understanding and mitigating the spread of CWD and other infectious diseases.

Methods

Deer Tracking

Our study was conducted within two areas of Wisconsin’s Disease Eradication Zone (DEZ; see [24,25]), between January and April, 2003–2008. We captured 173 individual white-tailed deer (113 females, 60 males; [24,25], using modified Clover and Stephenson box traps, rocket nets [26], drop-nets [27], and darting. We aged deer as fawns (<1 year), yearlings (≥1 year, <2 years), and adults (≥2 years) by tooth wear and replacement [28]. We chemically immobilized captured deer [25] and affixed VHF radio collars. We tested deer for CWD using tonsillar biopsy [29] and collected blood and tissue samples for genetic analysis. Five deer initially tested CWD-positive (4 females, 1 male) and were culled.

We triangulated locations of radio-collared deer using 3 to 5 azimuths collected from fixed telemetry stations and obtained locations on a 24 hr basis, using rotating start times. We located radio-collared deer roughly 3 times/wk from 2003 to 2008 (Range: 1–6, Mean: 3.2). We estimated locations using Location of a Signal (LOAS), Version 2.09 [30], for groups of azimuths obtained within 20 min of each other. Estimates of positional error were ≤0.05 km². We spaced relocations of individual deer ≥6 hrs apart to minimize temporal autocorrelation.

Ethics Statement

The University of Wisconsin-Madison (UW) College of Agriculture and Life Sciences’ Animal Care and Use Committee (ACUC, Permit No. A-3368-01), UW Research Animal Resources Center (Permit No. A01088309-02), and the Wisconsin Department of Natural Resources (WDNR, Scientific Collector’s permit No. SCP-SCR-018-0202) approved capture and handling methods. Landowner permissions were acquired for capture on private lands.

Inclusion Criteria

We selected a subset of radio-tracked deer for analysis in this study. We excluded all male deer older than 1 yr from analysis because they display less philopatry than females and often engage in long-range movements [9]. Male fawns were retained due to their close association with their mother prior to dispersal. In addition, to limit the number of possible deer-pairs with extremely low or no overlap, we used only deer-pairs trapped within 1.5 km of each other. To address the potential lack of spatial independence among deer-pairs, we assigned deer trapped within...
1.5 km of one another to one of 7 distinct capture groups which were treated as a random effect in our analysis. We used only deer with >50 estimated locations (n = 105) during a given year.

Overlap Modeling
We used volume of intersection (VI) of utilization distributions [6,31] as our measure of spatial overlap. This measure has previously been found to correlate strongly with contact rates in deer [6]. Utilization distributions are three-dimensional probability densities that indicate relative space use based on point locations [32,33]. The VI is the approximate spatial integral of the square root of the product of two fixed utilization distribution kernels. VI values range from 0 to 1, with 0 representing no overlap, and 1 indicating complete overlap. For included deer-pairs, VI values were calculated for all points collected within one year (years were defined to begin on 10 May). To avoid correlation from multiple observations of the same deer-pairs (example: Deer 1003 and 1004 in years 2003, 2004, and 2005), we selected a single VI calculated for each deer-pair, from the year with the most combined locations. Volumes of intersection (VI) were then logit-transformed to facilitate analysis using linear models.

Relatedness
Whole genomic DNA was extracted from deer samples using a Qiagen DNeasy extraction kit (Qiagen Inc., Valencia, CA) following the manufacturer’s protocol for either 100 ul of blood or 20 mg of tissue from ear punch samples (all samples frozen since collection). We amplified 13 highly variable microsatellite loci using PCR with the Qiagen multiplex PCR kit [34]. We re-genotyped 32 individuals to assess errors in genotyping. We calculated Hardy-Weinberg equilibrium (HWE) and expected versus observed numbers of heterozygotes and homozygotes for all loci (using GenePop on the web; [35]) to assess data quality and assumptions for population genetics models. We used probability

![Figure 1. Scatterplots showing the relationship between degree of overlap and relatedness for white-tailed deer-pairs. Figure 1a is generated from the dataset where each pair contains at least one adult (datasetadult), and figure 1b is generated from the dataset using only deer-pairs from capture groups 1 and 2 (datasetcapgroup). Figure 1a. Figure 1b. doi:10.1371/journal.pone.0056568.g001](image)
of identity statistics (PID and PIDsibs, performed in GenAlEx, 36) to ensure adequate power to identify closely related individuals in our dataset. In order to provide a more genetically representative background with which to test our hypotheses, we supplemented the current sample with 100 additional deer from the same general geographic area that were genotyped in a collaborating study (analyzed using the same genetic methods 21).

We calculated genetic relatedness and pedigree relationships using maximum likelihood 37 methods in program ML-Relate 38. Pair-wise genetic relatedness (Rxy) ranges from 0 to 1 representing the proportion of allelic composition shared between individuals x and y 39. Theory suggests first-order relatives (full siblings or parent-offspring pairs) should share half their genetic makeup (i.e., Rxy = 0.5). Half siblings or grandparent-grandchild pairs, termed second-order relatives, would be expected to share only a quarter of their ancestry (i.e., Rxy = 0.25). We evaluated the importance of relationship classes, in addition to continuous Rxy values, using three classes of relatedness: first order kin (Rxy of 0.51–1), second order kin (0.26–0.5), and unrelated (0–0.25).

Age and Sex

Age and sex categories were male fawn (B), female fawn (G), yearling female (Y), and adult female (A). Each deer-pair was
assigned an age-sex class category corresponding to the ages of each deer in the pair, for example: ‘AA’ for two adult females, and ‘BY’ for a male fawn-female yearling pair. We also created an alternate age-class variable consisting of only fawns (F) regardless of sex, yearling females, and adult females. We used May 10 to define a new year in the analysis, at which point fawns were transferred to the yearling class, and yearlings to the adult class.

Statistical Analyses

Overlap values derived from logit-transformed VIs were related to predictor variables including relatedness (continuous Rxy values and kinship categories) and age-sex classes using linear mixed effects models and maximum likelihood estimation (MLE) [40] with the nlme package in program R [41]. Fixed effects included relatedness and deer-pair age, while random effects were deer-pair nested within capture group (all deer captured within 1500 m). We used random effects for capture group and deer-pair to account for potential spatial autocorrelation and lack of independence for deer with multiple pairs, respectively. We first fit a global model using restricted maximum likelihood (REML), and tested the importance of fixed effects, with models selection based on Akaike’s information criterion adjusted for small sample sizes (AICc) [43] and AIC weights. We used odds ratios to determine effect sizes for predictor variables.

Initial models failed to converge due to very low representation of some age-class capture group combinations. Because of this, we created two subsets of the data in which all age-sex class-capture group combinations contained sufficient data. The first, dataset_adult, included only those deer-pairs with at least 1 adult, excluding deer-pair age classes such as fawn-yearling and fawn-fawn, which were absent in some capture groups. However, it does include representation from all 7 capture groups. The second, dataset_capgroup, included only deer from capture groups 1 and 2, which contained the majority (70.4%) of all deer-pairs, including at least 27 deer in each age-pair type.

Results

Over the 13 loci, no errors were found in the genotyping of the 32 repeated individuals. No deviations from Hardy-Weinberg equilibrium were found after Bonferroni correction for multiple loci tests. The locus set was highly variable, yielding sufficient power to distinguish among closely related individuals (PID = 7.18E-18, and PIDsibs = 1.30E-06).

Dataset_adult consisted of 668 deer-pairs with at least one adult. Dataset_capgroup consisted of 615 deer-pairs from capture groups 1 and 2. We tested a sequence of age and Rxy models, based on a priori knowledge of deer biology, for each of our two datasets (Table 1). Likelihood ratio tests applied to our global model indicated that both capture group and deer-pair (nested within capture group) contained significant explanatory power as random effects (capture group $\chi^2$ = 4.29, $p$ = 0.04; deer-pair $\chi^2$ = 23.84, $p$<0.001) so both were used in all subsequent models.

Deer relatedness (Rxy) had a positive association with spatial overlap (Figure 1, 2); however, this association had poor explanatory power (R$^2$<0.10). Based on AIC values, the best models were those containing both kinship categories and age classes. The second best model for dataset_adult contained kinship categories and age-sex classes that differed for male and female fawns. This model has lower support from the data, as it was separated by 1.95 AIC units, with 27% of the total model weight compared to 71% for the top model. All other models were separated by at least 7.6 AIC units with ≤2% of the total model weight indicating virtually no support. For dataset_capgroup, all other models were separated from the top model by ≥4.69 AIC units, with ≤8% of the total model weight, indicating very low support from the data.

For adult deer (dataset_adult), first order kin had significantly higher spatial overlap (Odds Ratio = 32.46±95% CI: 5.14–204.87) than unrelated deer (Table 2). Second order kin also had much higher overlap than unrelated deer (OR = 5.42, CI: 1.17–25.00). Adult females also had higher spatial overlap with fawns than with other adults (OR = 3.06, CI: 1.35–6.98) or than with yearlings, but adult-yearling pairs were not significantly

| Dataset | Variable | Estimate | SE | Odds Ratio | Lower 95% CI | Upper 95% CI |
|---------|----------|----------|----|------------|--------------|--------------|
| Adult   | Intercept| -2.76    | 0.55|            |              |              |
|         | Second Order Kin | 1.69   | 0.78| 5.42       | 1.17         | 25.00        |
|         | First Order Kin | 3.48   | 0.94| 32.46      | 5.14         | 204.87       |
|         | Age- AF  | 1.12     | 0.42| 3.06       | 1.35         | 6.98         |
|         | Age- AF  | -0.72    | 0.38| 0.49       | 0.23         | 1.03         |
| Capgroup| Intercept| -3.19    | 1.01|            |              |              |
|         | Second Order Kin | 1.64   | 1.03| 5.16       | 0.68         | 38.81        |
|         | First Order Kin | 3.86   | 1.23| 47.47      | 4.26         | 528.90       |
|         | Age- AF  | 0.88     | 0.64| 2.41       | 0.69         | 8.45         |
|         | Age- AF  | -1.05    | 0.55| 0.35       | 0.12         | 1.03         |
|         | Age- YY  | -1.05    | 0.74| 0.35       | 0.08         | 1.49         |
|         | Age- YF  | 0.20     | 0.68| 1.22       | 0.32         | 4.63         |
|         | Age- FF  | 2.43     | 1.00| 11.36      | 1.60         | 80.64        |

Table 2. Parameter estimates from top models used to predict white-tailed deer spatial overlap in Wisconsin.


Different (p>0.05) from adult female pairs (Table 2). For dataset

**capgroup (all ages),** first (OR = 47.47, CI: 4.26–528.90) and

second (OR = 5.16, CI: 0.68–38.91) order kin again had

significantly higher overlap than unrelated deer (Table 2).

Among the different age classes, fawn-fawn pairs had significantly

higher overlap than did adult females (OR = 11.36, CI: 1.60–80.64).

In addition, adult-fawn pairs tended to have higher spatial overlap

than adult-yearling or yearling-yearling pairs (Table 2).

**Discussion**

Social interactions, as well as group membership, may influence

transmission of wildlife diseases [1,2] and relatedness may be more

important for transmission than simple proximity [21]. However,

proximity can be a poor surrogate for relatedness [14] or group

membership [6]. While related female white-tailed deer form

social clusters on the landscape [9,21,44], social groups may

overlap in space, but not in time. Thus, proximity of deer alone is

not enough to discern relatedness, and by extension, the likelihood

of transmission of infectious diseases [21]. Even adult females and

fawns trapped in the same location are not always mother-

offspring pairs [45]. The mechanisms by which related deer

transmit infectious disease to one another are unclear, however.

Because volume of intersection is a useful predictor of both direct

and indirect contact rates in deer [6], it appears that related deer

are more likely to come into contact, and therefore drive the

dynamics of infectious diseases [21]. We identified a clear

relationship between overlap (as measured by a volume of

intersection) and relatedness for white-tailed deer in south-central

Wisconsin. In addition, we found that age of deer influenced

degree of overlap, with adult-fawn, yearling-fawn, and fawn-fawn

pairs overlapping more strongly, whereas adult-adult pairs, and

adult-yearling and yearling-yearling pairs exhibited lower overlap.

Kinship categories were stronger predictors than continuous Rxy

values, suggesting that deer beyond a certain degree of relatedness

exhibit higher amounts of overlap. Even within kinship

classifications, such as half-siblings or parent and offspring, there is

variation in the proportion of shared DNA, and thus degree of

relatedness may be less important than the nature of the social

relationship between individual deer (e.g., parent offspring vs.

cousins).

We found that first order kin had 32.5 times as much overlap as

unrelated deer. This value is somewhat larger than a previous

finding that deer in Illinois had 5.0–22.1 times greater odds of

direct contact when they belonged to the same social group, as

estimated by proximity [6]. However, our results may be closer to

a separate study in Wisconsin indicating that deer were >100

times more likely to become infected with CWD when a highly

related infected female was in close proximity, with much lower

effects from proximal unrelated animals [21]. This indicates that

probability of CWD infection is likely higher among closely related

deer, because they have much higher contact rates, as opposed to

unrelated deer that simply share space, but have lower contact

rates. In addition, a higher probability of transmission may occur

because of the more intense nature of contacts among related deer

[46,47]. Previous observational studies indicate that parent-

offspring pairs engage in significant contact during the first year

of life [46]. Studies that investigate the spatial dynamics of disease

transmission in wild populations should include direct observation

of deer behavior to more thoroughly address the heterogeneous

disease transmission that result from the social structure of deer

[22].

Our finding that adult-fawn pairs had higher overlap is not

surprising given patterns of maternal care in white-tailed deer [8].

The average VI of probable parent-offspring pairs (adult-fawn

pairs with Rxy values >0.5) were very high (0.64 in dataset

adult, 0.53 in dataset capgroup), compared to the overall mean (0.22 in

dataset adult, 0.20 in dataset capgroup). Adult-fawn pairs with moder-

ate relatedness value (0.26< Rxy <0.5) exhibited lower overlap

(0.36 in dataset adult, 0.36 in dataset capgroup). Those pairs with low

(Rxy ≤0.25) relatedness values had VI values approximating the

overall mean (0.24 in dataset adult, 0.22 in dataset capgroup,

respectively). In this system yearling females rarely disperse from

their natal home range [24,25]. However, yearling females

sometimes establish home ranges on the periphery of their

mother’s home range once they breed [5], which may help

explain slightly reduced overlap between adult and yearling deer.

Female white-tailed deer are highly philopatric, characterized

by stable home ranges with a high degree of overlap among

individuals within social groups [6,9,44,46]. However, social

structure of deer is less typical where rates of harvest are high

and age structure is biased towards young animals [14,48].

Nonetheless, we found that overlap (as measured by VI) closely

associated with degree of relatedness, providing evidence for social

structure at a local scale in spite of heavy harvest pressure.

While ongoing disease eradication efforts may have temporarily

increased deer harvest, this population has been subjected to

ongoing harvest for many years, and CWD control efforts are

unlikely to have produced the patterns observed. The strong

matriarchal social structure of female white-tailed deer likely

prevents homogenous mixing of individuals [6,9] and homoge-

neous CWD transmission among members of different social

groups [21].

The degree to which deer contact each another varies seasonally

[6, 56]. However, to ensure sufficient observations, our analyses

were based on annual data and provide no insight into seasonal

patterns. In addition to relatedness, hotspots of activity such as

scrapes, rubs, feeding/baiting sites, and mineral licks also likely

play a role in contact rates of cervids and potential disease

transmission [49,50,51]. We did not identify such features in our

study and have no basis to evaluate the contribution of these

behavioral hotspots to potential transmission of disease. Deer may

also be more likely to overlap in agricultural areas due to

concentrated food sources [3,16,25,52]. In fragmented systems,

deer would likely congregate closely in areas of remaining

resources, particularly in seasons when food is limited [52].

Unfortunately, accuracy of the spatial locations in this study was

insufficient to investigate the effects of habitat use, given that the

study area is a complex mosaic of forested and agricultural land

[25]. Our study focused on female deer because they are most

often targeted for population control and, unlike males, rarely

engage in long-distance movements [9,21,25]. However, males are

more frequently CWD positive than females (Greer et al. 2006),

and long-distance movements by males may be important in the

geographic spread of CWD.

CWD can be transmitted both directly (by deer-to-deer contact)

and indirectly (via contamination of the environment), though the

importance of these modes of transmission in the wild are

unknown [17,19,21]. VI provides a metric for both direct and

indirect contact, though the spatial-temporal resolution of our data

is insufficient to differentiate these specific events. While it is

possible for two deer who overlap in space to avoid direct contact,

indirect contact is virtually guaranteed, particularly given the likely

occurrence of congregation points such as scrapes, rubs, feeding/

baiting sites, and mineral licks. However, given previous findings

that contact rates vary predictably with VI [6], we believe our

findings likely apply for both direct and indirect transmission

scenarios. Our findings support previous research that suggest
CWD should spread more rapidly among related deer [21]. As such, control of large related social groups may be an effective strategy in slowing pathogen transmission, particularly given that there is little evidence that female harvest impacts movement behavior [16]. We also found limited overlap among unrelated deer suggesting that disease spread among social groups, which is needed to sustain disease, may occur between neighboring social groups. We believe the rate and mechanisms of disease transmission between adjacent social groups is an important area for future research.

We provide an important step in understanding the mechanisms underlying observed patterns of CWD transmission, namely, that related individuals are more likely to come into close proximity on the landscape, where disease transmission may occur either directly or indirectly. Further progress in understanding the specifics of disease spread will be necessary to devise practical strategies for deer management.

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Author Contributions

Conceived and designed the experiments: SM MS TVD SR NM. Performed the experiments: SM MS. Analyzed the data: SM SR. Wrote the paper: SM MS TVD SR NM.

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