Muskrats as a bellwether of a drying delta

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Wetlands worldwide are under threat from anthropogenic impacts. In large protected North American areas such as Yellowstone and Wood Buffalo National Parks, aquatic habitats are disappearing and wetland-dependent fauna are in decline. Here we investigate population dynamics of an indicator species in Canada’s Peace-Athabasca Delta (“the delta”), a World Heritage Site. Based on population surveys, habitat mapping and genetic data from 288 muskrats, we use agent-based modeling and genetic analyses to explain population expansion and decline of the semi-aquatic muskrat (Ondatra zibethicus). Simulations quantify a large population (~500,000 individuals) following flood-induced habitat gains, with decreased size (~10,000 individuals) during drying. Genetic analyses show extremely low long-term effective population size (Nₑ: 60–127), supporting a legacy of population bottlenecks. Our simulations indicate that the muskrat population in the delta is a metapopulation with individuals migrating preferentially along riparian pathways. Related individuals found over 40 km apart imply dispersal distances far greater than their typical home range (130 m). Rapid metapopulation recovery is achieved via riparian corridor migration and passive flood-transport of individuals. Source-sink dynamics show wetland loss impacts on the muskrat metapopulation’s spatial extent. Dramatic landscape change is underway, devastating local fauna, including this generalist species even in a protected ecosystem.
lobally, between 50 and 87% of wetlands are estimated to have been lost since 1700. In the 5500-km² Peace-Athabasca Delta (“the delta”) part of Canada’s Wood Buffalo National Park and a World Heritage Site with “In Danger” status pending, one of the world’s largest freshwater deltas has lost critical habitat. Previous work has attributed drying in the delta to the effects of climate change, upstream hydropower development on the Peace River, or a combination of both drivers. This habitat loss is concurrent with a dramatic decline in the delta’s muskrat (Ondatra zibethicus) population over the past half century, a trend long reported by local Indigenous trappers.

Muskrats serve as an ecolohydrologic indicator species. The muskrat shares habitat requirements of other wetland species, including fish and waterfowl, requiring vegetation maintained at early successional stages by flooding. Their numbers rise and fall rapidly with changes in the floodplain as they thrive on near-shore vegetation before declining rapidly in “die-offs”, often attributed to their intensive herbivory. Muskrats are key grazers in this system, influencing succession, diversity and productivity of aquatic and emergent vegetation and are important prey for numerous predators.

Despite their ecological significance, the population dynamics of muskrats at the scale of the floodplain remain poorly characterized. Indigenous land users in the delta have observed ecological changes in the delta that threaten their way of life, including fewer spring flooding events and increased desiccation of wetlands, with a decline in wetland species including the muskrat. For generations, muskrat have served as a source of food, fur and income for land users in the delta. Indigenous knowledge holders and experienced trappers are extremely interested in how ecosystem-scale flooding results in rapid muskrat population increases. Based on satellite remote-sensing temporal mapping of habitat, muskrat surveys, and genetic data from across the delta, we conduct independent investigations using agent-based (individual-based) modeling of muskrats and genetic analyses to reveal a likely mechanism for muskrat eruption and die-off. In addition, we investigate links between eruptive periods in population growth, longer-term drying, and intermittent flooding.

We developed a large-scale (computationally intensive), agent-based model of the muskrat population in the delta that simulates the combined life sequences of the total population ranging from 10,000 to >500,000 individuals for the period 1971–2017, with population survey data available for comparison in 21 years over the period 1970–2017. Life events of individual female muskrat are represented, including dispersal, home range establishment, reproduction, and mortality events, repeating annually. Muskrat reorganize their home range locations in the spring “shuffle” when they search for a mate and new territory for the coming year. The total population is twice the number of modeled females. Individual dispersal behavior was represented as a constrained random walk that penalized travel over land and favored travel through hydrologic features such as rivers and lakes.

Temporally dynamic habitat maps were created from an atlas of open water and shoreline habitat built from 30m- to 60m-resolution Landsat satellite imagery available for the period 1972–2017. Canny Edge Detection was used to create maps of land, shoreline and water classes. These maps were used to generate annual habitat suitability maps, consisting of a mosaic of viable riparian habitat regions interspersed with regions of no habitat value to muskrat. Habitat maps were discretized into a modeling grid containing over three million hexagons with individual hexagons of width 60 m (0.31 hectares).

Thirty model realizations were run over the period 1972–2017 to simulate recent periods of eruptive population growth and die-off. Modeling results were compared to population survey records comprised of 21 years of surveys counting muskrat houses at sites across the delta. Dispersal flux, represented as the number of individuals that have migrated through a given location, was mapped as the difference between dispersal flux in 2016 and 2015 for comparison to genetic data. Population sources and sinks, measured in productivity (the number of births minus deaths in a location) were mapped for three successive periods of eruptive population growth (1971–1972, 1996–1997, 2014–2015) and subsequent die-off (1975–1976, 1998–1999, 2016–2017). Model results were then compared to muskrat genetic data.

Microsatellite genetic analyses served as an independent measure of muskrat population history, structure, and dispersal patterns in the delta. Muskrat tissue samples were donated by trappers in the winter of 2015/2016 (hereafter referred to as 2015 samples) and the winter of 2016/2017 (hereafter referred to as 2016 samples), corresponding to habitat conditions mapped in the 2015 and 2016 open water seasons and model output for 2015 and 2016. Nine autosomal microsatellite loci were PCR-amplified (primers from Laurence et al., 2009) in 200 and 88 samples from the 2015 and 2016 trapping seasons, respectively.

Effective population size ($N_e$), the number of individuals in an idealized population producing the level of genetic diversity measured from the sample population was estimated using NeEstimator. $N_e$ is sensitive to past fluctuations in population size and reflects the harmonic mean of the population size over time.

First-order relatives (parent-offspring or full sibling pairs) were detected by cross referencing output from three different programs: Colony v 2.0.6.5, ML-Relate, and Coancestry. Population structure was assessed using the Bayesian clustering approach implemented in the program STRUCTURE v2.3.4. STRUCTURE identifies subpopulations and probabilistically assigns each sampled individual to one or more subpopulations. Pairwise genetic difference between sites within and between years was assessed by calculating $R_{ST}$ in Arlequin. An 872 bp region of cytochrome b was also PCR-amplified and sequenced from tissue samples. Number of haplotypes, number of polymorphic sites, and nucleotide and haplotype diversity were calculated using DnaSP v6.11.01. Median joining networks for cytochrome b haplotypes from each year separately as well as both years together were constructed using the program PopArt.

Combined agent modeling and genetic analysis results indicate that the muskrat population in the delta is a metapopulation with individuals migrating preferentially along riparian pathways. Genetic results showing related individuals found over 40 km apart imply dispersal distances far greater than the typical home range for muskrat (130 m). These findings suggest that rapid metapopulation recovery is achieved by the long-distance migration of individual muskrat, primarily through riparian corridors in the delta, as well as the passive transport of individuals during large flood events. Results further show the effects of wetland loss on the muskrat metapopulation’s total size and spatial extent. These findings suggest that landscape change is driving a reduction in both the total population size and spatial extent of the muskrat metapopulation of this inland delta.

**Results**

**Eruptive dynamics.** Agent-based simulations show that the number of muskrats in the delta is characterized by a series of peak population values followed by sharp declines in the
population, concordant with the population survey record (Fig. 1c). Total modeled population size—obtained by doubling model output of the number of females at each time step—ranges from a minimum value of 10,010 individuals in 2009, to a maximum population size of 546,619 in 1972, with values obtained by taking the median across 30 model realizations. Successive modeled periods of eruptive population growth (1971–1972, 1996–1997, 2014–2015) show temporally declining peak population values that reproduce declines in peak population values in the population survey record (Fig. 1c). However, based on genetic analyses, measures of effective population size, $N_e$, using NeEstimator are very low, at 103.7 individuals (95% confidence interval = 88.2–123.5) (6 sites) for 2015 and 61.4 individuals (95% confidence interval = 50.2–76.8) (4 sites) for 2016 (Supplementary Table 1).

Using agent model results, the total modeled population size at four randomly selected sites was examined in 2009, the year with the lowest total modeled population size (Supplementary Fig. 1). Results show sites whose local population has gone extinct (D, B), one site where the local population goes extinct in most but not all realizations of the model (E), and one site whose local population consistently survives, indicating site-dependent local extinction.

**Dispersal behavior and genetic relatedness.** Agent simulation results for dispersal show enhanced dispersal flux values in 2016 versus 2015 (Fig. 2b). Although there are some areas of decreased dispersal in 2016 compared to 2015 (red regions in Fig. 2b), we interpret this model output with a wider focus. Namely, that this area of the delta is predominantly experiencing greater dispersal in 2016 than in 2015 (blue regions in Fig. 2b), particularly in the area between sites sampled in 2016 (site A, B, D, and E). In this central region of the delta, this enhanced dispersal is coincident with genetic results showing an increase in the number of related individuals across sample sites and a decrease within sites (Fig. 2c, d). Genetic relatedness results indicate that in 2015, out of the 23 first-order relationships identified, 19 (82.6%) were between individuals at the same site, versus only 2 (18.2%) out of 11 such relationships in 2016. (Fig. 2c, Supplementary Table 2). Coincident with this increase in related individuals across sites, microsatellite data for 2016 relative to 2015 show a decrease in population structure and an increase in homogeneity (Fig. 2a, Supplementary Fig. 2). This trend is corroborated with RST values showing significant pairwise genetic differences between sites in 2015 but not in 2016 (Supplementary Fig. 3). Cytochrome b analysis shows a greater haplotype diversity in 2016 (0.88) than in 2015 (0.67) with a general mixing of haplotypes across all sampled locations (Supplementary Fig. 4).

Agent model simulation results in the des Rochers River region (indicated in Supplementary Fig. 5a) show enhanced dispersal in 2016. Dispersal occurs preferentially through hydrologic features, in this case the des Rochers River (Supplementary Fig. 5b). Results of genetic relatedness analysis, meanwhile, show three related pairs of individuals detected between site F, located in the des Rochers region, and two sites in the central region (Fig. 2d). These results, indicating the presence of related pairs of individuals tens of kilometers apart, suggest that long-distance dispersal by individual muskrats is occurring on the order of tens of kilometers per year. Since muskrat in the delta typically live for two years or less (less than 5% of individuals reach 3 years of age), these distances were likely covered in one or two discrete, long-distance spring dispersal events. This finding is consistent with the model depiction of long-distance migration of individual muskrat through preferential hydrologic pathways.

**Source-sink dynamics.** Productivity maps generated from agent model results, showing the number of births minus the number of deaths for muskrat in a given location, indicate locations contributing to net population growth (positive values of productivity) or to net population decline (negative values of productivity). Results show the delta landscape transitioning on an interannual basis from a state of having many landscape features acting as population sources (years of net population growth in 1971, 1996, and 2014, shown in the top panel of Fig. 3) to the landscape having many population sinks (years of net population decline in 1975, 1998, and 2016, shown in the bottom panel of Fig. 3). These modeled states correspond to eruptive population growth after flooding and post-flood die-offs, respectively. Comparing successive years of net muskrat population increase, the area of the delta serving as a source or sink has decreased from 79% to 55% to 42% in each of the periods 1971–1972, 1996–1997, 2014–2015, respectively (Fig. 3). This indicates a long-term decline in the area where muskrats are present, whether born or dying in a given location. Comparing the 2014 population increase year to prior years of increase (1971 and 1996), the last year of net population expansion was less productive for muskrat than in previous years. These comparisons also show that in years of net population increase or decrease, multiple locations serve as sources or sinks (Fig. 3).
**Fig. 2 Genetic population structure, agent model dispersal behavior, and genetic relatedness results for muskrat in the delta.**

a) STRUCTURE results assign fractions of the ancestry of individual sampled muskrat (shown with vertical bars) to population 1 (purple) or population 2 (green), with individuals ordered by year and sample site. The vertical axis represents the fraction of ancestry from either population, ranging from 0 to 1.

b) Dispersal flux in the agent model, the number of dispersing individual muskrat traveling through a given location, is shown as the difference between dispersal flux in 2016 and 2015 for a portion of the delta encompassing genetic sample sites A through E. The map indicates locations of increased dispersal (blue), locations of decreased dispersal (red), and locations where the amount of dispersal is unchanged across the two years (white). Colored circles show genetic sampling site locations. Colors indicate the year of sampling according to the color scheme in Fig. 1b. Dispersal flux data shown here is the mean of $n = 30$ model realizations.

c) In each year, the number of muskrat sampled and the number of first order relationships. First order relationships are made up of pairs of individuals found within the same sampling site (gray) or pairs of individuals from different sampling sites (blue).

d) The per mil of first order relationships observed out of all possible relationships within (gray) or between (blue, along arrows) sampled sites in each year. Distances between sites are not to scale. Sites not sampled in a given year are grayed out.

**Fig. 3 Productivity maps for muskrat in the delta during successive years of net population growth and decline.**

Maps of productivity (the number of births-deaths of muskrat in each 0.28 km$^2$ area) in the delta indicate the locations and strength of sources and sinks in the delta for three successive years of net population growth (1971–1972, 1996–1997, 2014–2015) and subsequent net population decline (1975–1976, 1998–1999, 2016–2017), averaged across $n = 30$ model realizations.
Discussion

Agent modeling results show muskrat population dynamics characterized by periods of eruptive population growth followed by die-offs that emerge from the simulated life sequence of up to 546,619 individuals on the floodplain (Fig. 1c). However, microsatellite analyses show extremely low effective population size values (on the order of tens to hundreds of individuals in 2015 and 2016), likely the consequence of repeated die-offs of muskrat resulting in genetic bottlenecks. The agent model and genetic results consistently illustrate a population characterized by rapid eruptive growth and massive die-offs.

We find greater modeled dispersal flux, greater relatedness across sample sites, and less genetic structure in 2016 compared to 2015. Interestingly, 2016 has a smaller modeled population size compared to 2015, with critical habitat and the on-the-ground population density decreasing over the same period10. This suggests that muskrat must disperse further under conditions of overcrowding and degraded habitat, with many dying off in the search for suitable habitat.

Our simulations indicate that the delta’s muskrat population is a metapopulation, comprised of many populations from permanent and ephemeral wetlands across the delta. Though local populations—populations of muskrat in subregions of the delta such as the subregions whose populations were tracked in the model (Supplementary Fig. 1b)—may go extinct (=sink) during dry periods, some “seed” populations persist. Following a flood event and ensuing increase in habitat availability and connectivity, the seed (=source) population repopulates sites where habitat has been replenished. While a small number of muskrat likely migrate into the delta during spring dispersal events annually via the rivers that drain into the delta, model results show that local surviving populations of muskrat within the delta numbering in at least the tens of thousands serve as the primary “seed” that disperses and supports rapid population resurgence following flood events. Sites with nonzero local populations thus sustain the delta’s muskrat metapopulation through dry periods between flood events (Fig. 1b, c, Supplementary Fig. 1b).

Simulation and genetic results independently support our conclusion that repopulation requires the occurrence of long-distance dispersal of muskrat in the delta. Both lines of analysis point to long-distance dispersal by muskrat such that newly replenished habitat following flood events results in ensuing eruptive growth (Fig. 1a, c). Genetic results show related muskrat pairs tens of kilometers apart, confirming long-distance dispersal behavior across the floodplain (Fig. 2d). Agent model results illustrate that muskrat use hydrologic features during long-distance migration (Supplementary Fig. 5). We conclude that the mechanism for rapid increases in muskrat abundance from low values after flood waters have subsided is a viable metapopulation that sends successful propagules from “seed” populations to extirpated sites that have been newly replenished. Given the high reproductive rate of muskrat (up to 15 female offspring per reproductive rate of muskrat (up to 15 female offspring per

The source-sink networks show a reduction in spatial extent of the species on the floodplain, even in years of peak abundance, due to recent dramatic declines in habitat availability (Fig. 3). This finding is consistent with recent hydro-limnological analysis showing reduced effects of 2014 flooding compared to past events2. The multitude of areas of the delta that serve as sources or sinks in any given year of substantial increase or decrease in muskrat numbers highlights that no single portion of the delta is most important for muskrat persistence. These results have a conservation implication: if it becomes necessary, reintroduction of muskrats may be successful in a multitude of water bodies in the delta given their long dispersal range as long as release occurs into a body of water with appropriate vegetation and as long as the water body persists through reproduction.

These findings have larger significance. Given that there are empirical and anecdotal reports of muskrat population declines across North America14–26, future work should investigate landscape change and wetland habitat loss as possible drivers of changing muskrat abundance at additional sites across the continent. In addition, the story of muskrats in the delta has broader implications for the delta ecosystem as a whole. Habitat loss, among other drivers, is implicated in the widespread documented decline of North America’s avifauna27. Over one hundred bird species, many migrating from across North America, breed at the delta and have similar habitat requirements to the muskrat28. While previous research on aquatic birds in the delta has investigated contaminant levels in colonial waterbird eggs, our results suggest that quantifying the effects of land cover change in the delta on waterbird populations would be a fruitful avenue for future studies29,30. These species are also likely being adversely affected by the overall decline of freshwater distribution and abundance in the delta, as are hundreds of native plants, fish, insects, and terrestrial animals. Our results have conservation implications for extremely large, remote regions where landscape change is putting local fauna at risk, even in protected ecosystems.

Methods

Agent modeling. The agent model for muskrat in the delta was developed using HexSim, an agent-based ecological model that allows for spatially explicit simulation of wildlife population dynamics12,31. The HexSim agent model of muskrat incorporated the entire delta in a modeling grid containing 1717 rows of hexagons by 1760 hexagons per row, for a total of 3,021,920 hexagons. Operating on an annual time step, the model tracked up to 273,310 females annually through their life cycles from 1971 to 2017. Given the computational intensity of the model (a runtime of ~16 h per realization), the number of realizations was limited to thirty after examination of model output for the ensemble. Boxplots showed good agreement across model realizations in the timing and magnitude of population peaks, die-offs, and years of low abundance, as well as normally distributed total population size in the majority of years simulated, suggesting that the central tendencies for total population size, dispersal and productivity maps were adequately captured (Supplementary Fig. 1a).

An initial population size for the delta was estimated using an observed muskrat “house” count at a well-studied site, Egg Lake. Records for 1971 show 179 houses, yielding an initial population size of ~484 females at that lake. This estimate was scaled up to a population estimate for the entire delta by accounting for the fraction of critical habitat in the delta occupied by Egg Lake in 1972 (4.88 km² out of 651.77 km²) to yield an initial population of 59,701 females for the entire delta.

Muskrat movement behavior. The delta model was developed to account for three broad categories of spring movement behaviors for individual muskrat:

(i) Local movements during spring dispersal

To represent the spring shuffle within the home ranges of muskrat at their home lake, an “exploration event” allows every individual to search their local surroundings (up to 500 hexagons, or 1.6 km²), with the goal of establishing a home range. Individuals that succeed establish a home range and finish the movement event. Individuals that are unsuccessful at establishing a home range, engaged in long-range dispersal, described in (ii) below. In the spring, muskrat home ranges typically shuffle within a given water body at the onset of breeding12,23. Home range adjustments are typically at the scale of several hundred meters away from previous territory13.

(ii) Long-distance movements during spring dispersal

The outcome of this process is a new disrtibution of individuals within the delta, with new, initialized home ranges and an estimate of the number of new home ranges established. The model then accounts for the number of new home ranges established and the number of existing home ranges lost.

(iii) Long-distance movements during spring dispersal

Lastly, the model simulates the establishment of new home ranges in addition to the movement of individuals within existing home ranges.

(iv) Long-distance movements during spring dispersal

The model simulates the movement of individuals within and between home ranges, resulting in the establishment of new home ranges and the loss of existing home ranges.

(v) Long-distance movements during spring dispersal

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The model simulates the movement of individuals within and between home ranges, resulting in the establishment of new home ranges and the loss of existing home ranges.
(ii) Long-range spring dispersal

For individuals that do not successfully establish a home range with local movements in (i), a long-range dispersal event occurs, and it is parameterized based on literature values for muskrat dispersal rates. Based on the highest values of muskrat emigration rates (not attributed to passive transport via flooding) of 60 km/year, we set a dispersal distance of 1000 hexagons, or about 60 km of travel\(^3\). In addition, such dispersal events are constrained by the fact that muskrat movement is more limited on land than on water. Muskrats are placed to move only in the order of magnitude of the shoreline, and individuals, in water they have been observed to travel much further distances irrespective of current; for instance, a single muskrat was observed to travel 50 km “against the current” in 15 days\(^4\). We therefore infer that higher reported rates of emigration for muskrats are made up primarily of travel through surface water features, combined with an ability of individual muskrat to travel over land up to 3 km.

To model this in the first model, we first used the annual water/shoreline/land maps of the delta to generate annual dispersal maps based on a dispersal metric for particular environment categories. For these maps, water and shoreline pixels received a score of 10, and land pixels received a score of zero. This yielded dispersal maps whose hexagons have values of zero when they entirely overlap land pixels, 10 when they entirely overlap water pixels, and values in the range (0.10) for shoreline regions. Then, at each step of muskrat travel along its dispersal path, the difference of the hexagon score from 10 is evaluated and added to that individual’s dispersal penalty. Land hexagons therefore have a resistance of 10, and water hexagons a resistance of 0, with shoreline regions incurring an intermediate resistance between 0 and 10. The resistance values of encountered hexagons are tracked cumulatively for each individual while it disperses. When an individual reaches a resistance threshold of 500, the individual must stop dispersing. This resistance threshold of 500 is equivalent to 3 km of overland travel. So, an individual dispersing with a path entirely over land can go 3 km per year from their prior home range, but if their dispersal is entirely through water, then there is a travel limit of 60 km in a year.

During long-range spring dispersal, individuals follow a constrained random walk to find a suitable location to settle. When selecting the adjacent hexagon to explore, individuals prefer hexagons with values between 2 and 10 (shoreline and water hexagons) at the expense of hexagons with values between 0 and 1 (land or mostly land hexagons), and they are influenced by their prior direction of travel with autocorrelation of 50%. At the completion of their long-range dispersal, individuals repeat the local movement exploration event to search for a suitable location to settle within their newly discovered home range. Individuals that do not succeed are removed from the simulation, representing death because they did not successfully establish a home range after long-range dispersal and succumbed to predation or starvation, or representing that they have migrated out of the delta.

(iii) Enhanced dispersal due to flooding

In years of known, large-scale flooding in the delta (1972, 1974, 1996, 1997 and 2014), a flood dispersal event is applied to simulate the effects of flooding on muskrat dispersal. A dispersal map is applied in which all hexagons in the delta have a value of 10, such that there is no resistance penalty for movement (a resistance value of 0) and the resistance threshold described in (ii) is never reached. When determining the range of distances for dispersal as a function of floodwaters, we drew on literature values. While some muskrat remain in the water and disperse during flooding, yielding emigration rates of up to 120 km/year, others find refuge in trees or on rafts that are swept into trees and floodwaters, we drew on literature values. While some muskrat tissue samples for this study consisted of ≤2 mm diameter pieces of tail tissue donated by trappers from muskrats trapped for fur and meat during the 2015 and 2016 trapping seasons (November to May). All trapping resulting in donated samples was done so legally under all necessary permits or by individuals with Indigenous trapping rights in the delta as recognized by Treaty 8. All muskrats were collected from the Peace-Athabasca Delta and the lake on creek of collection was noted by the trappers. A total of 200 muskrats were collected from 6 sites in the 2015 trapping season and 88 muskrats were collected from 4 sites in the 2016 trapping season (Fig. 1b). This study was designed to capitalize upon muskrat trapping that was already taking place in the delta, thus, we had no control over the number of samples donated nor the specific location within the delta where samples were collected.

Dispersal flux mapping. Dispersal flux, the number of individuals passing through a given location per year, was mapped as the difference in values for the two years in which genetics data were collected, 2015 and 2016 (Fig. 2b). This was done by first exporting hexagon-based dispersal flux tallies for all thirty realizations in the years 2015 and 2016. Then, the mean value of dispersal flux across all 30 realizations was calculated to produce a single average dispersal flux map for each year. Finally, the difference between these two maps was calculated to yield the difference map showing locations of increased, decreased, or unchanged dispersal flux shown in Fig. 2b.

Genetic analysis

Muskrat tissue samples for this study consisted of ≤2 mm diameter pieces of tail tissue donated by trappers from muskrats trapped for fur and meat during the 2015 and 2016 trapping seasons (November to May). All trapping resulting in donated samples was done so legally under all necessary permits or by individuals with Indigenous trapping rights in the delta as recognized by Treaty 8. All muskrats were collected from the Peace-Athabasca Delta and the lake on creek of collection was noted by the trappers. A total of 200 muskrats were collected from 6 sites in the 2015 trapping season and 88 muskrats were collected from 4 sites in the 2016 trapping season (Fig. 1b). Supplementary Fig. 6). This study was designed to capitalize upon muskrat trapping that was already taking place in the delta, thus, we had no control over the number of samples donated nor the specific location within the delta where samples were collected.

DNA extraction and amplification. DNA was extracted from tissue samples using the DNeasy Blood and Tissue kit following the manufacturer’s protocol (Qiagen, Valencia, CA). Partial mitochondrial cytchrome b sequences of 872 base pairs were amplified from samples using the same primers described in Mychall and Harrison (2014): OzbFW 5′CACCCTATTTCTGCACTCCCAAC3′; OzbREV 5′TGGG- TATGAGAATAATGATAATGGCAAAGTA3′.

Amplifications were carried out in a total volume of 10 µl made up of the following mixture: 1 µl DNA template, 5 µl GoTaq® G2 Hot Start Colorless Master Mix (Promega), 1 µl 5 µM of each primer, 2 µl water. PCR amplifications were performed in an Eppendorf Mastercycler Pro thermal cycler, with an initial denaturation at 95 °C for 15 min, followed by 30 cycles of 30 s of denaturation at 94 °C, 30 s of annealing at 57 °C, and 1 min of extension at 72 °C and a final hold at 4 °C. Negative controls were used in all extractions and PCRs as quality controls. The amplification products were electrophoresed in 2% agarose gels (200 Volts, 20 min). The DNA bands were visualized using SYBR Safe DNA Gel Stain (ThermoFisher Scientific) under UV light and product size was determined in relation to a 100 bp DNA size standard (ThermoFisher Scientific). PCR products were cleaned using ExoSAP-IT (ThermoFisher Scientific) and sequenced by Elim Biopharmaceuticals, Inc. (Hayward, CA). Sequences were aligned using Geneious 7.1.4.

Nine autosomal microsatellite loci (Oz06, Oz08, Oz16, Oz27, Oz32, Oz34, Oz41, Oz43, Oz44) were PCR amplified using primers from Laurence et al. in a total of 288 samples, 200 from 2015 and all 88 from 2016\(^5\). Amplification was carried out in a total volume of 5 µl made up of the following mixture: 0.5 µl DNA template, 2.5 µl GoTaq® G2 Hot Start Colorless Master Mix (Promega), 0.5 µl of each primer mixture (containing 2 µM of each primer), 1.5 µl water. PCR amplifications were performed in an Eppendorf Mastercycler Pro thermal cycler, with an initial denaturation at 95 °C for 15 min, followed by 30 cycles of 30 s of denaturation at 94 °C, 30 s of annealing at 60 °C, and 1 min of extension at 72 °C and a final hold at 4 °C. Microsatellite PCR product was sent to the Protein and Nucleic Acid Facility at Stanford University to conduct multiplexed fragment size analysis using an ABI 3130xl Genetic Analyzer. Fragment analysis was done with PCR products for four microsatellite loci (Oz06, Oz07, Oz32 and Oz43) multiplexed in one run and the other five microsatellite loci (Oz08, Oz16, Oz43, Oz41, Oz44) multiplexed in a second run.

Statistics and reproducibility

Agent-based model. Statistics within the model were carried out using HexSim tools with code specific to the muskrat simulation to generate model output files, all available at https://doi.org/10.17605/OSF.IO/CHNR62\(^{1,12}\). Statistics on model output to calculate mean and
median values and boxplots on model output for n = 30 realizations reported in the paper were carried out using R version 3.4.3.51.

Cytochrome b. To determine how many sequences to sample to capture all cytochrome b haplotype diversity, we plotted haplotype accumulation curves using the package Spider in R with random ordering of samples and 10,000 permutations24,45. Based on accumulations curves, we sequenced 150 samples from 2015 and all 87 samples from 2016; one sample from 2016 did not amplify (Supplementary Fig. 7).

Microsatellite calling. For all microsatellite loci, samples were PCR amplified, fragment analyzed and scored twice to assure genotyping accuracy. All scoring was conducted in Geneious v7.1.4. Three samples from 2015 were removed due to missing data and two discrepancies between the two genotyping runs. In our final dataset, 90.3% of alleles from 2015 were confirmed by two identical genotypes and 96.5% of alleles from 2016 were confirmed by two identical genotypes. The remainder were alleles that were successfully captured in only one fragment analysis and were missed in the other run due to allelic dropout. The final microsatellite dataset used for all analyses consisted of n = 197 for 2015 and n = 88 for 2016 and is provided as Supplementary Data 1.

Microsatellite analyses. We investigated the presence of null alleles with a 95% confidence interval using MICROCHECKER v2.2.3 and estimated the frequency of null alleles in each sampling site using genepop v1.0.5 in R.36,48. We also tested for null alleles in the dataset for each year considered as one population. Null alleles were indicated at numerous loci (Supplementary Table 3); however, there was no consistency in loci displaying null alleles across sampling sites and thus all alleles were kept in the analyses.

We calculated the number of alleles and allelic richness (corrected for sample size) for each sampling site in each year using FSTAT V2.9.3.2.48. We calculated the observed (H2) and expected (H2e) heterozygosity and deviations from Hardy-Weinberg equilibrium using Arlequin 3.5.2.220. When identifying significant deviations from Hardy-Weinberg equilibrium we corrected for multiple comparisons by applying the Benjamini & Hochberg method in R to control for false discovery rate46. We assessed pairwise genetic difference between sites with 14 or more samples (excluding G) by calculating R̄s in Arlequin. R̄s is similar to FST but takes allelic size into account and is considered to be more suitable for the high mutation rate of microsatellite data48.

Structure. Population structure was assessed using the Bayesian clustering approach implemented in the program STRUCTURE v2.3.4 run using the admixture model with correlated allele frequencies. 100,000 burn in and 1,000,000 MCMC iterations19.

These are the same priors used in previous studies to successfully identify population structure in muskrat using the same microsatellite markers used here45,46. We ran 20 independent runs for K from 1 to 10. STRUCTURE was run on a Linux machine and parallelized using the program StrAuto50. We used the Evanno method to identify the most probable K as the one with the largest Delta K value41. The rate of change in the log probability of the data between successive K values (Delta K) was calculated and visualized using STRUCTURE HARVESTER Web v0.6.94. Structure results were visualized using GLUMPA v2.225. We conducted these structure analyses on each year separately as well as both years together (Supplementary Fig. 2). In all STRUCTURE runs, no prior information on sampling location or sampling year is included. The Evanno method indicated that two populations (K = 2, as shown in Fig. 2a), was the most probable (Supplementary Fig. 2c).

Estimation of relatedness. We identified putative first-order relatives (parent-offspring or full siblings) by cross referencing output from three different programs—Colon v 2.0.6.5, ML-Relate, and Coancestry18–20. We ran Colony assuming polygamy in both males and females and assuming a genotyping error rate of 0.05.24 We only considered first-order relatives with a probability of 95% or higher. We further verified relatedness by only keeping relationships that were also supported by ML-Relate with a maximum likelihood estimate of relatedness of 0.495 or higher and by Coancestry with a TrioML 95% confidence interval overlapping 0.5 but not 0.125. In Coancestry, we used the triad likelihood estimator (TrioML) which uses a triad of individuals for estimating pairwise relatedness24,44.

Effective population size. Effective population size (Nₑ) with 95% confidence intervals using the parametric method was estimated using the linkage disequilibrium method in NeEstimator15,55,58. We ran NeEstimator using a random mating system and considered results when excluding alleles with frequencies less than 0.01. This estimation of Nₑ represents the effective population size of the parental generation of the samples. We also estimated Nₑ with 95% confidence intervals using the moment and likelihood methods implemented in MLNE57. We ran MLNE without assuming drift-migration equilibrium, set a maximum Nₑ value of 35750 (maximum based on computing limitations), and used both timepoints (2015 and 2016) to estimate Nₑ for both breeding seasons. These two methods of estimating Nₑ have been found to be the most accurate when considering numerous Nₑ estimation methods24.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The agent-based model and modeling results, including raw data associated with Figs. 1, 2 and 3, are available at https://doi.org/10.17605/OSF.IO/CHNR6. All cytochrome b sequence data have been submitted to the GenBank database under accession numbers MT215718 - MT215954. Figure 2 has associated raw data from the genetics analyses; that microsatellite dataset is included as Supplementary Data 1. Muskrat tissue samples are available from E.A. Hadly upon reasonable request.

Code availability

Custom code developed for the muskrat agent-based model in HexSim Version 4.0.13.0 is available at https://doi.org/10.17605/OSF.IO/CHNR6.

Received: 28 October 2020; Accepted: 2 June 2021; Published online: 24 June 2021

References

1. Riordan, B., Verbyla, D. & McGuire, A. D. Shrinking ponds in subarctic Alaska based on 1950-2002 remotely sensed images. J. Geophys. Res. Biogeosci. 111, (2006) https://doi.org/10.1029/2005JG000150.
2. McMenamin, S. K., Hadly, E. A. & Wright, C. K. Climatic change and wetland desiccation cause amphibian decline in Yellowstone National Park. Proc. Natl. Acad. Sci. 105, (2008) https://doi.org/10.1073/pnas.0809909105.
3. Remmer, C. R., Klemt, W. H., Wolfe, B. B. & Hall, R. L. Inconsequential effects of flooding in 2014 on lakes in the Peace-Athabasca Delta (Canada) due to long-term drying. Limnol. Oceanogr. 63, (2018) https://doi.org/10.1002/lo.10787.
4. Davidson, N. C. How much wetland has the world lost? Long-term and recent trends in global wetland area. Mar. Freshw. Res. 65, (2014) https://doi.org/10.1016/j.mfr.2014.07.007.
5. Wolfe, B. B. et al. Reconstruction of multi-century flood histories from oxbow lake sediments, Peace-Athabasca Delta, Canada. Hydrolog. Process. 20, (2006) https://doi.org/10.1002/hyp.6423.
6. Beltsos, S. Comparing the impacts of regulation and climate on ice-jam flooding of the Peace-Athabasca Delta. Cold Reg. Sci. Technol. 108, (2014) https://doi.org/10.1016/j.coldregions.2014.08.006.
7. Beltsos, S. & Peters, D. L. Commentary on "Past variation in Lower Peace River ice-jam flood frequency" by Wolfe et al. (2020). Environ. Res. Lett. 25, 560–566 (2020).
8. Wolfe, B. B., Hall, R. L., Wiklund, J. A. & Kay, M. L. Response to Commentary by Beltsos and Peters on "Past variation in Lower Peace River ice-jam flood frequency" by Wolfe et al. (2020). Environ. Res. Lett. 25, 567–568 (2020).
9. Straka, J. R. et al. "We Used to Say Rats Fell from the Sky After a Flood": Temporary recovery of muskrat following ice jams in the Peace-Athabasca Delta. Arctic 71, 218–228 (2018).
10. Ward, E. M. & Gorelick, S. M. Drying drives decline in muskrat population in the Peace-Athabasca Delta, Canada. Environ. Res. Lett. 13, 124026 (2018).
11. Ward, E. M., Wysong, K. & Gorelick, S. M. Drying landscape and interannual herbivory-driven habitat degradation control semiaquatic mammal population dynamics. Ecolography 13, (2020) https://doi.org/10.1002/eco.2169.
12. Westworth, D. A. Ecology of the Muskrat on the Peace-Athabasca Delta, Wood Buffalo National Park. (Department of Zoology, The University of Alberta, 1974). https://doi.org/10.7939/R3NZ8113R.
13. Errington, P. L. Muskrat Populations. (The Iowa State University Press, 1963). ISBN: 978-0-59836-540-8.
14. Laurence, S. et al. Isolation and characterization of polymorphic microsatellite loci in muskrat, Ondatra zibethicus. Mol. Ecol. Resour. 9, 654-657 (2009).
15. Do, C. et al. NeEstimator v2: re-implemention of software for the estimation of contemporary effective population size (Ne) from genetic data. Mol. Ecol. Resour. 14, 209–214 (2014).
16. Kalinowski, S. T., Wagner, A. P. & Taper, M. L. ML-RELATE: a computer program for maximum likelihood estimation of relatedness and relationship. Mol. Ecol. Notes 6, 576–579 (2006).
17. Jones, O. R. & Wang, J. COLONY: a program for parentage and sibship inference from multilocus genotype data. Mol. Ecol. Resour. 10, 551–555 (2010).
18. Wang, J. COANCESTRY: a program for simulating, estimating and analysing relatedness and inbreeding coefficients. Mol. Ecol. Resour. 11, 141–145 (2011).
19. Pritchard, J. K., Stephens, M. & Donnelly, P. Inference of Population Structure Using Multilocus Genotype Data. Genetics 155, 945–959 (2000).
20. Excoffier, L. & Lischer, H. E. L. Arlequin suite ver 3.5: a new series of programs to perform population genetics analyses under Linux and Windows. Mol. Ecol. Resour. 10, 564–567 (2010).

21. Rozas, J. et al. DnaSP 6: DNA Sequence Polymorphism Analysis of Large Data Sets. Mol. Biol. Evol. 34, 3299–3302 (2017).

22. Leigh, J. W. & Bryant, D. POPART: full-feature software for haplotype network construction. Methods Ecol. Evol. 6, 1110–1116 (2015).

23. Kiflawi, A., Behar, M., Huzurbaz, V., Holloran, J. M. & Miller, S. N. Identifying Greater Sage-Grouse source and sink habitats for conservation planning in an energy development landscape. Ecol. Appl. 25, 968–990 (2015).

24. Roberts, N. M. & Grimmings, S. M. Do Trends in Muskrat Harvest Indicate Widespread Population Declines? Northeast. Nat. 17, 229–238 (2010).

25. Brietzke, C. Muskrat Ecology in the Mackenzie Delta: Insights from Local Knowledge and Ecological Field Surveys. Arctic 68, 527–531 (2015).

26. Ahlers, A. A. & Heske, E. J. Empirical evidence for declines in muskrat populations across the United States: Muskrat Population Declines. J. Wildl. Manag. 81, 1408–1416 (2017).

27. Rosenberg, K. V. et al. Decline of the North American avifauna. Science 366, 120–124 (2019).

28. Timoney, K. P. The Peace-Athabasca Delta: Portrait of a Dynamic Ecosystem. (University of Alberta Press, 2013). ISBN: 978-0-88864-603-3.

29. Hebert, C. E. et al. Mercury Trends in Colonial Waterbird Eggs Downstream of the Oil Sands Region of Alberta, Canada. Environ. Sci. Technol. 47, 11785–11792 (2013).

30. Hebert, C. E. The river runs through it: The Athabasca River delivers mercury to aquatic birds breeding far downstream. PLOS ONE 14, (2019) https://doi.org/10.1371/journal.pone.0260192.

31. Schumaker, N. H. & Brookes, A. HexSim: a modeling environment for ecology and conservation. Landsc. Ecol. 33, (2018) https://doi.org/10.1007/s10463-017-0605-9.

32. Ward, F. A. Agent-based model of muskrat population in a large northern river delta. (2021) https://doi.org/10.17605/OSF.IO/CHNR6.

33. Sather, J. H. Biology of the Great Plains Muskrat in Nebraska. Wildl. Monogr. 2, 1–35 (1958).

34. Artimo, A. The dispersal and acclimatization of the muskrat, Ondatra zibethicus (L.), in Finland. Ristitieteilijätä Suomenlaista 21, 1–101 (1960).

35. Lay, D. W. Muskrat Investigations in Texas. J. Wildl. Manag. 9, 56–76 (1945).

36. Ahlers, A. A., Schooley, R. L., Heske, E. J. & Mitchell, M. A. Effects of flooding and riparian buffers on survival of muskrats (Ondatra zibethicus) across a flashiness gradient. Can. J. Zool. 88, 1011–1020 (2010).

37. Bellrose, F. C. & Low, J. B. The influence of flood and low water levels on the survival of Muskrats. J. Mammal. 24, 173–188 (1943).

38. Pulliam, H. R. & Danielson, B. J. Sources, sinks and habitat selection: a simulation study. J. Wildl. Manag. 68, 2611–2620 (2005).

39. Earl, D. A. & von Holdt, B. M. STRUCTURE HARVESTER: a website and program for visualizing STRUCTURE output and implementing the Evanno method. Conserv. Genet. Resour. 4, 359–361 (2012).

40. Kopelman, N. M., Mayzel, J., Jakobsson, M., Rosenberg, N. A. & Mayrose, I. CLUMPAK: a program for identifying clustering modes and packaging population structure inferences across K. Mol. Ecol. Resour. 15, 1179–1191 (2015).

41. Waples, R. S. Bias correction for estimates of effective population size based on linkage disequilibrium at unlinked gene loci*. Conserv. Genet. 7, 167–184 (2006).

42. Waples, R. S. & Do, C. Linkage disequilibrium estimates of contemporary N e using haplotype variable genetic markers: a largely untapped resource for applied conservation and evolution. Evol. Appl. 3, 244–262 (2010).

43. Wang, J. & Whitlock, M. C. Estimating effective population size and migration rates from genetic samples over space and time. Genetics 163, 429–446 (2003).

44. Gilbert, K. J. & Whitlock, M. C. Evaluating methods for estimating local effective population size with and without migration. Evolution 69, 2154–2166 (2015).

Acknowledgements

We are grateful for financial support through an Environmental Venture Project Grant from the Stanford Woods Institute for the Environment. We thank Larry Marten, Archie Antoine, Ross Baumann, and Robert Grandjambe Jr. for donating muskrat tissue samples used in the genetic portion of this study. We thank the Mikisew Cree First Nation (MCFN), Athabasca Chipewyan First Nation, and Fort Chipewyan Metis Local 125 for helpful discussions about the delta ecosystem and for permission to work on this land; we thank Melody Lepine at the MCFN GIO office in Fort McMurray for expertise and advice; Jocelyn Marten at the MCFN GIO office in Fort Chipewyan for expertise and assistance in publicizing the 2016 genetics sample collection effort; and Queenie Gray, Jason Straka, and Jess Lankshrief, Parks Canada staff in Fort Chipewyan, for logistical support for muskrat tissue sample collection in Fort Chipewyan. We thank Stuart Macmillan from Parks Canada and Melody Lepine for support in seeking grant assistance for the project. We thank Elizabeth and Philip Ward for muskrat sampling equipment support in Canada. We thank Gili Greenbaum for advising us on microsatellite analyses. Further graduate student support was obtained through a George P. Schultz Fellowship in Canadian Studies from the Stanford Freeman Spogli Institute for International Studies. We are also grateful for support by the Stanford Graduate Fellowship program and travel support from a McGeek Grant from Stanford University’s School of Earth, Energy, and Environmental Sciences.

Author contributions

E.M.W., K.A.S., A.V., S.M.G., and E.A.H. designed the project; E.M.W. developed the agent model for muskrat in the delta and carried out the first year of muskrat tissue collection. A.V., E.M.W., and K.A.S. carried out the second year of genetic sample collection. A.V. generated the first round of microsatellite and cytochrome b data from 2015 samples, and conducted preliminary data analysis; K.A.S. generated all genetic data for 2016 samples and the second round of microsatellite data for 2015 samples and conducted all final genetic analyses; S.M.G. and E.A.H. provided expertise and advice on agent model development, genetics analysis and review of results; E.M.W. wrote the first draft of the manuscript except for the genetics method section, written by K.A.S.; all authors contributed substantially to all revisions and figure development.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s42003-021-02288-7.

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Peer review information Communications Biology thanks the anonymous reviewers for their contribution to the peer review of this work. Primary Handling Editors: Michelle Laving and Luke R. Grinham.

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